By heating benzole said, NaCl, and K.S.O. at AcCl in the cold, and then distilling the mixture. 200° (Beketoff, A. 109, 256).—6. By the action of The isonitroso-acetophenone need not be sepa-tock, on benzene in presence of Al.Cl. (Friedel, rately prepared, but amyl nitrite (1 mol.) can Crafts, a. Ador, B. 10, 1855).—7. From benzoic acid or benzoates and S.Cl. (Carius, A. 106, 800) .- 8. By heating henzoic acid with ZuCl. and adding benzotrichloride (D. P. J. 239, 157).

C.H.CCl₁ + C.H., CO.H - 2C.H., COCl + HCl.

Properties. - Colonriess pungent oil. Decomposed into HCl and benzoic acid slowly by cold, quickly by hot, water. Alcohol reacts vigorously, forming benzoic ether and HCl. Ether and CS.

dissolve it without decomposition.

Reactions. - 1. Aqueous KOH gives KOBz and KCl. -2. Dry BaO at 150° forms Bz.O (Gal, A. 128, 127) .- 3. Dry NII, or ammonium carbonate forms benzamide; other bases not similarly. -4. Sodium has no action in the cold, but in presence of other at 100°, 'dibenzoyl' [140'] is slowly formed (Briegel, Bl. [2] 5, 278). 5. Hydride of Copper forms Cu,Cl. and benzoic aldehyde (Chicaza, A. 85, 232), 6. KI forms BzI.-7. HgCy, forms BzCy - 8. KSCN forms benzonitrile, CO., and CS. (Limpricht, A. 99, 117).—9. Pb(SCN) forms B2SCN. - 10. KNCO forms benzonitrile (Schiff, A. 101, 93) and cynphenine (Clocz, Iil. 1859, 100). -11. NaOBz gives Bz.O (Gerhardt). -12. Sodium formate forms CO, NaCl, and benzoic acid .- 13. Potassium oxalate forms Bz O. KCl, CO, and CO .. -14. By the dry nitrates of Pb, Ag, Hg, or Cu, it is converted into benzoic anhydride, with formation of the chloride of the metal, N.O. and oxygen (Luchowicz, B. 18, 2990). 15. Conc. H.SO, forms, apparently, Bz.SO, II, which on henting becomes benzene-supphonic acid (Oppenheim, Z. [2, 7, 21). 16. PCl, at 200° gives C.H., CCl., C.H., Cl.CCl., C.H., Cl., CCl., and at a higher temperature CCl, and chlorimated benzenes (Schischkoff a. Rösing, J. 1858, 279; Limpricht, A. 134, 55; Claus n. Hoch, B. 19, 1194).—17. KHS forms BzSH.—18. BaO, gives Bz.O. -19. S dium amalgam in acid solution torms benzoic aldehyde and benzyl alcohol (Lippmann, A. 137, 252). -20. KN11, forms benzamide and dibenzamide (Bannert a. Landolt, A. 111, 1) .- 21. Succinic ether at 200° gives succinic anhydride, EtOBz, and EtCl (Kraut, A. 137, 254).

Combination. - TiCl, BzCl. [652]. Yellow erystals (Bertrand, Bl. [2] 34, 631).

BENZOYL-CHLORO-TOLUIDE v. Culoso-TOLUDINE.

BENZOYL-CHOLIC ACID v. CHOLIC ACID. BENZOYL-CROTONIC ACID C₁₁11₀O₂ i.e. C₂H₃.CO.C(CH₄):CH.CO₂H. [113°]. Long pointed crystals. Prepared by the action of Al.Cl. on n mixture of benzeno and citraconic unhydride. By alkalis it is resolved into phered ethyl ketono and glyoxylic acid (Pechmann, B. 15, 891).

BENZOYL-CUMIDIC ACID v. PHENYL-XYLYL-METONE DI-CARBONTLIO ACID.

BENZOYL CUMIDINE v. CUMIDINE.
BENZOYL-CYANACETIC ETHER v. CYANG-

BENZOYL-ACETIC KTHER.

BENZOYL CYANIDE C.H., CO.CN. [33°]. (208°). Formed by distilling BzCl with HgCy, or AgOy (Liebig a. Wöhler, A. 3, 267; H. Strecker, 4, 90, 62; Hübner a. Buchka, B. 10, 480; Kolbe, 4. 90. 68; 98, 847). Formed also by mixing isonitroso-acetophenone C.H. CO.CH; NOII with

be allowed to drop slowly into a warm mixture of acetophenone (I mol.) and acetyl obloride (3 mols.), and the product distilled; yield: 65-70 p.o. of the theoretical (Gausen n. Munnsse, B. 20, 2196). Pungent crystalline mass. Decompased slowly by water, more rendily by KOHAq into HOBz and HCN. Funning HCl forms C.H., CO.CO.NH, whence phenyl glyoxyllo acid. Zino and HCl reduce it to benzoic alde. hyde. NII, gives benzamide and NII,CN.
Aniline gives benzamilde. PCI, nupears to January Res Benzammer Poly appears to form C₆H, COL, CN (2212) (Claisen, B. 12, 626). ZuEt₂, diluted with other forms 3 p.e. of benzeymidino' C₂₄H₁₉NO₂₁ [1217], needles (from alcohol); unother product (200°-220°), either contains phenyl othyl ketone or yields that body on oxidation (Frankland a. Louis, C. J. 37, 742).

BENZOYL CYANIDINE P. BENZOYL OYAN-

BENZOYL CYANURATE v. CYANERIO ACID. BENZOYL - CYMENE - SULPHAMIDE v. CYMENE SULPHONIC ACID.

BENZOYL-CYMENOL v. CYMENOL. BENZOYL CYMIDIDE P. CAMIDINE. BENZOYL DESOXALIC ACID P. DESOXALIC

BENZOYL-ISODURENE v. PHENYL TETRA-METHYL-PHENYL DATONES

O-TRI - BENZOYLENE . BENZENE C, H, O, i.e. Ca(CallaCO), [above 360°]. Formed together with methyleno-phthalyl by heating phthelic subviride with malonic ether and sodium acetate, or by the netion of H2SO, on phthalyl acctic acid (Gabriel a. Michael, B. 10, 1557; 11, 1007, 1679; 14, 925). Yellow crystals. Soff-ble in nitrobenzene, nearly insoluble in other solvents. Potash-fusion converts it into phenenyl-tri-benzoic acid O.H.(C.H.CO.H).

DI . BENZOYL . ETHANE v. DI . PHUNYL. ETHYLENE-DI-RETONE.

BENZOYL . ETHYL - ACETIC ACID v. Ben-ZOYL ACETIC ACID.

BENZOYL . ETHYL . ANILINE v. ETHYL. ANILINE.

BENZOYL . ETHYL . BENZENE v. PHENYL ETHYL-PHENYS REPONE.

BENZOYL ETHYL-o-CARBOXYLIC ACID v. PHENYL RUBYL RETONE O-CAUBOXYLIC ACID.

BENZOYL FORMIC ACID v. PHENYL GLY.

OXYLIC ACID. BENZOYL FLUCRIDE CH.CO.F. From HKF, and BzCl (Borodin, A. 126, 60). Pungent liquid; attacks glass. Decomposed by

water into HF and HOBe. DI-BENZOYL-FUMARIC ETHER

CO.Et.CBz:CBz.CO.Et. Formed by the action iodine dissolved in ether upon the disodium compound of di-benzoyl-succinate, CO.Et.CBzNa.CBzNa.CO.Et (Perkin, C. J. 47. 262).

RENZOYL-GALLIC ACID v. GALLIO ACID. BENZOYL-GLYCOCOLL v. Harpenic acid. BENZOYL-GLYCOLLIC ACID v. GLYCOLLIC ACID.

a BENZOYL . ISO . HEXOIC ACID v. Leebutyl-BENZOTL-ACETIO ACID.

BENZOYL HYDRIDE v. BENZOIO ALDEHYDE. DI-BENZOYL-IMIDE v. p. 475.

DI.BENZOYL-INDIGO v. Indigo. BENZOYL IODANILINE v. Iodo-aniline

BENZOYL IODIDE C.H., CO.I. Easily-fusion ble crystalline mass obtained by heating BzCl with KI (Liebig a. Wölder, A. 3, 260).

BENZOYL-ISATIN v. ISATIN.

BENZOYL-ISETHIONIC ACID v. ISETHIONIO

AC(I) BENZOYL-LACTIC ACID v. LACTIC ACID.

BENZOYL-LEUCINE o. LEUCINE. TRI-BENZOYL-MELAMINE v. MELAMINE.

BENZOYL . TRIMELLITIC ACID. Benzo. phenone tricarborylie acid C. II. O. i.e. C₀11₂.CO.C₀11₂(CO₂11)₃ [5:1:2:1]. From phenyl ψ-cumyl ketone by oxidation with dilute HNO, or KMnO, (Elbs, J. pr. [2] 35, 494). Salts. BatlA".

BENZOYL-MESIDIDE v. MESIDINE.

BENZOYL-MESITYLENE v. PHENYL TRI-METHYL PHENYL KETONE.

Dibenzoyl · mesitylene v. Dr · PHENYL TRI-METHYL-PHENYLENE DIRECTORS.

Tri-benzoyi-mesitylene Call ... O. i.s. (C.H.,Ct)), C.Me., Tri-phenyl tri-methyl-phe-menyl tri-kelone. [216]. Formed by heating benzoyl mesitylene or di-benzoyl-mesitylene with BzCl and Al,Cl, at 198°. Crystals (from ulcohol), v. sl. sol. cold absolut, v. sol. a mixture of chloroform and acctone. When BzCl acts on mesitylene in presence of Al₂Cl₄ below 118⁵ only benzoyl-mesitylene is formed; at 150° dibenzayl-mesitylene is the shief product (Louiso, C. R. 98, 1440; A. Ch. [6] C. 237).

O-BENZOYL MESITYLENIC ACID C., II., O.

i.e. Calla.CO.CallaMe.COOH. Phenyl rylyl ketone carborytic acid. [185]. Prepared in the same way as the p-acid (r. infra) (Louise, Rt. [2] 44, 418). Colouriers crystals, Asol. cold water, sl. sol, boiling water, sol, CHCl, meetore, ether, and benzene. Its salts do not crystallise well. - AgA'. CuA'

p-Benzoyl-mesitylenie acid C, H, O, [160]. Prepared by oxidising phonyl tri methyl phonyl ketone (benzoyl-mesitylene) (Louise, Bl. [2] 44, 418; A. Ch. [6] 6, 218). Nacroous scales, very sol. ether, CHCl, acetono, &c., sol. boiling water.

Salts. A'NH,: small brilliant crystals. -A'Ag; white pp. sol. boiling water. A'Ba 2aq; long needles. A'.Ca: long white lilaments. A'Mg 6aq: crystals, sol. hot water.—SrA'.

BENZOYL METHANE C. ACETOPHENONE Di-benzoyl-methane c. Di-PHENYL METHYLENE DIRE FORK.

zoyl-methane (Baeyer & Perkin, B. 16, 2135; C. J. 47, 210). Small needles. Sublimable. V. 41, sol. alcohol, v. sol. dilute alcoholic KOH. Converted by NaOEt and BzCl into a substance {260°-270

BENZOYL METHYLAMINE C. METHYL.

BENZOYL METHYL-ANILINE o. METHYL-ARIT.INP

BENZOYL - DIMETHYLANILINE r. Di-MRTHYL-AMIDO-BENZOPHENONE

BENZOYL-TRIMETHYLENE C. PRENTI. TRI-METUYLLNE KETONE.

DI . S. BESZOYL - DI - METHYL . MALOWIC AGID (C.H., CO.CH.), C(CO.H). Di-phenacyl-malonic acid. [1849]. Formed by saponifi-cation of its ether, which is obtained by the action of w-bromo-aceto-phonone npon sodio-malonio ether. Large colourless prisms. V. sol. alcohol, ether, and acetic acid, sl. sol. water, insol. benzone and ligroin. Reacts with phonyl-hydrazine. Evolves CO2 on heating, giving di-benzoylisobutyric neid .- "A"K2: white needles or plates,

v. sol. water.—"A M₂: nearly insol. white pp.
Diethyl ether A"Et.: [119°]; large white
glistening prisms or long flat needles; v. sol. water, benzene, neetic acid, and CS, less in alcohol, insol. ligroin. Reacts with phenyllydrazine but not with hydroxylamine (Kues a. Paal, B. 19, 3144).

BENZOYL-METHYL-p-NITRANILINE v. p-

NITRO-PRENYL-W-AMIDO-ACETOPHENONE. BENZOYL - METHYL - PHENYL - NITROS -AMINE v. PHENYL-AMIDO-ACETOPHENONE.

BENZOYL-NAPHTHALIDE v. NAPHTHYL-

BENZOYL - NAPHTHYLAMINE - IMIDE -CHLORIDE v. w. CHLORO-BENZYLIDINE-NAPHTHYL-AMINE

BENZOYL-NAPHTHYL-THIO-UREA v. NAPH-THYL-THIO-DREA

BENZOYL-NITRANILIDE v. NITRO-ANILINE. BENZOYL-NITRITE Bano, (?). formed together with m-nitro-benzoic aldehyde by the action of 20 vols, of a mixture of HNO. (i vol.) and H.SO, (2 vols.) apon I vol. of benżoie aldehydo (Lippinann a. Hawliczek, B. 9, 1463). It is decomposed by distillation.

BENZOYL - NITRO - AMIDO - DIPHENYL v. NITHO-AMIDO-DIPHENYL

BENZOYL . NITRO . AMIDO . PHENOL v.

NITRO-AMIDO-PHENOL. BENZOYL NITRO-ANISIDINE v. NITRO-AMIDO-PHENOL

BENZOYL NITRO - CUMIDINE v. NITRO .

BENZOYL . NITRO - NAPHTHALIDE NUTRO-NAPHTHYLAMINE.

BENZOYL . NITRO . DIPHENYLAMIDE v. NITRO-DIPHENYLAMINE.

BENZOYL NITRO TOLUENE SULPHAMIDE NITRO-TOLUENE SULPHONIC ACID.

BENZOYL-NITRO-TOLUIDE P. NITRO-TOLU-IDINE

BENZOYL PEROXIDE C, H,O, i.e. Bz,O, [101"]. BzCl is mixed with hydrated BaO, and the resulting solid cake washed with water and No.CO₃, and crystallised from CS₂ (Brodie, Pr. 9, 361; 12, 655; Sperlich a. Lippmann, Sits. Tri-bensoyl-methane (C.H., CO), C.H. Mo. 9, 361; 12, 655; Sperlich a. Lippmann, Sitz. thenyt tri-phenyl tri-ketone [225]. Formed by B. 62, 613). Trimetric crystila, insol. water, v. the action of benzoyl-chlorido on sodio-di-ben sol. ether and benzene. Decomposed by heat, giving off CO2 with slight explosion. Boiling hold of forms O and KOBz. Benzoyl peroxide acts as an oxidising agent, splitting up into Bz O and O: thus it oxidises p toluidine to tolucue azo toluene

BENZOYL-PHENOL C.H.,OBz ". PHENOL:

C,II, CO.C. H,OH r. OXY-BENZOPHENONE.
BENZOYL PHENOL SULPHONIC ACID p. PRENOL SULPHONIC ACID.

BENZOYL-PHENYL-AMINE

C.H., CO.C.H., NH. Benzanitide (18 c.), BzCl (14g.), and ZnCl, give the p-benzoyl derivativa [150°], together with a little of the c-benzoyl-

WATTS

DICTIONARY OF CHEMISTR

REVISED AND ENTIL

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ASSISTED BY EMINENT CONTRIBUTORS

IN FOUR VOLUMES VOL. I.

NEW AIMPRESSION

LONGMANS, GREEN, AND CO PATERNOS THE BOW, LONDON BOUBLE AVENUE A COM STREET, NEW YORK BOMBAY, CALCUTE AND MADRAS

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PREFACE.

EWENTY-FIVE years have passed since the publication of the first edition of Watts' Dictionary of Chemistry began, and it is now seven years since the second part of the last supplement was prolished.

Some time before his death, Mr. Watts had agreed to prepare a new edition of its Dictionally, which should give as complete an account of the present state of the science as might be found compatible with the appearance of the book in four volumes of about 750 pages each. Mr. Watts had prepared Instructions to Contributors, and had written sixty-three pages for the new edition, when his death—which all chelinists so deeply lamented—stopped the work. It has fallen to us to take up the task dropped from worthier hands, and to endeavour to bring it to a satisfactory conclusion.

MR. WATTS' MS. is printed very much as he left it, subject only to some necessary condensation. In preparing a new edition we have found it necessary to rewrite the whole book. Our instructions were that we should give as complete and satisfactory an account of the present state of chemical science as we could, consistently with the size to which we were required to confine the book. We have been obliged, therefore, to adopt a very condensed style; the descriptions of individual bodies are given in few words, abbreviations are freely used, and formula are frequently employed instead of names in order to save space.

The original edition was called 'A Dictionary of Chemistry and the Allied Branches of other Sciences; the new edition deals with chemistry only. Considerable space was devoted in the original work to processes of chemical technology, the new edition gives no special information with regard to these matters. Technical chemistry will be treated in a companion volume to be published under the editorship of Professor Thorpe. The great importance of the application of physical methods to chemical questions has made it necessary to consider these methods and the results gained by applying them. Hence in our enumeration of the properties of each element and compound we have included those physical constants which are of most importance to the chemist; and we also intend to describe the leading physical methods of investigation employed in chemistry, and to give a short account of the chief results obtained, in an article entitled Physical Methods used in Chemistry. This article will be divided into sections, each of which will be written by a specially qualified author.

After much consideration, it was decided to omit details regarding analytical processes. In certain cases, e.g. Arsenic methods of detection are given rather

fully. But the new edition is not intended for the use of the analyst in the laboratory. A sketch of the principles of analytical chemistry, and some account of the chief classes of analytical methods, are given in the article Analysis. We have been especially anxious to arrange the matter in a methodical manner, so as to make the task of finding the chief facts about any specified body as little laborious as possible. Cross-references are freely used.

As mere descriptions of individual bodies in strictly alphabetical order cannot states to give a fair notion of the present position of chemistry, we have supplemented times descriptions by short general articles on classes of elements and compounds, e.g. Alkali Metals, Carbon Group of Elements, Oxides, Hydroxides, Hydrates, and Amines. We have also devoted considerable space to articles on important theories, hypotheses, and principles. Some of these articles may be found to overlap, e.g., Chemical Change and Equilibrium, Chemical; but the great importance of the subjects treated in such articles is, in our opinion, sufficient warrant for devoting much space to their consideration, and for inviting different authors to treat parts of the same subject from different points of view.

One of the editors is responsible for the inorganic and general, and the other for the organic, chemistry in this work. This division was absolutely necessary if the book was to appear in a reasonable time; and moreover the nature and arrangement of a Dictionary enables various writers to co-operate in its production without material injury to the unity of the work.

We have been fortunate in securing the help of many contributors—English, American, and Foreign—whose work and position enable them to speak with authority on the subjects of which they treat.

We have had the advantage of the advice and a sistance of Proff G. Carey Foster, F.R.S., and Dr. W. J. Russell, F.R.S. To these gentlemen, and to all our contributors and abstractors, we return our sincere thanks.

Each editor contributes an introduction to his special part. It is hoped that the reader will not pass over these introductions, as they give the necessary explanations of the plan on which the book has been written. The table of abbreviations used is also important.

H. FORSTER MORLEY.

M. M. PATTISON MUIR.

March, 1888.

INTRODUCTION

TO THE PORTION OF THE BOOK DEALING WITH INORGANIC CHEMISTRY.

Each element is described in its alphabetical position. The account of the element is allowed by accounts of its binary compounds and those compounds which may be called ouble binary, in alphabetical order; e.g. bromides, chlorides, chloriodides, sulphochlorides, c.; but cyanides are placed together in one article. There are also short article on Bromides, Chlorides, Oxides, &c.; and an article is devoted to cach clase of elements, e.g. Likali metals, Carbon Group of elements, &c. Ammonium is treated as an element of ar as the description of the Ammonium compounds is concerned. Each group of salts, with he exception of those mentioned above, is described under one heading; e.g. all carbonates redescribed under the heading Carbonates, all nitrates under the heading Nitrates, and o on. The ealts of any specified metal are not as a rule enumerated in the article devoted to he metal; but in a section of this article is given a short account of the salts of the metal considered as a class. Whon some salts belonging to one class are marked off from the whole, &c.; thus there is an article on Alums, and each alum is described in the article Sulphates.

The nomenclature adopted is generally that used in the Journal of the Chemical Society, but it has not been thought expedient to attempt great strictness in this department. Structural formulæ are seldom used for inorganic compounds.

The term molecular weight is generally used only of those elements and compounds which have been gasified, and the specific gravities of which in the gaseoue state have been determined.

• The term valency ie only applied to atoms, and is used to denote the maximum number of atoms of hydrogen, fluorine, chlorine, bromine, or iodine with which one atom of a epecified element is known to combine to form a gaseous molecule.

The symbol Aq is employed to denote an indefinite quantity of water; when Aq is added to the symbol of an element or compound it means an aqueoue colution of this body.

The following gentlemen have been so good as to prepare abstracte of the paper dealing with inorganic chemistry which have appeared in the various journals since the publication of the last supplement to the first edition of this Dictionary:—Messrs Coeno I. Burton, William Burton, G. J. Hill, H. A. Lawrance, Chas. Slates, and Alfred E. Tutton, I am much indebted to these gentlemen, and also to Miss Ida Freund, Lecturer in Chemistry at Newmann College, Cambridgo, who prepared a translation of Prof. Octwald's article on Affinity, and I beg to tender them my best thanks.

INTRODUCTION

TO THE ARTICLES RELATING TO ORGANIC CHEMISTRY.

ORGANIO chemistry probably includes a greater number of observed phenomena than any other science; it is, clearly, not possible to arrange the description of these in such a way that any one, ignorant of the method of arrangement, could readily obtain the information he required. The reader is therefore requested to look through this introduction before referring to any of the organic articles.

The general idea is to devote a separate article to each compound and to arrange these articles in strictly alphabetical order; exceptions are made in the salts of acids and of bases, the ethers, chlorides, amides, anilides, and anhydrides of acids, the acetyl and benzoyl derivatives of compounds containing hydroxyl (OH), amidogen (NH₂), or imidogen (NH), the alkyl derivatives (ethers) of compounds containing hydroxyl, and the oxims and hydrazides of ketones and aldehydes; all these are described in the same article as the parent substance.

The headings of separate articles are in thick BLACK CAPITALS, the salts are in spaced type, the alkyl and alkoyl derivatives are in spaced italics; derivatives of derivatives are in spaced type. Subsidiary articles are in black type. In describing a compound the physical constants (e.g. melting-point, boiling-point, solubility, refractive index) are first given, then follow the modes of formation and preparation of the body, then such properties as cannot be expressed numerically, and finally a list of the chief reactions in which it plays a part. Inasmuch as organic substances are chiefly characterised by their melting or boiling points, it has been thought desirable to give these immediatly after the name and formula of each compound, so that they may be most readily found. The melting-points are inclosed in square brackets, the boiling-points in round brackets. The modes by which salts, ethers, acid chlorides, and amides are formed from the parent acid are only given in particular cases or when the method used is not general; a similar remark applies to the acetyl- and benzoyl-derivatives of compounds containing hydroxyl or amidogen, and to the oxims and hydrazides of ketones and aldehydes. Information on the preparation and properties of such derivatives will be found in general articles.

Nomenclature.

Constitutional names are usually employed, except when the constitution of a body is doubtful; cross-references will be found under the trivial names. Many trivial names that have been almost universally adopted are nevertheless retained, e.g. aniline, aspartic acid, cinnamic acid, pyrocatechin, hydroquinone, resorcin.

The names of hydrocarbons usually end in ene or ane, of phenols in ol, of bases in ine, and a indifferent bodies in in.

In naming several substituting alkyls, that with less carbon comes first, and when there is an equal number of carbon atoms the unsaturated alkyl comes before the saturated: e.g. mathlyl-ethyl-succinic acid; phenyl-naphthyl-amine; allyl-propyl-malonic acid. Radicles containing a blosed ring, however, precede fatty radicles, unless there is great danger of ambiguity; in the latter case cross-references will be given.

Ethers, acetyl and benzoyl derivatives of hydroxylic compounds are placed under the parent substance. Thus anisole and phenyl acetate are described under 'Phenol,' as its methyl ether and acetyl derivative respectively. So also methoxy-benzaldehyde is described under 'Oxy-benzoic aldehyde' as its methyl derivative.

Tetra-alkylated ammonium compensate are usually described under the tertiary ammeriron which they are derived. This phenyl tri-methyl ammonium iodide is described under 'di-methyl-aniline' as ite methyl iodide.

Acetyl and banzoyl derivatives of Ameines are described under the amines to which they belong, thus acetanilide is described under 'Aniline' as its acetyl derivative. Derivatives of aniline, methylamine, &c., containing other alkoyls are usually described as the anilide, methylamide, &c., of the acid from which they are derived; thus C₈H₂.SO₂.NEtH is described as the ethylamide of 'Benzene sulphonic acid.'

Sulphonic and carboxylic acids (whenever they are so named) are represented as derivatives of the hydrocarbon, not of the radicle, thus $C_2H_2(CO_2H)_4$ is called ethane tetracarboxylic acid, not acetyrene tetra-carboxylic acid; and $C_2H_4(SO_3H)_2$ is called ethane disulphonic acid, not ethylene disulphonic acid.

When a compound contains several substituents they are named in the following order: Chloro-, Bromo-, Iodo-, Cyano-, Nitro-, Oxy-, Amido-, Sulpho-, Carboxy-. In choosing the naming group (i.e. the group that is not to be represented as a substituent, but in the termination of the name) the following is the order of preference: CO₂H, SO₂H, CHO, SH, OH and NH₂. Amidogen has precedence over hydroxyl in fatty compounds, but the reverse is the case with aromatic compounds; thus we say oxy-propyl-amine, but amido-phenol.

Examples: chloro-bromo-phenol, not bromo-chloro-phenol; chloro-nitro-oxy-benzoic acid, not nitro-oxy-benzoic acid, nor nitro-oxy-chloro-benzoic acid, nor oxy-chloro-benzoic acid, nor oxy-nitro-benzoic acid, nor chloro-oxy-nitro-benzoic acid; eulpho-benzoic acid, not carboxy-benzenc sulphonic acid; amido-phenyl mercaptan, not sulphydro-phenyl-amine, nor sulphydro-aniline.

Prefixes indicating position.

The latters ω , a, β , γ , &c., are employed to denote the position of substituents in an open chain of carbon atoms. If the substituent is attached to the terminal carbon atom it is preceded by ω , while a, β , γ , indicate its attachment to the first, second, or third, atom of carbon reckoned along the chain from the terminal atom. There are at least two ends to an open chain; the end to be reckoned terminal is determined by the nature of the compound: in monobasic acids it is the carboxyl, in alcohols the group CH_2OH , and in general the group represented in the termination of the name. Thus $CH_2CL.CHI.CHBr.CO_2H$ is called γ -chloro-a-bromo- β -iodo-butyric acid.

When a, β, γ , &c., are used in any other sense than that just explained, they are inclosed between brackets; e.g. (β)-naphthol.

Exo- indicates substitution in an open chain, Eso- denotes substitution in a ring; these prefixes are used when the exact position of the substituent is unknown. The prefixes o-, m-, p-, (ortho, meta, para) indicate isomerism of the di-derivative of benzene (v. p. 454); s- and u- are employed as contractions for symmetrical and unsymmetrical. Thus s-diphenyl-ethane is C_0H_3 .CH₂.CH₂.CH₂.CH₂, while u-di-phenyl-ethane is $(C_0H_3)_0$ CH.CH₃.

In derivatives of quinoline (B.) signifies the benzene ring and (Py.) the pyridine ring. In anthracene, accidines, and azines (B.) signifies the benzene ringe, (A.) denotes the sentral ring.

Alphabetical Order.

In determining the alphabetical order, the following prefixes are discarded: monodi-, tri-, tetra-, penta-, hexa-, hepta-, octo-, &c., per-, ortho-, meta-, pana-, poly-, exo-, eto-, prim-, sec., tert-, iso-, pseudo-, allo-, a-, β -, γ -, ω -, ν -, n-, o-, m-, p-, ψ -, s-, c-, u-, i-, (B)-, (Py.), (A.)-, and all numbers. Of course when the entire name is numeral, e.g. hexadecane, hexane, &c., this rule does not hold. Thue di-bromo-benzene is in the same article as bromo-benzene; paraldehyde is associated with aldehyde, isobutyric acid with n-butyric acid, &c. The prefixes pyro- and proto- do not belong to this class.

The presence or absence of hyphens between parts of a name in no way affects its alphabetical position; thus 'Benzylidene' precedes Benzyl iodide.'

Formulæ.

Formulæ, to save space, are written as much as possible in one line. A pormuo of a formulæ inclosed in brackets is usually supposed to represent a group

INTRODUCTION.

of atoms more intimately connected with the groups represented by the preceding symbols, which are not in brackets, than with those following, e.g. $CH_1(CO_2H).CH_1.CO_2H$ is succinio acid. When numbers within square brackets follow a formula they refer to the substituents taken in the order in which they occur in the formula: thus $C_0H_3Br(NO_2)(CO_2H)[1:2:0]$ is used as an abbreviation for $C_0H_0Br(NO_2)(CO_2H)[Br:NO_2:CO_2H_0E:0]$. The system here adopted differs, therefore from that sometimes employed, according to which the above symbol would mean $C_0H_3Br(NO_2)(CO_2H)[CO_2H:Br:NO_2=1:2:6]$. Constitutional formula are looked upon by the majority of chemists as nothing more than a short way of indicating which atoms in a molecule are directly combined, and which are only indirectly combined with one another. The followers of Van 't Hoff and Wislicenus, however, suppose that constitutional formulæ can be constructed in the form of solid figures which give some notion of the actual relative positions of the atoms in a molecule. All agree that it is by the use of constitutional formulæ that the remarkable development of organic chemistry has been made, and that they cannot be abandoned until something better can be found to take their place.

It is not possible to find space for discussing the reasons which have led to the adoption of each constitutional formula; where these reasons are not given, a careful consideration of the methods of formation and the reactions of the compound will probably reveal them.

Special Articles.

In a few articles a number of compounds are grouped together, in violation of the foregoing rules. The longest of these are the articles on 'azo-' compounds. Other such articles are on the anunonia derivatives of 'Benzoic aldehyde,' on 'Benzil,' on the organic derivatives of 'Antimony,' 'Arsenic,' and 'Bismuth,' on 'Camphor' and on 'Cellulose.' The following general articles, amongst others, will also be found in this volume: 'Acids,' 'Alcohols,' 'Aldehydes,' 'Alkaloids,' action of 'Aluminium chloride,' 'Amides,' 'Amide,' 'Amides,' 'Armines,' 'Analysis,' 'Anhydrides,' 'Aromatic Series' (see also 'Benzene'), 'Azo-colouring matters,' 'Diazo-compounds,' and 'Bromo-compounds,'

Contracted Expressions.

Since the date to which Watts had brought the record of chemical discovery, the number of organic compounds known has doubled, nevertheless the space allotted to them in the present dictionary is little more than a quarter of that devoted to organic chemistry in the original dictionary and its supplements. It is evident that there must be extreme compression, and this compels the free use of abbreviated expressions; it is hoped, however, that a reader who has once made himself acquainted with the nature of these abbreviations will find that they are very convenient. In the first place, the symbols of a fow common reagents are used in the text with purely qualitative meaning, although when connected in an equation they are used in the ordinary sense. The great saving of space (about 200 pages) has compelled the use of this convention, which would be reprehensible under any other circumstances. The use of the contractions 'v. sl. sol.;' 'sl. sol.,' 'in. sol.,' 'v. sol.,' 'v. e. sol.,' and 'sol.,' for 'very slightly soluble in,' 'slightly soluble in,' 'moderately soluble in,' 'very soluble in,' 'very easily soluble in,' and 'soluble in,' cnables solubilities to be given in the case of many hundred compounds where space would otherwise have compelled their omission. Of course these terms are vague; where numerical data have been determined, they are usually given in the dictionary, preceded by the letter S. Particular attention should be paid to the exact meaning of these numbers; they denote the rumber of grammes of a liquid or solid dissolved by a hundred grammes of the solvent, but the number of volumes of a gas dissolved by one volume of the solvent. Soluble, used as an adjective, the menstruum not being named, means soluble in water.

Constants.

Numerical constants are not given in the form $a+bt+ct^2$, &c., since such expressions not only take up a great deal of room, but are usually worthless, because slight errors of experiment produce an enormous effect upon the constants b, c, &c.; in such cases one of two actual observations, of a kind likely to be useful in identifying the substance, have usually been selected.

It is unfortunate that there is a want of uniformity among authors in the method of recording physical constants. Specific gravities are given by most authors without any mention of the temperature of the water that is taken as standard. Some take water at 0°. some at 4°, and others compare the substance with water at the same temperature as itself. Taking the specific gravity of water at 4° as unity, that at 21° will be 998; that is to say, for a Substance whose specific gravity is about 1 we may make on error of '002 by assuming that the author used water at 4° as a standard, whereas he really used water at 21°. Under such circumstances it would be preposterous to give four places of decimals, and such indefinite specific gravities have been cut down to three decimal places, and even then the last figure is somewhat doubtful.

Heats of formation are usually calculated on the assumption that the heat of formation of 44 g. of carbonic acid is 96,960, and that of 18 grms. of water is 68,860; Stohmann, Rodatz, and Herzberg, however, use 94,000 and 69,000 respectively, hence their heats of formation are not directly comparable with those of other observers.

Molecular refraction is the value of the expression $M(\frac{\mu-1}{d})$, where M is the mole cular weight, μ the index of refraction, and d the specific gravity of the liquid at 20 compared with water at 4° (Landolt, P. 123, 595; Brtihl, A. 200, 139). Other constants, such as $\left(\frac{\mu^2-1}{\mu^2+2}\right)\frac{M}{d}$, have also been used; these are of course not comparable with those first mentioned (cf. Brühl, A. 235, 1).

The specific rotation is given by most observers for a tube of liquid 100 mm. long, but many French chemists use a 200 mm. tube as a standard, and some even 50 mm. When the length of tube is stated it is easy to apply the correction, but when, as is often the case, an author does not give the longth of tube, his numbers are indefinite.

The rotation measured for the neutral tint is of course not the same as that measured for the sodium line, yet authors occasionally fail to mention the kind of light employed. The angular rotation ought to be divided by the specific gravity of the liquid during the experiment, in order that the effect of equal weights of material may be compared; yet it is to be feared that many authors neglect to perform this division, and also to mention that they have not done it.

Authors frequently fail to state whether their melting and boiling-points have been corrected for the exposure of part of the stem of the thermometer. This may make a difference of 5°. The immersion of the whole of the mercury in the liquid or vapour is indicated by i.V.

References.

Where the same paper is referred to several time in the course of one article, the full reference is given once, and in other places there will be found the first letter or the first two letters of the author's name, inclosed within brackets; thus, if (Perkin, C. J. 45, 890) and (P.) are found in the same article, the (P.) is a contraction for (Perkin, C. J. 45, 890).

Short Article Expanded.

In order to make sure that the contractions employed are thefoughly understood, a hort specimen article will be expanded by simply exchanging the contractions for their quivalents:-

Bromo-di-exy-bancele acid C.H.Br(OH) CO.H [z:2:6:1]. [184°, anb drous]. From c-di-exy-sanzoid acid in ether and Br (Zehenter, M. 2, 480). Prisms (containing aq); v. sol. alcohol, . al. 10 water. Fe Cl. gives a violet colour to its aqueous solution. AgA'aq. - PAA' 71aq. KA' 1 aq.

May be expanded thus :-

 $\{C_{s}H_{s}Br(OH)_{s}CO_{s}\}_{s}Ba_{s},7\\ \\ \frac{1}{2}H_{s}O_{s} \text{ and } C_{s}H_{s}Br(OH)_{s}CO_{s}K,1\\ \\ \frac{1}{4}H_{s}O.$

Nomenclature of Rings.

Besides the hydrocarbon rings, represented by benner, naphthalend, phenanthrene, CH, anthracene, indonaphthene $C_0H_1 < CH_2 > CH$, tri-methylene $CH_2 - CH_2$, tetra-methylene OH₂—OH₂
i i , penta-methylene CH₂<OH₂·CH₂>, &c., there are a great many rings conOH₂—OH₂ taining other elements. Some of these are collected hore for convenience of reference. It will be noticed that glyoxaline and metapyrazole differ only in regard to the position of one stom of hydrogen. The exact structure of rings containing five or six atoms is not known; some alternative formulæ will be found on p. 446.

Nitrogen ring compounds.

The di-oxy-derivative of the second form of triazol has been named 'Urazole' by Pinner.

and C,H,<CH>N (pseudo)

$$C_{10}H_{\bullet}$$
 \bigvee_{N}^{N} $C_{10}H_{\bullet}$ Nanhthazine.

'Pyrroline' has been used by some authors for Pyrrole-dihydride. 'Pyrroline' in the abstracts in the Journal of the Chemical Society means Pyrrole.

$$\begin{array}{c|cccc} \operatorname{CH_2--CH_2--CH_2} & \operatorname{CH_2--CH_2} & \operatorname{Piperidine.} \\ \operatorname{CH_2-NH--CH_2} & \bullet \\ \operatorname{CH_2-NH--CH_2} & \bullet \\ \operatorname{CH_2-NH--CH_2} & \bullet \\ \end{array}$$

The numbers indicating position in compounds of naphthalene are as follows:--

The positions 1, 4, 1', 4' are termed (a), while 2, 3, 2', 3' are called (3). Quinoline is numbered thus:—

Thus (B. 4)-bromo-(Py-8)-oxy-quinoline would be

Pyridine is numbered thus:

One of the assumptions made by the recent doctrine of tautomerism is that a lactam CO.NH can readily change into a lactim C(OH):N, and that the group CO.CH₂ can change • into C(OH):CH. It is obviously expedient to describe two compounds which are mutually interchangeable, if not identical, in the same article, hence rings containing CO.NH or CO.CH₂ are named as if they were hydroxylic compounds of the form C(OH):N and C(OH):CH.

Lactones and Anhydrides.

Lactones and anhydrides are usually described under the substance from which they may be derived by the abstraction of water; thus, butyro-lactone will be described under oxy-butyrio acid.

Prefixes discarded.

The prefixes home, hydro, and mono are not used. The nyuro-compounds of unsaturated bodies are, if saturated, hamed in the usual way; thus hydro-sinnamic soid is phenyl-propionic soid. The hydro-derivatives of ring compounds are described as hydrides of the simpler compounds from which they are derived: e.g. di-hydro phthalic soid as

phthalic acid dihydrids. Compounds beginning with home- must be re-named; thus home-salicylic acid is oxy-toluic acid.

Hyphens.

Hyphens are placed between each significant part of a name; absence of the hyphen asually indicates close connection between two groups of atoms; e.g. phenylethyleuron is C₂H₂.O₂H₂.NH.CO.NH₂ while phenyl-othyl-ures is C₂H₃.NH.CO.NHC₂H₃.

Ambigueus names.

A number of names have been used in several senses by different authors; it may therefore be well to mention the names chosen in some of these cases. The terms cyanide and isocyanide are altogether discarded, carbanine and nitrile being used instead. Cyanate is used for ordinary potassium cyanate and the ethers that may be derived therefrom; the corresponding sulphur compounds are described as sulphocyanides and thio-carbinides (mustard oils). Cinnamyl is C_0H_5 .CH:CH.CH.2, the acid radicle C_0H_5 .CH:CH.CO being cinnamoyl and C_0H_5 .CH:CH is termed styryl.

Tolyl is used only for CH₃,C₆H₄, and not for benzyl C₆H₅,CH₂, nor for CH₃,C₆H₄,CH₂.

Cresyl is not used as a name. Xylyl is only used for (CH₃)₂C₆H₃., not for CH₃,C₆H₄,CH₂, nor for (CH₃)₂C₆H₃,CH₂.

Durene is used as synonymous with tetramethyl-benzene.

Discarded names.

As it commonly happens that several names have been given to the same compound, it may be well to give a list of the names that have been chosen in a few cases.

Carbamic ether	is	used	instead	of	Urethane
Urea		,,	**		Carbamide
Thio-carbimide		,,	,,		Mustard oil
Tolylene		,,	**		Toluylens
Methyl-pyridine		**	77		Picoline
Di-methyl-pyridino		**	***		Lutidine
Tri-methyl-pyridine		**	**		Collidine
Methyl-thiophene		**	***		Thiotolens
Di-methyl-thiophene		"	**		Thioxene
Oxy-pyridine		,, .	71		Pyridone
Methyl-quinoline		**	"		Quinaldine
Diquinoline		.,	**		Diquinolyl
- hydrazide		**	**		izine
(B. 1)-		1,	11		ana-

Acknowledgments.

I have been fortunate in securing the assistance of Mossrs. A. G. Green, V. H. Veley, G. N. Huntly, E. E. Gravos, Cecil H. Cribb and Cosmo I. Burton, and of Drs. Samuel Rideal and T. A. Lawson. I am also greatly indebted to Mr. A. G. Green for assistance in revising the proof-sheets. Without the assistance of these gentlemen, it would have been impossible to have done anything like justice to the multitudes of original researches that appear every month, and I have therefore great pleasure in publicly thanking them for the zeal they have shown in endeavouring to render the portion of the Dictionary dealing with Organic Chemistry as far as possible complete.

H. FORSTER MORLEY.

INITIALS OF SPECIAL CONTRIBUTORS.

- C. F. C. . | C. F. CROSS, Esq., Consulting Chemist. Contributes Cellulose.
- W. D. . WILLIAM DITTMAR, Ph.D., F.R.S., Professor of Chemistry at Anderson's College, Glasgow. Contributes Analysis.
- A. G. G. ARTHUR G. GREEN, Esq., F.I.C., Research Chemist to the Atlas Works, Hackney Wick. Contributes Diazo-compounds.
- J. J. H. . J. J. HOOD, Esq., D.Sc. Contributes CHEMICAL CHANGE.
- W. D. H. W. D. HALLIBURTON, M.D., B.So. Assistant Professor of Physiology at University College, London. Contributes Blood.
- F. R. J. . FRANCIS R. JAPP, M.A., Ph.D., F.R.S., Assistant Professor of Chemistry at the Normal School of Science, South Kensington. Contributes Benzil, Ammonia derivatives of Benzil, Ammonia derivatives of Benzil, Ammonia derivatives of Benzil, Ammonia
- E. R. L. E. RAY LANKESTER, M.A., F.R.S., Professor of Zoology at University College, London. Contributes Bacteria.
- L. M. . LOTHAR MEYER, Ph.D., Professor of Chemistry in the University of Tubingen.
 Contributes Allotroff.
- R. M. . RAPHAEL MELDOLA, F.R.S., Professor of Chemistry at the Finsbury Technical College. Contributes Azo-colouring matters.
- W. O. . WILHELM OSTWALD, Ph.D., Professor of Physical Chemistry in the Landwirthschaftliches Institut, Leipzig. Contributes Appinity.
- R. T. P. RICHARD T. PLIMPTON, Ph.D., Assistant Professor of Chemistry at University
 College, London. Contributes Amylamines.
- W. R. . WILLIAM RAMSAY, Ph.D., Professor of Chemistry at University College, London Contributes Acros and Alloys.
- C. O'S. . C. O'SULLIVAN, F.I.C., Burton-on-Trent. Contributes Arabic acid, Bassorin, and Creasin.
- T. S. . THOMAS STEVENSON, M.D., Lecturer on Forensic Medicine at Guy's Hospital.

 Contributes Detection and Estimation of Poisonous Alkaloids.
- J. J. T. J. THOMSON, M.A., F.R.S., Professor of Experimental Physics in the University of Cambridge. Contributes Adgregation, STATES OF.
- T. E. T. . T. E. THORPE, Ph.D., F.R.S., Professor of Chemistry at the Royal School of Mines. Contributes Atmosphere.
- R. W. MARINGTON, Esq., F.R.S. Contributes Ash of organic bodies.
- C. J. W. CHARLES J. WILSON, Esq., F.I.C. Contributes Caoutchouc.
- H. W. HENRY WATTS, B.A., F.R.S. (the late). Contributes many special articles, and the

Articles by Mr. MUIR are initialed M.M. P. M. URSIGNED ARTICLES are by Dr. MORLEY.

ABBREVIATIONS

I. JOURNALS AND BOOKS.

When an author has been mentioned in an article, he is usually referred to thereafter in that article by his initial only.

	in that article by his initial only.
d	Liebig's Annalen der Chemie.
A. A.	Annales de la Sociedad Cientifica Argentina.
A. Ch.	Annales de Chimie et de Physique.
P. Am. A.	
Am	American Chemical Journal.
Ann. M.	Annales des Mines.
Am. S	American Journal of Science.
A. C. J.	Journal of the American Chemical Society.
Am. Ch.	American Chemist.
Am. J.	American Journal of Pharmacy.
Pharm.	•
An	The Analyst.
A. Ph. S.	Proceedings of the American Philosophical Society.
Ar. N.	Archives nécrlandaises—The Hague.
Acad.	Mémoires de l'Académie des Sciences.
Ar. Ph	Archiv der Pharmacie.
Ar. Sc	Archives des Sciences phys. et nat.
\overline{B}	Berichte der deutsehen ohemischen Gesellschaft.
B. A	
Bl	Dollar de la Carte d'altre la Danie
* *	
B. B.	Biedermann'e Centfalblatt für Agricultur-Chemie.
B. J	Berzelius' Jahresberichte.
	Berliner Monatsberiohte.
C.S. Mem.	Memoirs of the Chemical Society of London.
C. J	Journal of the Chemical Society of London.
C.J. Proc.	Proceedings of the Chemical Society of London.
C. N	Chemical News.
C. R	Comptes-rendus hebdomadaires des Séances de l'Agadémie des Scieuces
	Paris.
C. C	Chemisches Central-Blatt.
D. P. J.	Dingler's polytechnisches Journal.
Fr.	Fresenius' Zeitschrift für analytische Chemie.
G	Gazzetta chimica italiana.
G. A	Gilbert's Annalen der Physik und Chemio.
\underline{H}	Hoppe-Seyler's Zeitschrift für physiologische Chemie.
<u>r.</u>	Proceedings of the Royal Irish Academy.
J	Jahresbericht über die Fortsehritte der Chemie nnd verwandter Theik
	anderer Wissenschaften.
J. C. T	Jahresbericht für Chemische Technologie.
J. M	Jahrbuch für Mineralogio.
J. de Pla	Journal de Physique et des Sciences accessoires.
J. Ph. • ₹	Journal de Pharmacie et de Chimie.
J. pr.	Journal für praktische Chemie.
J. Th.	Jahresbericht über Thlerchemie.
J. R.	Journal of the Russian Chemical Society.
J. Z	enaische Zeitschrift für Medicin und Naturwissenschaft. Landwirthschaftliche Versuchs-Stationen.
L. V. e	Monatshefte für Chemie und verwandte Theile anderer Wissenschaften.
M.	MODERATE IN CHAMIS WHITE AND AND THE AND
M. D.	Le Moniteur Scientifique.
Mem. 8.	Mémoires de la Société d'Arcuell.
d'A.	Mémoires couronnés par l'Académie de Bruxelles.
Mêm. B.	Tromonia contoution has a position as a series

```
N. Ed.P.J.
N. J. P.
N. B. P.
N. J. T.
N. Z. R.
              New Edinburgh Philosophical Journal.

Neuer Jahre bericht der Pharmacie.
              Neues Repertorium für die Pharmacie.
              Neues Journal von Trommsdorff.
              Nene Zeitschrift für Rübenzuckerindustrie.
              Philosophical Magazine.
Poggendorff's Anualen der Physik und Chemie.
 P. M.
P. M.
P. B.
Pf.
Pr. E.
Ph. C.
              Beiblätter zu den Annalen der Physik und Chemie.
              Pflüger's Archiv für Physiologie.
              Proceedings of the Royal Society of Edinburgh.
              Pharmaceutical Journal and Transactions.
              Pharmaceutisches Central-Blatt.
 Pr. P. R. I.
              Proceedings of the Royal Society.
              Proceedings of the Royal Institution of Great Britain.
 P. Z.
              Pharmaceutische Zeitschrift für Russland.
 R. T. O. .
              Recneil des travaux chimiques des Pays-Bas.
 R. P.
Q. J. S. .
              Repertorium für die Pharmacie.
              Quarterly Journal of Science.
Schweigger's Journal der Physik.
 Scher. J.
              Scherer's Journal der Chemio.
 8. C. I. .
Sits.W. .
              Journal of the Society of Chemical Industry.
Sitzungsberichte der K. Akademie zu Wien.
 T. or Tr.
              Transactions of the Royal Society.
              Transactions of the Royal Society of Edinburgh.
Wiedemann's Annalon der Physik und Chemie.
 T. E.. .
 17.
 W. J.
              Wagner's Jahresbericht.
 z.
              Zeitschrift für Chemie.
Zeit.ang.
              Zeitschrift für angewandte Chemie.
   Oh.
Z. B. .
Z. f. d. g.
Natur-
              Zeitschrift für Biologie.
              Zeitschrift für die gesammten Naturwissenschaften.
   wiss.
Z. K.. .
Z. P. C.
Z. V.. .
              Zeitschrift für Krystallographie nud Mineralogie.
              Zeitschrift für physikalische Chemie.
Zeitschrift des Vereius für die Rübeuzuckerindustrie des deutschen
Bn.
              Handbuch der organischen Chemie: von F. Beilstein, 2te Auflage.
E. P.
              English Patent.
G. P.
              German Patent.
              Gmelin's Handbook of Chemistry-English Edition.
Gm.
Gm.-K. .
              Gmelin-Kraut: Handbneh der anorganischen Chemie.
Gerh.
              Traité de Chimie organique: par Charles Gerhardt.
Lehrbuch der organischen Chemie: von Aug. Kekulé.
              Graham-Otto: Lehrbuch der anorganischen Chemie [5th Ed.]
Stas,
              Stas' Recherches, &c.
   Rech.
                                                    Aronstein's German translation is re-
Sms,
             Stas' Nonvelles Recherches, &c. )
                                                         ferred to as Chem. Proport.
 Nouv. R.
             Thomsen's Thermochemische Untersuchungen.
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II. TERMS AND QUANTITIES, &C., FREQUENTLY USED.

```
Water; e.g. NaOHAq means an aqueous solution of canstic soda.
18 parts by weight of water.
aq
A"
                         Residues of mono-, di-, and tri-basic acids. Thus, in describing the salts of a monobasic acid NaA', CaA', AlA', may be written, HA's tending for the acid. For a dibasic acid we should write Na<sub>2</sub>A'', CaA'', Al<sub>4</sub>A'', &c. Stand for bases of the ammonia type, in describing their salts. Thus the hydrochloride would be B'HCl or B''2HCl, according as the base is
 A#
B' B" &c.
                                    monacid or discid, &c.
                          Concentrated.
conc.
dil.
                         Dilute.
                         gram.
g. .
                         milligram.
mgm.
mnn.
                         millimetre.
mol.
                         molecule
```

```
oil, . .
            liquid, nearly, or quite, insoluble in water.
рp.
             precipitate.
             precipitate.
to ppt. .
            precipitating.
ppg. .
ppd. .
            precipitated.
      *
insol.
            insoluble in.
v. e. eol..
            very easily
v. sol.
                           eoluble in.
m. sol. .
            moderately
            slightly
al. sol.
v. sl. sol.
            very slightly
v. . . .
            sce.
cf. . .
            oompare. .
(°) .
            ahout.
            a melting-point.
(°)
            a boiling-point.
            Hardness (of minerals).
Åt. w.
            Atomio weight.
Mol.w.or
            Molecular weight.
 M. w.
D. . .
            Density.
cor. .
            corrected.
uncor. .
            uncorrected.
i.V.
V.D.
            iu vapour.
            vapour-density, i.e. density of a gas compared with hydrogen or air.
S.G. 19
S.G. 19
S.G. 11
            Specific gravity compared with water.
                             at 10° compared with water at 0°.
                       **
                ,,
                       **
                              ,, 12°; compared with water of which the temperature is
                       ••
                not given.
S.H. .
            Specific heat.
S.H.v.
                      " of a gas at constant volume.
S.H.p.
            Quantity of heat, in gram-units, produced during the complete combustion of the mass of a solid or liquid body represented by its
H.C.
                 formula, taken in grams.
H.C.v.
            Heat of combustion in gram-units of a gram-molecule of an element or
                 compound, when gaseons, under constant volume.
H.C.p.
            The same, under constant pressure.
H.F.
            Quantity of heat, in gram units, produced during the formation of the
mass of a solid or liquid body represented by its formula, taken in
                 grams, from the masses of ite constituent elements expressed by
                 their formulæ, taken in grams.
H.F.v. .
            Heateof formation of a gram-molecule of a gaseous compound from the
                 gram-molecules of its elements under constant volume.
            The same, under constant pressure.
Heat of vaporisation of a liquid, i.e. gram units of heat required to change
H.F.p.
H.V.
                 a gram-molecule of the liquid compound at B.P. into gas at same
                 temperature and pressure.
T.C. . .
            Thermal conductivity (unit to be stated).
S.V. .
            Specific volume; or the molecular weight of a gaseous compound divided
                 by the S.C. of the liquid compound at its boiling-point compared with
                 water at 4°.
3.V.S. .
            Specific volume of a solid; or the mass of the solid expressed by its
                 formula, taken in grams, divided by its S.G.
Electrical conductivity (the unit is stated in each case).
            Coefficient of expansion (between 10° and 20°).
  to 20°)
l (alco-
                                    of a gas = volume dissolved by I volume of water.
            Solnbility in water
                                   of a liquid or solid = number of grms. dissolved by
                       " alcohol 100 grms. of water. In both cases the temperature is stated.
  hol)
            Index of refraction for hydrogen line 8.
            " ", sodium ", de. Molecular refraction for sodium light, i.e. index of refraction for line m
 D, de.
 lo 16. *
                 minus one, multiplied by molecular weight, and divided by S.G. at 150
                 compared with waterat 0.
            The same; S.G. being determined at 150-20° and referred to water at 4°.
            The same for line of infinite wave-length, index being determined by
                Cauchy's formula (Brühl's R.).
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ARBREVIATIONS.

```
( ) .
             Specific rotation for sodium light,
                       ... " neutral tint. [a] = \frac{100}{p} \times \frac{a}{d}. a = observed rotation for
 (0)
                  100 mm. of liquid. d=S.G. of liquid. p=nc. of grammes of active
                  substance in 100 grammes of liquid.
 M, M
             Molecular magnetic rotatory power = d x d x m', where m = molecular
                  weight of the body of S.G. =d, a angle of rotation under magnetic influence, a' angle of rotation of water under same influence, and
                  m' = molocular weight of water (18).
             Acetyl C.H.O.
 Αc
Bz
             Benzoyl C,H,O.
Cy
             Cyanogen CN.
             Ethyl C.H.
Me
             Methyl CH.
 Ph
             Phenyl C.H.
                                               in formula.
             Normal Propyl ClI<sub>2</sub>. CH<sub>2</sub>. CH<sub>3</sub>. Isopropyl CH(CH<sub>3</sub>).
Pr
Pr
R, R' &e.
             Alcohol radiclee or alkyis.
             primary.
prim .
sec
             secondary.
tort
             tertiary.
15 .
             normal.
m, 0, p
             meta-ortho-para.
             consecutive.
             irregular.
             symmetrical.
  .
72
             unsymmetrical.
             pscudo.
  ٠
             attached to nitrogen.
            Employed to denote that the substituent is attached to a oarbon atom
                 which is next, next but one, or next but two, respectively, to the
                 terminal carbon atom. The end to be reckoned from ie determined
                 by the nature of the compound. Thue CH<sub>2</sub>.CHBr.CO<sub>2</sub>H le a-bromo-
             denotes that the element or radicle which followe it is attached to a ter-
                 minal oarbon atom.
a,β,γ,&c.
             indicate position in an open chain, only.
1,2,3,&c.
             indicate position in a ring only.
(a), (β),
             Used when a, B, &c. are employed in a cense different from the above,
  de.
                 e.g. (a)-di-bromo-camphor.
             Baeyer's Nomenclature :
(B.) . (Py.) .
                 benzenc ring.
                 pyridine ring.
                     Thus (B. 1:3) diehloroquinoline, means a meta-dichloroquinoline in
                  which the chlorine atoms are both in the benzono ring.
                     While (Py. 1:3) dichloroquinoline, means a similar body, only the
                 chloring atoms are in the pyridine ring. The numbere are counted
                 from two carbon atoms which are in different rings, but both united
                 to the same carbon atom.
(A.) . .
            denotes the central ring in the molecule of anthracenc, acridines, and
                 azinee.
            meane that the element or radicle it precedes is in a closed ring.
680-
6220-
                                                               not in a benzene ring.
alic.
            denotes isomerism that ie not indicated by ordinary formulæ; thus malele
                 acid may be called allo-fumaric acid.
thio. .
            denotes displacement of oxygen by sulphur.
                    the group SO,H, except in the word sulphocyanide.
sulpho. .
sulphydro-
                    the group SH.
            Tribromonitrobenzene eulphonic acid [1:2:3:4:5] means that the three
                 bromines occupy pocitione 1, 2, and 3; the nitro- group the position 4,
                 and the sulpho- group the position 5.
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analysis. But it by no means follows that formulæ without this mark are those of analysed compounds.

All temperatures are given in degrees Centigrade unless when specially stated otherwise.

Wave-lengths are given in 10.7 mm.

Rermulæ, when used inetcad of names of substances, have a qualitative meaning only.

Exomsen's notation is used in thermoghemical data.

DICTIONARY OF CHEMISTRY

ABIES .- The needles of A. pectinata contain a sugar called Abietite, C.H.O., very much like mannite, but differing therefrom in composition and in solubility. The same plant contains a tannin identical with the solublo tannin of the horse-chestnut, C18H12Oe, and convertible by liydrochlorio acid into an anhydride C52H46O23, insoluble in cold water, but soluble in boiling potash-lye, slightly in water and alcohol (Rochleder, J. pr. 105, 63, 123).—The fruits of Abies Regina Amalia, indigenous in Arcadia, yield, by distillation with water, about 18 p.o. of a colourless volatile oil Cto II115, smelling like lemons, S.G. 868 (156-159°); slightly levorotatory. Resinifies quickly in the air, exerting an ozonising influence stronger than that of turpentine-oil. Dissolves iodine, and absorbs hydrogen chloride, forming a liquid compound C₁₀H₁₆·HCl (Buchner Phiel. J. nr. 92, 109).

H. W. a. Thiel, J. pr. 92, 109).

ABIETENE C.H16.-The heptane of Pinus

sabiniana (v. Heptanes).

ABIETIC ACID C4,He4O5 [139°] or [165°].-Caillot, J. Ph. 16, 436; Maly, A. 129, 94; Emmerling, B. 12, 1441; Kelbe, B. 13, 898.—Occurrence. The clear liquid turpentine of various species of pine contains abietic anhydride $C_{i1}H_{c2}O_{i}$, which, on exposure to the air, absorbs moisture and is converted into abietic acid, the liquid thon coagulating to an opaque granular pulp. The anhydride is the chief constituent

of common resin or columny.

Preparation. - 1. Coarsely pounded colophony is digested for two days with weak spirit; the liquid is decanted from the white crystalline pulp, and squeezed in a press; the press-cake dissolved in hot strong alcohol, and the solution left to itself at ordinary temperatures; a white orystallino crust is thus obtained; the motherliquor, when cooled by ice, usually solidifies to a loose mass of white lamines, which constitutes the greater part of the product. The crystalline crust consists of sylvic acid C20H30O2, tho laminæ of abictic acid (M.).—2. Colophony is digested for two days with spirit of 70 p.c., and the undissolved portion, after washing with weak spirit, is dissolved in the smallest possible quantity of glacial acetic acid. From this solution the acid separates in orusts, and on adding a little water to its solution in hot alcohol and stirring, it is obtained in crystalline scales (E.).—3. Sodalye which has been used for purifying crude resin-oil is mixed with common salt, and the soap which separates is dried at 70°-80°, and prified by exhaustion with ether. The readue for ether; pale green. dissolves in alcohol, and the solution, on evaporation, deposits needle-shaped crystals of sodium posing silver abietate with ethyl iodide diluted states, the aqueous solution of shich yields, with ether, forms a yellowish mass, having an Vol. I.

on addition of hydrochloric acid, a white pp. of abietic acid, which melts to a resinous mass if the mixture is boiled (K.).

Properties. - Separates from hot alcoholic solution in irregular transparent pointed tri-clinio crystals melting at 165° (M., K.); 139° (E.); 135° (Flückiger). Sol. alcohol, ether, cenzene, glacial HOAc, CHCl, and CS.

Reactions .- 1. Abietio acid distilled with sine chloride yields a heavy oil (70°-250°) containing heptylene (E.).—2. Strong hydrochloric and hydrodic acids at 145° abstract the elements of water from it, leaving the anhydride (E.); but when treated in alcoholic solution with gaseous HCl, it yields sylvio and sylvinolio acids:

 $C_{44}H_{34}O_5 + H_2O = C_{20}H_{30}O_2 + C_{24}H_{30}O_4$ (?) Sylvio acid is also formed when a hot alcoholic solution of abietic acid is mixed with sulphnrie acid (M.).-3. Triturated with PCls it yields on distillation a volatile oil C, H, called by Maly abictone, together with HCl and POCl .oxidation with KMnO, abietic acid yields carbonie, acetio and formio acids.-5. Boiled with chronic mixture, it yields large quantities of acetic and formic acids, and, after removal of those by distillation, ether extracts from the liquid a small quantity of trimellitic acid $C_6H_3(CO_2H)_3(E.)$.—6. The anhydride (colophony), oxidised with nitric acid, yields isophthalio aoid, together with trimellitic acid (Schreder, B. 6, 413).—7. Abietio acid fused with potash yields propionio, but no protocatechuic, acid (M.) .-- 8. Sodium-amalgam added to a warm alcoholis solution of abieticacid converts it into hydrabietic acid C,4H,8O, a dibasic acid which forms white unctuous lamine melting at 160° (M.).—9. Abietic acid with acetic chloride or anhydride at 160° yields an oily acetyl-compound (E.).-10. Bromine added to a solution of abietic acid in CS, forms a bromo-derivative, probably C4 H62Br.O5, which separates from alcohol as a red powder melting at 134° (E.).—11. Distilled with zinc dust it yields tolnene, m-ethyltoluene, naphthalene, methyl-naphthalene, and methylanthracone (Ciamician, G.4, 305, B. 11, 269).

Salts.—Abietio acid is dibasic, mostly forming normal, rarely acid, salts. The alkaline salts are difficultly crystallisable. The normal abie-tates of the other metals C₁, H₆₂M'O₂ are sparingly soluble in water, and are obtained byprecipitation. Na,A", needles (from alcohol).—MgA", flocculent, v. sol. alcohol.—MgH,A,".—CaA".—BaA".—ZnA", sl. sol. a Rohol.—CuA", v. sol. CS,

Ethyl Abistats Et.A", Otsined hy decom-

etherio odour; insoluble in water, slightly so-table in alcohol, easily in ether and CS.

Abietin CasH7,00, i.e. Ci.Ha1(CHMe: CH),0, is deposited from a mixture of glycerin and a concentrated alcoholic solution of abietic acid, after exposure to a low temperature for several days, in small white orystals melting at 125°, soluble in ether and aicohol (M.). H. W. ABIETIC ANHYDRIDE C., H., O, is not

formed by direct dehydration of the acid, but exists, as already observed, in the clear fresh turpentine of certain conifers, and forms the essential part of colophony.

ABIETIN. V. supra.
ABIETITE C₁H₂O₃.—Abietol. The sugar of

Abies pectinata.

ABROTINE Cal Hand. O. - An alkaloid from Artemisia abrotanum (P. Giacosa, J. 1883, 1356). White crystalline powder or white needles. sol. hot water. Its solutions fluoresee blue, Salts: B"H.PtCl_o.-B₂"H_SO₄Gaq. Needles, ABSINTHIN or Absynthiin C₄₀H₃₀O₂ [120°-

125°].—(Mein, A. 8, 61; Luck, A. 78, 87; Kromayer, Ar. Ph. [2] 108, 129).—The bitter principle of wormwood (Artemisia absynthium). Prepared by exhausting the dry herb with cold water; absorbing the bitter principle from the concentrated oxtract with boneblack; oxtracting with alcohol; purifying by treatment with basic lead acctate, precipitating the lead with H.S. and evaporating the filtrate.

Properties.—Yellow powder, composed of minute crystals. V. sl. sol. cold water, sl. sol. hot water, v. sol. alcohol or ether. Very bitter. Neutral to litmus. Smells like wormwood.

Reactions.-1. Conc. H.SO, forms a brown solution, turning greenish-blue. A little water turns the colour to a splendid blue, destroyed by more water .- 2. Boiling dilute H.SO, acquires a yellowish-green fluorescence, and deposits a brown resin. - 3. Does not reduce Fehling's solution.-4. Gives a mirror with warm ammoniacal AgNO₃. - 5. An alcoholic solution gives a sticky pp. with tannin. 6. Gives no pps. with metallic galts. H. W.

ABSINTHOL.—C₁₀H₁₆O (195°) or (204°).— **Beils**tein a. Kupffer, B. 6, 1183, A. 170, 290; Wright, C. J. 27, 1 and 319). Isomeric with commos camphor. Forms the essential principle of wormwood-oil, in which it is associated with a terpene (b.p. below 160°) and a deep-blue oil (270°-300) identical with the blue chamomile oil examined by Kachler (B. 4, 36). Absinthol boils at 195° (B. and K.), at 200°-205° (W.), 217° (Gir.lstone). Differs essentially from camphor in chemical reactions, not being converted into camphoric acid by oxidation with nitric acid, nor into campho carboxylic acid, $C_{11}H_{10}O_{2} = O_{10}H_{11}(OH)$. CO₂H, by sodium and CO₂, and yielding with melting potash a large quantity of resin but no acid. Heated with P.S. it yields cymene C.H., and cymyl hydrosulphide C. H., SH, boiling at 230°-240° (W.). Cymene is also formed, though in smaller quantity, by treating absinthol with zinc chloride (W.).

ABSORPTION OF GASES BY LIQUIDS AND

SOLIDS v. GASES.

ABSORPTION-SPECTRA v. PHYSICAL ME-WHODS: Sect. OPTICAL.

ACACIN or Acacia gum v. Arabin.

ACAJOU. - The pericarp of the nuts of the

Acajou or Cashew-nut tree, Anacardium occu-dentale, growing in the West Indies and South America, contains a large quantity of a red-brown resinous esicating substance, which may be extracted by ether, the solution when evaporated leaving a network of small crystal of anacardic acid soaked in an oily liquid called cardol, to which the resin owes its aerid properties (Stadeler, A. 63, 137). A catechin $C_{12}H_{31}O_{16}$ [165°] may be get from acajou-wood (Gautier, BL 30, 568).

ACAROID RESIN .- Resin of Xanthorrhea hastilis, a liliaceous tree of Australia: also called resin of Botany Bay. Yellow, fragrant, soluble in alcohol, ether and caustic potash. The potashsolution treated with HCl deposits benzoio and cinnamic acids. Nitrio acid readily oxidises if to pieric acid. Yields on distillation phenol and small quantities of benzene and styrone (Stenhouse, A. 57, 84). By potash-fusion it gives p-oxy-benzoio seid, rosorein, and pyrocatochin

(Hlasiwetz a. Barth, A. 139, 78). H. W. ACECHLORIDE OF PLATINUM v. ACETONE. ACECONITIC ACID C.H.O. .- The ethyl ether is formed, together with the (probably isomeric) citracctic ether, by the action of sodium on ethyl bromo-acetate:

 $3\text{EtC.H.BrO}_2 + 3\text{Na} = \text{Et}_3\text{C}_6\text{H}_3\text{O}_9 + 3\text{NaBr} + \text{H}_3$ (Baeyer, A. 135, 306). The product is distilled in vacuo, and the ethers saponified by baryta. Baric accomitate crystallises, leaving the gummy baric citracetate in solution.

Properties.-Nodular groups of needles, sol. other. Gives no crystalline sublimate.

Salts .- Barium salt forms small, sparingly soluble crystals. A solution of the calcium salt becomes turbid when heated.—Ag₃A"aq. Ethyl cther.—Et₃A". Lighter than water. H. W. ACEDIAMINE C2H6N2 i.e. NH2 CMe: NH v. Acet-amidine.

ACENAPHTHENE C12H10 i.e. C10H4: C2H4

M.w. 154. [95°] (Behr a. Dorp, A, 172, 265), [103°] (Schiff), (278° i. V.). V.D. 5-35 (for 5-38). S.V.S. 149-16 (Schiff, A. 223, 263).

Occurrence .- In coal-tar oil (Berthelot, Bl. [2] 8, 226).

Formation.-1. By passing a mixture of ethylene and benzene or naphthalene through a red-hot tube (Berthelot).—2. PBy passing (α)-ethyl-naphthalene through a red-hot tube.— 3. By treating (a)-ethyl-naphthalene with Br at 183° and decompesing the product, C₁₀H₁₀C₂H₄Br, with alcoholio KOH at 100° (Berthelot a. Bardy, C. R. 74, 1463).

Preparation .- Heavy coal-tar oil (260°-290°) is carefully fractioned, and the fraction 260°-270° cooled strongly till it solidifies. Recrystal lised from alcohol (Terrisse, A. 227, 184).

Properties.—Long needles (from alcohol). V. sol, hot alcohol, v. sl, son cold alcohol.

Redctions. - 1. A mixture of alcoholic solutions of acenaphtene and picric acid deposits orange-yellow needles of the picrate, C₁,H₁,C₂H₂(NO₂),OH [162°].—2. Corb. H₂SO₄ forms a sulphorate whose salts are very soluble. A little HNO turns the solution in H.SO, green. B. Cold fino, forms di-nitro-acenaphthene. Yellow needles (from banzoline); insol. in aloohol.—4. CrO₃ and H₂SO₄ give naphthalio aoid, O₁₆H₆(CO₂H)₂ (B.a. D.).—5. Bromine added to an ethereal solution forms bromo-acenaphthene, C,H,BrC,H, [53']; tables (from alcohol); oxilises to bromo-naphthalio acid (Blumenthal, B. 7, 1095) .- 6. A further quantity of bromine added to a solution in CS₂ forms C₁₂H₃Br₆; white needles (from alcohol).—7. Iodine at 100° jolymerises it.—8. Conc. III at 100° forms a iydrocarbon (? C₁₂H₁₂) (o. 270°).—9. Conc. III (20 pts.) at 280° produces naphthalene di hydride and ethane.—11. Potassium gives off hydrogen, forming C₁₂H_aK (Berthelot).

ACENAPHTHYLENE C₁₂H_a i.e. C₁₀H_a: C₂H_{.;}

probably

[93°] (265°-275°).

Preparation.—Accomphthene (6g.) is put into a combustion tube, and the rest of the tube filled with litharge. The acenaphthene is heated strongly, and the vaponrs pass over the litharge, which must not be red hot (Blumenthal, B. 7, 1092; Behr a. Dorp, B. 6, 753).

Properties.—Largo golden plates (from alcohol). Is partly decomposed by boiling. V. e. sol.

alcohol, ether or benzene.

Reactions.-1. Sodium amalgam roduces it, in alcoholio solution, to acenaphthene. -2. Chromic mixture oxidises it to naphthalic acid .-3. Combines, in ethereal solution, with bromine, forming

$$C_{10}H_{d} < CHBr$$

This forms white needles (from [1210_1230]. benzene mixed with alcohol). Chromic mixture oxidises it to naphthalic acid. Alcoholic KOH oonverts it into bromo-acenaphthylene,

$$C_{i0}H_{\bullet} \stackrel{CBr}{\underset{CH}{\longleftarrow}}$$

This is a liquid, but its picrate forms yellow needles. Resmo-accnaphtlyleno is converted by bromine into orange-rod platos of di-bromo-acenaphthylene,

Picrata.—C₁₂H.C.H.(NO₂),OH[202°]. Yellow needles. V. sl. sol. cold alcohol.

AGETACETIC ACID v. AGETO-AGETIC ACID.

ACETAL C.H.,O. t.e. CH.,CH(OEt),—Diethyl-acetal, di-ethyl aldehydate (v. Aldenyde), M.wr 118. (104°) (Staf); (108-2°) at 752 mm.

(R. Schiff, A. 220, 104); (21°) at 22 mm., (50·5 at 121 mm., (102·22°) at 760 mm. (Kabibaum S.G. 3° 8314 (Brāhl); 15 8319, 35 8233 (Perkin 1932 7364 (Sc.). V.D. 4:141. Critical temperatur 254 4° (Pawlewski, B. 16, 2633). S. 4 6 at 25 S.V. 159 88 (Sc.). μ_{θ} 1 386. R_{cr} 52 52 (B. M.M. 6 968 at 16 1° (F.).

Occurrence .- In crude spirit, after filterin through charcoal (Geuther, A. 126, 63).

Formation.-1. By the imperfect exidation

of alcohol (Doebereiner; Liebig, A. 5, 25; 1. 156; Stas, A. Ch. [3] 19, 146; Wurtz, A. Ci. [3] 48, 370; A. 108, 84). Hence its occurrence in raw spirit and in old wines.—2. By action collaboration of the collaboration o chlorine on alcohol:

 $3C_2H_6O + Cl_2 = C_0H_{14}O_2 + 2HCl + H_2O.$

3. One of the products of action of alcohol o ethyl di-bromo-acctate (Kessel, B. 11, 1917 4. By passing non-inflammable PH, into a mix ture of equal volumes of aldehyde and alcohe at -21° (R. Engel a. De Girard, C. R. 91, 692 C. J. 38, 458).

Preparation.-I. From Alcohol.-1. By im perfect oxidation under the influence of plati num-black. Fragments of pumice are moistened with nearly absolute alcohol in a wide-mouther flask, the upper part of which is filled with shallow glass capsules containing platinum black, and the flask, covered with a glass plate is left in a room at 20° till nearly all the alcoho is converted into acetic acid. Alcohol of 60 p.o is then poured in, and the flask, again covered with the glass plate, is exposed to the same temperature for a fortnight or three weeks, by which time the liquid above the pumice will have be-come viseid. This liquid is then poured off, more alcohol is added, and this course of proceeding is repeated till a few litres of very acid liquid have been obtained. This product is saturated with potassium carbonate, dried with calcium chloride, and about a fourth of it is distilled off; the distillate is treated with calcium chloride; the lower layer of liquid-consisting of aldehyde, ethyl acetate, and alcohol-is again mixed with calcium chloride, and distilled till the distillate no longer reduces silver nitrate; and the residne is treated with potash-lye, washed, dried with calcium chlorido and rectified (Stas). 2. By distilling alcohol (2 pts.) with manganese dioxide (3 pts.), sulphurio acid (3 pts.), and water (2 pts.), and rectifying the product, which consists of acetal mixed with aldehyde, ethyl acetate, &c., as abovo.-3. By passing chlorine through alcohol of 80 p.c. cooled to between 10° and 50 till . portion becomes turbid on addition of water, indicating the formation of substitution-products. One fourth of the acid liquid is then distilled off; the distillate is mentralised with chalk; fourth part again distilled off; and the distillate, consisting of alcohol, ethyl acetate, aldehyde, and acetal, is treated as above to separate the acetal (Stas). According to Lieben (A.Ch. [3] 52, 313), the ohief products of the action of ohlorino on 80 p.o. alcohol are mono- and dlchloracetal.

II. From Aldehyde.—1. By passing gaseous hydrogen chloride into a mixture of 1 vol. aldehydo and 2 vol. absolute aloohol, cooled by a freezing mixture, whereby the compound C.H.ClO is obtained, as an ethercal liquid floating on the

aqueous hydrochloric acid, and treating this

compound with sodium ethylate: $C_2H_1O + C_2H_2O + HCl = H_2O + C_4H_4ClO$; and $C_1H_2ClO + C_2H_3ONa = NaCl + C_6H_{14}O_2$ (Wnrtz-a. Frapolli, C. R. 47, 418; A. 108, 223) 2. By treating aldehyde with PBr., whereby it is converted into ethylidene bromide, and acting on this compound with sodium ethylate:

 $CHMeBr_1 + 2NaOEt = 2NaBr + CHMe(OEt)_2$

(W. a. P.).

Properties.-Colourless liquid, less mobile than ether, having a peculiar agreeable odour and refreshing taste, with an after-taste like that of hazel-nute. Separated from aqueous solution by ealcinm obloride and other soluble ealts. Miscible with ether or alcohol.

Reactions.—1. Not altered by mero exposure to air, but quickly oxidised in contact with platinum-black to aldeliyde and acetic acid. Oxidised also by nitric and by chromic acid, -2. Not decomposed by caustic alkalis if air is excluded. 8. Forms substitution-products with chlorinc. 4. Strong sulphuric and hydrochloric acids dissolve and decompose it, the mixture turning black .- 5. Dilute acids, even in the cold, split up acetal into alcohol and aldehyde.-6. A solution of acetal does not give the iode form reaction, nnlese it be first acidified (Grodzki, B. 16, 512). 7. PCl, forme CH, CHCLOEt, EtCl and POCL (Bnchanan, A. 218, 38).—8. Heated with glacial HOAe it forms acetie other, thus:

 CH_3 .CH(OEt)₂ + 2AcOH = $CH_2CHO + H_2O + 2AeOEt.$

9. Does not reduce AgNO₃Aq.—10. Chromic mixture forms acetic acid.—11. Heated with MeOH it is almost completely converted into EtOH and CH₂CH(OMe)...-12. Heated with ProH it is mostly unchanged, but some CH2.CH(OEt)(OPr) and some CH3CH(OPr)2 are formed .- 13. Heated with iso-amyl alcohol it bohaves as in 12.

References .- Homologues of acetal are described under the aldehydes, to which they correspond. Bromo- and chloro-acetals are described under brome and chlore acetic aldehyde. For oxy-acctal v. glycollic aldchyde.
ACETALDEHYDE v. ALDEYYDE.

ACETAMIDE C.H.NO i.e. NII₂Ac CH. COSNH -- Amide of acetic acid. M.w. 59. [83°] (Hofmann, B. 14, 2729) (222° cor.). S.G. $_{x}$ 1·159 (Schröder, B. 12, 562). R_{$_{\infty}$} 24·35 in a 7 1.159 (Schröder, B. 12, 562). R_{∞} 24.35 in a 34-p.c. aqueous solution (Kanonnikoff, J. pr. [2] 81, 347). Discovered by Dumas, Malaguti, and Leblano in 1847 (C.R. 25, 657).

Formution.—1. By heating ethyl acetate with

strong aqueous ammonia at 120°

AcOEt + NH2 = AcNH2 + HOEt.

2. By action of ammenia on acetic anbydride: $Ac_2O + 2NH_3 = NH_2Ao + AcONH_4$ 3. By distillation of ammonio acetate:

 $AcONH_4 = AcNH_2 + H_2O$ (Kundig, A. 105, 277). 4. When dry NaOAc (580 g.) is distilled with NH.Cl (225 g.) very little acetamide (70 g.) is got: the distillate is chiefly NH, and soid ammonic acetate, which boils at 145°

Preparation .- 1. Accio ether and aqueous ammonia are left in a closed vessel until the ether has disappeared. The product is distilled.—
2. Glacial acetic soid (1 kilo.) is saturated with dry NH, and the product distilled in a current of dry NH.. Above 190° acetamide (460 g.) comes

over; the first distillate (below 190°) is treated in the same way: it gives more acetamide (170 g.). A third repetition of this operation gives mors scetamid() (110 g.). Total yield: 740 g. (Keller, pr. [2] 31, 864).—3. Ammorio obloride and eodie acetate are hated in an enamelled iron digester for six hours at 230°. The product is distilled (Hofmann, B. 15, 981).—4. A mixture of ammonio acetate (20 g.) and acetic anhydride (26 g.) yields on distillation 96 p.e. (12 g.) of acctamide (Sohulze, J. pr. [2] 27, 512).—5. Ammonie sulphocyanide (1 mol.) is boiled for four days with glacial acetic acid (21 mols.): NH₄CNS + 2AcOH = 2AcNH₂ + COS + H₂O (S.).

Purification.-Acotamide can be freed from ammenic acetate by drying over lime (Menschut-

kin, J. R. 17, 259)

Properties.-Whito hexagonal seales, smelling like excrement of mice. Deliquescent. V. e. sol. water. Conducts electricity and is easily electrolysed.

Reactions .- 1. Resolved by distillation with P.O. into water and acetonitrile, C.H.N.-2. With P.S. it also yields acctonitrile, giving off H2S, and leaving a blackish tumefied residue.-3. Heated in dry HCl-gas it yiolds: a. A liquid distillate consisting of acetic acid with a small quantity of acetyl chloride; b. A crystalline distillate of (C₂H₅NO)₂HCl, and a compound of acetamide and diacetamide C₂H₅NO.C,H₇NO₂, the latter of which may be dissolved out by ether; c. A nonvolatilo residuo of acctamidine hydrochloride mixed with sal-ammoniac:

 $2C_2H_3NO + HCl = C_2H_6N_2.HCl + C_2H_4O_2$ (Strecker, A. 103, 328).-4. Acetamide heated in sealed tubes with saturated hydriodic acid yields ammonia, acetic acid, and ethane:

2C₂H₃NO + 3H₂ = C₂H₄O₂ + 2NH₃ + C₂H₄ (Berthelet, Bl. [2] 9, 183).—5. With CS₂ at about 210° it gives of H₂S, COS, CO, and probably ethane, leaving ammonium sulphocyanide mixed with undecomposed acetamide:

2C₂H₅NO + CS₂ = NH₂SCN + COS + CO + C₂H₆ (Ladenburg, Z. [2] 4, 651). V. Aldenburgs.

6. Nascent hydrogen (coppor-zine couple or sodium-amalgam) forms some alcohol and aldehyde (Essner, Bi. [2] 42, 98).-7. Heated with NaOEt at 180' it forms ethylamino (Seifert, B. 18, 1357).-8. With ethyl orthoformate at 180° acetamide yields ethyl alcohol and diacetylformamidine:

 $2NH_zAc + CH(OEt)_s = 3EtOII + N_z(CH)Ac_zH$. Another reaction, however, takes place at the same time, producing alcohol, ethyl acetate, and formamidine:

 $2NH_2Ac + CH(OEt)_s =$ EtOH + 2EtOAc + N2(CH)H.

(Wichelhaus, B. 3, 2).—9. Acetamide heated in sealed tubes with benzaldehyde is converted into benzylidene-diacetamide:

 $2NH_2Ae + PhCOH = H_2O + PhCH(NHAe)_2$ With aldehyde in like manner, it yields MeCH(NHAc), in large prisms [169°], partly decomposed by distillation, and giving off aldebyde when treated with soids (Tawild row, B. 5, 477). With anisaldehyde the compound O12H18N3O is formed in nodular groups of needles [1808], soluble in water, insoluble in alcohol and ethers decomposed by HOl, not altered by boiling with potash (Schuster, Z. [2] 6, 681). With salicylic aldehyde a yellow neutral body is formed (Credper. ib. 80). With chloral acetamide unites directly, forming the crystalline compound C.H.NO.C.HCl.O. (v. CHLORAL).—10. Heated with mesityl oxide it forms a basic substance, C.H., NO. oxy-hydro-collidine. A yellowish liquid (175°-180°) (Canzoneri a. Spica, G. 14, 349).

Combinations. - Acetainide unites directly with the stronger ands. The hydrochloride (NH,Ao),HCl is formed by passing gascous HCl into its solution in ether-alcohol. Long needles (from alcohol); insol. in ether.—NH, AcHCl (Pinner a. Klein, B. 10, 1896).—The nitrate, NH, AcHNO, [98°], separates from a solution of acetamide in strong HNO₃. It is very acid, and is deliquescent. Sl. sol. ether. Gives off CO2, N2O and HNO2 when heated.

Salts. -- AcNIIAg. Scales. -- (AcNII) Hg. Sixsided prisms [195°]. Both formed by dissolving the oxides in acetamide.—(AcNII) Zn. From ZnEt, and acctamido. Amorphous. (Frankland.)

Chloro-acetamides .- The amides of the chloroacetic acids are described under those acids. Aceto-chloro-amide NAcClH [110°] is formed by passing chlorino into fused acetamide, or by pouring aqueous HCl upon accto-bromo-amide:

2NAcBrH + HCl = NAcClH + NAcH2 + Br. (Holmann, B. 15, 410). Sol. ether. Split up by HCl into chlorinc and acctamide.

Bromo-acetamides v. Bromo-acetic acids.

Aceto-bromo-amide

NHBrAc [108°]. NHBrAc aq. [70°-80°]. Formed by adding aqueous KOII to a solution of Br (1 mol.) in acetamide (1 mol.). Striated rectangular plates (from ether).

Reactions .-- 1. Boiled with water it forms acetamide, Br, HBrO, methyl-acetyl-urca, and mcthylamine.-2. Heated with Ag.CO, it forms methyl cyanate:

2CH CO.NIIBr + Ag2CO2=

 $2CH_3NCO + 2AgBr + CO_2 + H_2O$. 3. Boiled with KOHAq it forms HBr, CO2, and methylamine, the methyl cyanate formed according to the last reaction being decomposed in the usual way.—4. Acetamido and NaOHAq form methyl-acetyl-urca.—5. Ammonia reacts violently, thus:

 $3NAcHBr + 5NH_3 = 3NAcH_2 + 3NH_4Br + N_2$. 6. Aniline forms scetanilide and tri-bromoaniline .- 7. Phenol gives tri-bromo-phenol and

acetamide (Hofmann, B. 15, 407).

Salts.-NAcBrNa. Hair-like needles, ppd. by conc. NaOH. NAcBrNaBraq. Made by adding conc. NaOH to a mixture of acetamide (1 mol.) and bromine (1 mol.). Rectangular plates. Decomposed by water into NaBr and aceto-dibromo-amide.

Aceto-di-bromo-amide NAcBr, [100°]. Made by adding aqueous KOH to a dilute solution of bromine (1 mol.) and bromo-acctamide (1 mol.) (Holmann B. 15,413). Golden needles or plates; sol. warm water, alcohol, or ether. Boiled with water, it gives HBrO, NAcBrH, and NAcH... Potash decomposes it into retrogen, acetic acid and potassic hypobromite. H.W.

Bromo-chlere-acetamide v. Силоко-пвомо-

ACETIO ACID.

Icdo-acetamide v. Iopo-ACETIO ACID. Di-acetamide NAo.H. M.w. 180 [82°] {210° 215°).

Preparation .- 1. The ethereal solution of the crystalline compound of accommide and di- [1850]. Obtained as above. Nodules of small

acetamide got by heating acetamide in a current of HCl (v. Reaction 3), deposits, whon gaseons HCl is passed through it, spicular crystals of acetamide hydrochloride, and the filtrate yields, by evaporation over H2SO, orystals of diacetamide.—2. By heating acctonitrile with glacial HOAc, or acetamide with Ac_2O at 250° (Gautier, Z. 1869, 127).—3. By boiling methyl-acetyl-ursa with Ac₂O (Hofmann, B. 14, 2731)

Properties. — Long needles (from ether). Neutral. V. e. sol. water, v. sol. alcohol or ether. Does not combine with acids, so that HCl gives

no pp. in an othereal solution.

Reactions.—1. By boiling with acids or by heating with ZnCl₂ it is resolved into acetic acid and acetonitril .- 2. Fuming HNO, reacts, giving off N2O.

Tri-acetamide NAc, [79°].—Formed in small quantity when a mixture of acetie anhydride and acetonitrile is heated to 200°, and may be dissolved out by ether after the excess of Ac2O has been distilled off. White fiexible needles [780-79°]. Neutral. Gently warmed with silver oxide it yields silver acetate; so likewise do aceta-mide and diacetamide (Wichelhaus, B. 3, 847). H. W.

Tri-acet-di-amide N₂Ac₂H₂ [212°-217°]. — This is the compound of acetamide and di-acetamide mentioned under acetamide (Reaction 3) and di-acctamide (Preparation 1).

Di-azo-acetamide v. Azo compounds. Ethyl-acetamide v. ETHYL-AMINE. Methyl-acetamide v. METHYL-AMINE.

Phenyl-acetamide v. Aniline.

ACET .- If compounds whose names begin with acet or aceto are not here described, remove this prefix and look for the remaining word, changing the termination ide, if present, into ine.

AČEŤAMIDINE C₂II₆N₂ i.e. CH₃.C(NH).NH₄ Acediamine, Ethenyl-amidine, Acet-imid-amide (Strecker, A. 103, 328; Hofmann, B.17, 1924).— The hydrochloride of this base is left as a residue when acetamide is distilled in a current of HCl (v. Acetamide, Reaction 3). The mass is extracted with alcohol, which leaves NH,Cl behind.

Properties.—When liberated from solutions of its sales, it splits up into ammonia and am-

monic acctate.

Salts. - B'HCl: prisms (from alcohel), [165°], -(B'HCl)₂PtCl₄: yellowish-red prisms.

-B'2II,SO.: pearly lamine.

Reactions.-1. The hydrochloride boiled with Ac O aud NaOAc for 11 hours forms anhydrodi-acetyl-acct amidine and anhydro-di-acetyl-acet-amidil (Pinner, B. 17, 173).—2.27. Aceto-ACETIC ETHER, Reaction 25.

Anhydro - di - acetyl - acetamidine C.H.N.O [253].—Prepared as just stated, the product being treated with agreeous NaOH and the pp. boiled with water, which dissolves the 'amidil,' but not the amidino.

Silky needles (from alcohol). Insol. water, sl. sol. cold alcohol, v. sol. hot alcohol, v. e. sol. dilute acids. Forms a platino-chloride.

Anhydro-di-acetyl-acet-amidil CaH, NaO2aq. CH, CON CH, Possibly NH.CO.CH

prisms. Loses 2ag over H.SO. Sl. sol. cold jar, Much of the alcohol is, however, converted water, v. sol. hot water, v. e. sol. alcohol and in

dilute scids. Forms a platinum salt.
ACETAMIDOXIM v. ETHENYL-AMIDOXIM. ACETANILIDE v. Aniline Acetyl deriva-

ACETIC ACID C.H.O. i.e. CH.CO.OH or Active Active C₂H₁O₂ A.E. CH₃.CO.OH of AcOH.—Methane carboxylic acid, Pyroligneous acid.—M.w. 60 [16·5°] (Zander), [17·5°] (Sonstadt, C. N. 87, 199). (118·29°) (Z.), (117·5°) (Schiff.). Critical temperature 321·5° (Pawlewsky, B. 16, 2684). S.G. Solid. § 1·0701 (Z.); ½ 1·0607 (Mendeléefi, J. 1806, 7). S.G. Liquid. 1.5 1·0576, 13 1·0542, 18 1·0562 (Patternative Active Act 10543, 19 1.0503 (Pettersson, J. pr. [2] 24, 801); 29 1.0495 (Brühl); at boiling-point 9325 (Ramsay, C. J. 35, 463). V.D. 29.7 at 250° and upwards. C.E. (0°-10°) 00106 (Z.). H.F.p. 105,290. H.F.v. 104,130. S.H. (between 0° and 100°) 497. Latent heat of fusion for 1 mol. (at 1.5° to 4.2°) 2619. μ_{θ} 1.3765. R_{∞} 20.69 (B.). M.M. 2.525 (Perkin). S.V. 64.3 (R.).

Occurrence. In the juices of plants, especially of trees, and in certain animal eccretions.

Synthesis.—1. From acetylene (i.e. from C and H) by converting that hydrocarbon into ethylene by direct addition of hydrogen, then the ethylene into alcohol, and oxidising tho alcohol; or more simply by heating acetylene dichloride with aqueous potash at 230° or with alcoholic potash at 100° for ten hours:

 $C_2H_2Cl_2 + 3KHO = C_2H_2O_2K + 2KCl + H_2O$ (Berthelot, Z. [2] 5, 683).—2. When a mixture, I vol. acetylene and 2 vols. air, is exposed to daylight over dilute potash-lye, the acetylene is slowly oxidised to acetic acid, which is absorbed by the alkali: $C_2H_2 + O + KOH = C_2H_3O_2K$ (Berthelot, A. Ch. [4] 23, 212).-3. From sodiummethyl and oarbonic acid:

CH2Na + CO2 = CH2.CO2Na

(Wanklyn, A. 111, 234).—4. By boiling acctonitrile (methyl cyanide) with potash:

 $CH_2CN + KOH + H_2O = CH_2CO_2K + NH_3$ Frankland a. Kolbe, A. 65, 298).- 5. By passing CO over sodium methylate at 160°:

 $CH_2ONa + CO = CH_1CO_2Na$

(Fröhlich, A. 202, 294).

Formation .- 1. By dry distillation of organic bodies, especially wood .- 2. By the action of atmospheric oxygen, chromio acid, nitric acid, hypochlorous acid, and other oxidisere, on alcohols and other organic bodies, especially under the infinence of ferments which act as carriers of oxygen. e-3. By the action of KOH or NaOH at a high temperature on various organic bodies, e.g. tartarie, citric, and malic acids, sugar, alcohol, &c.—4. In various processes of fermentation

and putrelaction (J. 1878, 1917, 1019, 1023).

Preparation.—1. By oxidation of ethyl alcohol, the alcohol being first converted into aldehyde: C₂H₆O + O = H₂O + C₂H₄O, and the aldehyde then oxidised to acetic acid. The oxi-

dation may be effected:

a. By the influence of spongy platinum. If a tray containing this substance be placed over a dish containing a little alcohol, the whole being covered with a bell-glass open below, as well as at the top, on gently warming the dish the alcohol will be rapidly oxidised, acetic soid sondensing in abundance on the inside of the

into aldehyde and lost by volctilisation.
b. Under the influence of ferments. This is

the ordinary process of making vinegar fron alcoholic liquids, wine being generally, used to the purpose in Franchand GermanyPand maltir England. The most favourable temperature is The experiments of Pasteur have 25°-80°. shown that the oxidation of alcohol in the ordinary process of vinegar-making depends essen tially on the presence of a fungoid plant callso Mycoderma vini, Mycoderma aceti, or 'mother of-vinegar,' and is invariably precoded by its development on the surface of the liquid. It appears to act like platinum-black, as a carrier of oxygen. The plant may be sown on the surface of the liquid by introducing a small portion of it from another vinous liquid already in the fermenting state, or by simply exposing the liquid to the air in which the germs of this fungus, as of many others, are always floating. Like all other plants, it requires food for its development, and this it finds in the albuminous matter and mineral salts contained in ordinary vinous liquore. If these are absent the plant cannot grow, and acetification cannot take place. Thus, pure aqueoue alcohol may be exposed to the air for any length of time without turning acid, because the germs of the mycoderma which fall into it from the air remain barren for want of nutriment. Moreover, pure aqueous alcohol may be acetified without the aid of any albuminous matter, provided the myeoderma have access to it, and he supplied with the nitrogen and ealine matters necessary for its growth. Pasteur has in fact shown that this nutriment may be supplied in the form of alkaline and earthy phosphates and ammonium phosphate, the latter furnishing the nitrogen. Under these circumstances the inycoderm grows, though less quickly than in ordinary vinoue liquids, and the alcohol is clowly converted into acctic acid. If the mycoderma be allowed to remain in the liquid after the acctification ic complete, the whole of the acetic acid may be destroyed and the liquid rendered perfectly neutral. (Pasteur, Etudes sur le Vinaigre, Paris, 1868; also Annales Scienti-fiques de l'École normale supérieure, tome i. 1864; Bl. 1861, p. 94; J. 1861, 726.)

Malt Vinegar ie prepared from a fermented wort obtained by mashing malt, or a mixture of malt and raw barley, with water, as

in brewing.

Quick Vinegar Process.—The oxidation of the alcoholic liquor may be greatly accelerated by allowing it to trickle down in a fine shower over chips of wood covered by the mycoderma, and exposed to an npward current of air.

Wood Vinegar -- Pyroligneous Acid .- The greater part of the acetic acid now used in arts and manufactures is obtained by the destructive distillation of wood. The wood is heated in large iron cylinders connected with a series of condensers. The watery liquid which condenses in the receivers, consisting of water, tar, methyl alcohol or wood-spirit, methyl acetate and acetic acid, is redistilled after separation of the tar, the wood-spirit passing over among the first portions of the distillate and the acctic or pyroligneous acid afterwards. The acid thus obta tained is coloured and has a strong tarry flavour,

act removable by distillation. To purify it, the crude liquor is saturated with lime, which removes part of the tarry matter, the rest remaining in solution with the calcium acetate. The liquid clarified by repose or by filtration, is evaporated inean iron pot to half its bulk, and mixed with enough hydrochloric acid to give a slight acid reaction, whereupon the greater part of the tarry matter separates, and may be skimmed off the surface. The hydrochloric acid also decomposes certain compounds of lime with creosote and other volatile substances, which msy then be expelled by heat. The calcium acetate thus purified is completely dried and distilled with hydrochlorio acid. The density of the acetic acid thus obtained is about 1.06. If it contains hydrochloric acid it may be purified by redistillation with addition of a small quantity of sodium carbonate, or, better, 2 or 3-p.c. potassium dichromate, this latter at the same time destroying certain erganic impurities which give the acid a peculiar odour (Völckel, A. 82, 49). Crude wood vinegar contains small quantities of propionic, n-butyrio, n-valerio, and two crotonic acids (Grodzki a. Krämer, B. 11, 1356).

Crystallisable or Glacial Acetic Acid -the pure acid, C2H4O2, so-called because it crystalliscs at ordinary temperatures-is obtained: 1. From the ordinary aqueous acid by fractional distillation, repeated till the residue solidifies on cooling. 2. By distilling certain dry metallic acetates with strong sulphurio acid or with hydrogen potassium sulphate,

2C,H,KO₂ + H,SO₄ = K,SO₄ + 2C,H,O₂; and C₂H,KO₂ + HKSO₄ = K,SO₄ + C,H,O₂. 3. Together with acetone and other products, by dry distillation of cupric acctate (Spiritus

Eruginis or Sp. Veneris).

Physical Properties.—The selid acid ferms prismatic or tabular crystals. The diquid acid is transparent, colourless, and mobile. Vapourdensity at 250° and upwards is 2.08 (air =1) er 29.7 (H = 1), which is nearly half the molecular weight of the acid, showing that at these high temperatures the vapoer exhibits the normal condensation. But at temperatures nearer to the boiling-point the density of the vapour is much greater, showing a cendensation to ? vol. or even less (Cahours, C. R. 19, 771; 20, 51). The pressure of the vapour of solid acetic acid is 1.3 mm. at -5.7°, 2.0 mm. at 0°, and 9.5 mm. at 16.4°; the vapour pressure of liquid acetic acid being 3.2 mm. at 0°, 6.3 mm. at 10°, 11.8 mm. at 20°, 19.9 mm. at 30° (Ramsay a. Young, C. J. 47, 45).

Glacial acetic acid has a pungent sour taste and odour and blisters the skin. It does not redden litmus paper per se, but reddens it strongly when mixed with water. It does not attack CaCo, until water is added. It is hygrescopic.

Aqueous Acid .- Acetic acid mixes with water in all proportions. The density of the aqueous acid does not vary in proportion to the amount of real acid present; and consequently the strength of any sample cannot be inferred from its density, but must be determined by itration with standard alkali. The following table has been constructed in this manner by Oddemans (Fr. 5, 452) for the temperatures 0° 15°, and 40°.

Density of Aqueous Acetic Acid (Ondemana)

Density of Aqueous Acetic Acid (Oudemans).				
O.H.O.		Density		
p. c.	at 0°	at 150	at 409	
0	0 9999	0.9992	0.9924	
1 2	1.0016 1.0033	1.0007 1.0022	0.9936	
3	1.0051	1.0022	0.9948 0.9960	
4	1.0069	1.0052	0.9972	
5	1.0088 1.0106	1.0067	0.8984	
6 7	1.0100	1·0083 1·0098	0.9996 1.0008	
8	1.0142	1.0113	1.0020	
9	1.0159	1.0127	1.0032	
10 11	1.0176 1.0194	1·0142 1·0157	1·0044 1·0056	
12	1.0211	1.0171	1.0067	
13	1.0228	1.0185	1.0079	
14 15	1.0245 1.0262	1.0200	1.0090	
16	1.0202	1·0214 1·0228	1·0101 1·0112	
17	1.0295	1.0242	1.0123	
18	1.0311	1.0256	1.0134	
19 20	1·0327 1·0343	1·0270 1·0284	1.0144	
21	1.0359	1.0298	1·0155 1·0166	
22	1.0374	1.0311	1.0176	
23 24	1.0390	1.0324	1.0187	
25	1·0405 1·0420	1·0337 1·0350	1.0197 1.0207	
26	1.0435	1.0363	1.0217	
27	1.0450	1.0375	1.0227	
28 29	1.0465 1.0479	1·0388 1·0400	1·0236 1·0246	
30	1.0493	1.0412	1.0255	
31	1.0507	1.0424	1 0264	
. 32 33	1·0520 1·0534	1·0436 1·0447	1.0274	
34	1.0547	1.0459	1·0283 1·0291	
35	1.0560	1.0470	1.0300	
36 37	1.0573	1.0481	1.0308	
38	1·0585 1·0598	1·0492 1·0502	1·0316 1·0324	
39	1.0610	1.0513	1.0332	
40	● 1·0622	1.0523	1.0340	
41 42	1.0634 1.0646	1·0533 1·0543	1·0348 1•0355	
43	1.0657	1.0552	1.0363	
44	1.0668	1.0562	1.0370	
45 46	1.0679 1.0690	1·0571 1·0580	1·037 ? 1·038 4	
47	1 0700	1.0589	1.0391	
48	1.0710	1.0598	▶0397	
19	1.0720	1.0607	1.0404	
50 51	1·0730 1·0740	1.0615 1.0623	1·0410 1·0416	
52	1.0749	F0631	1.0423	
53	1.0758	1.0638	1.0449	
54 55	1·0767 1·0775	1.0646 1.0653	1·0434 1·0440	
56	1.0783	1.0660	1.0445	
57	1.0791	1.0666	• 1°0450	
58	1.0798	1.0678	1.0455	
69 60	1·0806 1·0813	1.0679 1.0685	1·0460 1·0464	
61	1.0820	1.0691	1.0468	
	1.0826	1.0697	1.0472	
63	1.0832	1.0702	• 1.0475	

O.H.O.		Density	•
p. c.	at 0°	at 15°	at 40°
64	1.0838	1.0707	1.0479
85	1.0845	1.0712	1.0482
86	1.0851	10717	1.0485
67	1.0856	1.0721	1.0488
68	1.0861	1.0725	1.0491
69	1.0866	1.0729	1.0493
70	1.0871	1.0733	1.0495
71	1.0875	1.0737	1.0497
72	1.0879	1.0740	1.0498
73	1.0883	1.0742	1.0499
74	1.0886	1.0744	1.0500
75	1.0888	1.0746	1.0501
76	1.0891	1.0747	1.0501
77	1.0893	1.0748	1.0501
78	1.0894	1.0748	1.0500
79	1.0896	1.0748	1.0499
80	1.0897	1.0748	1.0497
81	1.0897	1.0747	1.0495
82	1.0897	1.0746	1.0492
83	1.0896	1.0744	1.0489
84	1.0894	1.0742	1.0485
85	1.0892	1.0739	1.0481
86	1.0889	1.0736	1.0475
87	1.0885	1.0731	1.0469
88	1.0881	1.0726	1.0462
89	1.0876	1.0720	1.0155
90	1.0871	1.0713	1.0447
91		1.0705	1.0438
92		1.0696	1.0428
93	_	1.0686	1.0416
94		1.0674	1.0403
95		1.0660	1.0388
96		1.0644	1.0370
97		1.0625	1.0350
98		1.0504	1.0327
99		1.0580 1.0553	1.0301
100		1.0008	1.0273

The maximum density corresponds at 0° to about 81 p.c., and at 40° to about 76 p.c. Orthoacetic acid, CH₂C(OH)₃, would contain 77 p.c. of HOAc.

Reactions .- 1. Vapour inflummable, burning with blue flame to water and CO2-2. Partly decomposed by passing through a red-hot tube yielding carbon and combustible gases, together with acetone, benzene, phenol, and naphthalene (Berthelot, A. Ch. [3] 33, 295).—3. Dropped upon hot ZnCl2 it gives CO, CO2, C2H4, C3H8, isobutylene, and a little CH₄ (Lebel a. Greene, Am. 2, 26).—4. Passed over zine dust at 300°-350° it gives hydrogen, acetone, CO, and somo propylene (Jahn, M. 1, 683) .- 5. Mixes with strong sulphuric acid without evolution of gas, but the mixture becomes hot, and if further heated gives off CO2 mixed with SO2. Dissolves SO3 without giving off gas, forming sulpho-acetic acid. Not sensibly altered by nitric acid.-6. Periodie acid converts it into carbonic or formio acid, with formation of iodic acid and separation of iodine.—7. With chlorine in sunshino it forms mono- and tri-chloro-acetic acids (q. v.), tho one or the other predominating according as the acetic acid or the chlorine is in excess. -8. Heated with bromine in a scaled tubo it forms mono- and di-bromo-acetic acids. Not acted upon

by iodins, even in sunshine.—9. With PCl₂ it forms AcOl, HCl, and POCl₂ With PCl₂ it reacts thus: 3AcOH+2PCl₂ = 3AcOl+P₂O₃ + 3HCl.—10. With P₂S₃ the products are thio-acetics oid and phosphoric oxide: 5AcOH₂ P₂S₂=P₂O₅ + 5AcSH.
11. With chromyl-dichloride OrQ.Cl₂ it forms the compound Cr₂O₃(C₂H₂O₂)₁₀3H₂O (Etard, A. Ch₂ fig. 22, 286).

Ch. [5] 22, 286).

Detection.—The solution supposed to contain acctio acid or an acctate is acidified with H₂SO₄ and distilled. The distillate, if acid, is neutralised with KOH and should then give the following tests: (1) FcCl₃ a brown-red colour, and a pp. on boiling. (2) AgNO₃ a white flocculent pp., sol. hot water, separating in spangles whon the solution cools. (3) Evaporate to dryness, mix with As₂O₃ and heat: a disgusting odour of caeedyl is perceived.

Acetic Acid Dibromide C.H.40, Br. [37°] is formed on treating acetic acid with bromino in presence of a small quantity of carbon bisulphide. Orange-red needles or thick roseate prisms, very deliquescent; dissolving in water with great fall of temperature and separation of bromine, in alcohol, benzene, and glacial acetic acid with partial formation of substitution-products. At 100° dissociation first takes place, but finally HBr and C.4H,BrO, are formed.

Hill and C₁H₂BrO₂ are formed.

Compounds of acetic acid with Br and HBr.
On adding bromine to well-cooled glacial acetio acid saturated with HBr, the whole solidifies to a mass of thick, rather large, tabular, crystals, which when dried have the composition (C₂H₄O₂)₂Br₂HBr; they fume in the air, molt and decompose at +8°, and are decomposed by water and by potash-lyc, yielding (C₂H₄O₂)₂Br₂ and KBrO₃. Heated in a sealed tube, they yield bromacetic acid (Steiner, B. 1874, 184). The compound (C₂H₄O₂)₂Br₃HBr has also been prepered by Hell a. Mühlnäuser (B. 1878, 241), who by using larger quantities of bromine have further obtained (C₂H₄O₂)₂Br₄(BrH)₂ in radiate groups of hard roscate crystals, which may be dried in the lime exsicenter.

Acctates.—Acetic acid is monobasio, the general formula of its normal salts being:

 $R^{(n)}(CH_3,COO)_n = R^{(n)}A'_n$ the symbol $R^{(n)}$ denoting an n-valent radiclo metallic or alkylic, and A' standing for $C_2H_2O_2$.

METALLIC ACETATES.—The normal acctates all dissolve in water, and most of them readily. The least soluble are the silver and mercury salts, so that solutions of other acctates added to mercurous nitrate or silver nitrate throw down white shining scales of mercurous or silver acctate. But for the most part acctates are formed not by precipitation, but by the action of acctic acid on metallic exides or carbonates; many carbonates, however, those of barium and calcium for example, are not decomposed by acctic acid in its most concentrated state. All acctates are decomposed by heat, most of them yielding carbon dioxide, acetone and an empyroumatic oil. Those which are easily decomposed, and likewise contain bases forming stable carbonates, are almost whelly resolved into acctone and carbonate, e.g.:

G Ba(O.CO.Me)₂ = COMo₂ + BaCO₃. Those which, like the potassium and sodium salts, require a higher temperature to decompuse them, yield more complex products, but always a

certain quantity of acetone. Among the products are found methylethyl ketone and methyl propyl ketone, together with dumasin C.H.O (Fittig, A. 110, 17). Acetates containing weaker bases give off part of the acetic acid undecomposed, the remaining portion being resolved into acetone and carbonic anhydride, or if the heat bestrong, yielding empyreumatio oil and charcoal: the residuo consiste sometimes of oxide, sometimes, as in tha case of copper and silver, of reduced metal; in this cass part of the acetic acid is burnt by the oxygen abstracted from the metal. The decomposition of silver acetate may be oxprossed by the equation

 $4CH_3.CO_2Ag = 3CH_3.CO_2H + CO_2 + C + 4Ag$ (Iwig a. Hecht, B. 19, 238). Acetates hoated with a large excess of fixed caustic alkali, are resolved at a temperature below redness into marsh gas and alkalino earbonate, e.g.:

 $KC_2H_3O_2 + KOH = K_2CO_3 + CH_4$ Acetatos distilled with sulphuric acid and alco-hol yield ethyl acetate. The acetates of the alkali-metals, and probably others also, treated with phosphorus oxychloride, yield acetyl chlo-ride, together with a tribasio phosphate:

 $3NaOAo + POCl_3 = 3AcCl + Na_3PO_4$ Many acetates may be decomposed by water into acetic acid and metallic oxide. This decomposition in the case of aluminio and ferrio acctates occurs at 100°, while at 175° the acctates of Mn, Co, Ni, Zn, Ur, Cu, and Ag, as well as ferrous and mercurio acetates, are slowly de-

composed (Riban, C. R. 93, 1140).
Aluminium Acetates.—The normal salt AlA', exists only in solution, being decomposed on evaporation. The solution, which is much used as a mordant in dyeing and oalico-printing, and is called 'red liquor' because it yields madder reds and pinks may be formed by dissolving freshly precipitated aluminium hydroxide in strong acetic acid, or by precipitating a solution

of normal aluminium sulphate with lead acetate: $Al_2(SO_4)_2 + 3PbA'_2 = 3PbSO_4 + Al_2A'_8$. When quickly evaporated at a low temperature, by spreading it out in thin layers ou glass or porcelain, it leaves a soluble basic acetate:

Al2O2.2C4H2O2.4H2O or Al2O(AcO)4.4H2O. forming a gummy mass perfectly soluble in water; but, if hoated, or left to evaporate at erd. temp., it deposits insoluble basic salts, containing in the first case two, and in the second five molecules of water, instead of four. The soluble acetate exposed in diluto solution to the temperature of boiling water for several days, undergoes a remarkable change, the whole or nearly the whole of the soid being expelled and a peculiar modification of alumina remaining dissolved (v. ALUMININM) (Walter Crum, C. J. 6, 216). dilute solution of aluminio acetate, free from alkali, maybe boiled without a pp. being formed (Reinitzer, M. 3, 259).

Ammonium Acetates .- Thenormal salt NH, A [89°], obtained by saturating eglacial acotic acid with dry ammonia-gas, is white, odourless, extremely soluble, and difficult to crystallise, its aqueous solution when evaporated giving off ammonia and leaving the acid salt (Barthelot, Bl. 22, 440; Smit, Bl. 24, 539; Bahrmann, J. pr.

agneous solution known in the Pharmacopoeia as Spiritus Minderer is prepared by saturating aqueons acetic acid with ammonia or ammonium carbonate. The acid salt NH, HA' (145°) is obtained as a crystalline sublimate with evolution of ammonia by heating powdered ammonium chlo-ride with potassium or calcium acetate (v. Acar-ANIDE). When commercial ammonium acetate is dissolved in its own weight of glacial acetio acid, an acid salt is obtained in long needles, having the composition 2NH, A'3HA' (Berthelot, Bl. 24, 107).

Barium Acstate BaA', aq, prepared by decomposing the earbonate or sulphide with acetic acid, is obtained, on ovaporating the solution at a gentle heat, in flattened prisms; and on cooling to 0°, in monoclinic prisms, BaA', 3aq. The crystals dried at 100° yield the anhydrous salt as a white powder, resolved at a high temperaturs into barium carbonate and acetone. S.G. (of BaA'2aq) 2.02; (of BaA'2) 2.47 (Schröder). V. e. sol. water, insol. alcohol.

Acid Salts.—BaA', HA'2aq.—BaA', 2HA'2aq. (Villiers, Bl. 30, 177; C. R. 85, 1234).

Double Salt .- BaA'(NO3)4aq (Lucius, A. 103,

Bismnth Acetate separates in micaceous laminæ from a warm mixture of bismuth nitrate and potassium acetate. Acetic acid mixed with a solutiou of bismuth nitrate prevente the precipitation of that salt by water.

Cadmium Acetats CdA'23aq. - Monoclinic prisms. V. e. sol. water, deliquescent and difficult to crystallise (Hauer, Sitz. B. 16, 131). S.G. 2.01 (dry, 2.31, Schröder).

Calcium Acetate CaA' eq. Small efflorescent needles. V. sol. water, sl. sol. alcohol. S.G. of aqueous solutions of CaA'2 at 17.5° (Franz, J. pr. .[2] 5, 298) :

P.C.	S.G.	P.C.	S.G.	P.C.	8.G.
1	1·0066	11	1.0527	21	1·0925
3	1·0198	13	1.0597	23	1·1027
5	1·0330	15	1.0666	25	1·1130
7	1·0394	17	1.0750	27	1·1248
9	1·0458	• 19	1.0834	29	1·1366

Calcie acetate splits up on distillation into CaCO, and acetone.

Acid salt CaA'2HA'12aq. Hygroscopic. Double salt CaA'2CaCl210aq. Monoclinic prisms, permanent in air.

Carena Acetate Ce, A Baq forms radiate groups of small needles, which become anhydrous in dry air without losing their crystalline form; after drying at 115° they carbonise at a higher temperature without fuelig, and when strongly heated leave a residue of cerous oxide Carge, J. pr. 82, 129).

Chromium Acetates .- The chromous salt CrA'aq, obtained from the chloride by decomposition with potassium or sodium atetate, forms red transparent crystals which when moist absorb oxygen very rapidly from the air, sometimes taking fire (Peligot, A. Ch. [3] 12, 541).—Normal Chromic Acetate Cr.A', 2aq is obtained by evaporating a solution of chromic hydroxide in [2] 27, 296). When dietilled with phosphorio evaporating a solution of officinic hydroxide in alchydride it losss 2 mol. water, and gives off acctic acid, as a green crystallino mass, insoluble acctonitrile C.H.N. NH.C.H.O. 2H.O. The in alcohol. Its aqueous solution, green by acceptance of the control of

flected, red by transmitted light, is not decemposed either by boiling or by addition of limewater; but ammonia throws down from it a green precipitate of chromic hydroxide, soluble in excess./H. Schiff, A. Ch. [3] 71, 140; Schützenberger, Bl. [2] 4, 86). The solution of the normal acetate heated for several days with excess of chromic hydroxide loses its acid reaction, and yields by evaporation a green powder soluble in water, consisting of a basic acetate Cr.A',(OH),

Schiff, A. 124, 168).

Chromic Diacetotetrachloride, Cr. A', Cl, is obtained by dissolving Cr.OCl, in strong acetic acid. It is an unstable salt, which gives off acetic acid when heated above 100°. The chlorine is but very slowly precipitated from it by silver nitrate at ordinary temperatures, but, on the other hand, the salt easily yields acetic ether when heated with sulphuric acid and alcohol (Schiff). - Chromic Diaceto-sulphate Cr₂A'₂(SO₂)₂, obtained by dissolving chromic disulphate in acetic acid, is a crystalline salt which becomes anhydrous at 100, and gives off acetic acid at a higher temperature (Schiff).-Chromic Pentacelo-nitrate Cr.A', NO, 4aq is obtained by mixing a solution of chromio hydroxide in a slight excess of acetic acid with a solution of the same quantity of chromic hydroxide in the exact quantity of nitric acid required to The concentrated solution, when dissolve it. left to itself, deposits an abundant crystallisation of a dark green salt, which may be purified by recrystallisation from water or from glacial acetio acid. It forms dark green bulky lamine, which give off nitrous fumes at 100°, the chromium being at the same time converted into trioxide. (Schützenberger).

Cobalt Acctate CoA', 4aq. - Red needles.

Copper Acetates .- The euprous salt Cu.A' sublimes towards the end of the distillation of normal cupric acetate. According to Berzelius. it is contained in green verdigris and sublimes en distillation. Soft loose white flakes which redden litmus and have a caustic astringent taste. Decomposed by water, yielding normal

enprie acetate and cuprous oxide.

The normal cupric salt CuA' aq is prepared by dissolving cupric oxide or common verdigris in hot acetic acid, or by decomposing normal lead acetate with cuprio sulphate. Dark-green monoclinic crystals (S.G. 19), efflorescent, soluble in 14 pts. cold and 5 pts. boiling water, sparingly also in alcohol, insoluble in ether. The solution boiled with grape-sugar yields a rod precipitate of cuprous oxide. Cupric acetate crystallised at s temperature noar 8° yields crystals containing CuA',5H,O. A solution of cupric acetato heated in a sealed tubeat 200° forms crystallised enpress oxide and cupric glycollate (Caseneuve, C. R. 89, 525).

Acid Cnpric Acetate CuA'2HA'aq (Vil-

liers, C. R. 85, 1234).

Basic Cupric Acetates .- These salts may be regarded as compounds of the normal salt with CuO, as compounds of Ac₂O with CuO, or, by taking account of water of crystallisation, as aceto-hydrates, e.g. :

Sesquitasio • On O.2 CuA', 6aq = 3CuO.2Ac, O.6Aq = 2CuA' (OH). CuA', 5aq. Dibasio: CuO.CuA's6Aq=2CuO.AcsO.6aq= 2(HO.CuA)5aq.

Tribasio: 20u0.CuA',2ag = 80u0.Ac,0.2aq

=2(HO.CuA')Cn(OH);.
They are contained in common verdigris plates of copper to the air in contact with acetic acid, and much used as a pigment and as a mordant in dyeing wool black. There are two varieties of this substance, the blue and the green, the former consisting almost wholly of dibasic cupric acetate, the latter of the sesquibasic salt mixed with smaller quantities of the dibasic and tribasic acetatss. The dibasic salt or blue verdigris is prepared at Montpellier and in other parts of the south of Francs, by exposing copper to the air in contact with fermenting wine-lees. The same compound is obtained by exposing copper plates to damp air in contact with normal cupric acetate made into a paste with water. It forms delicate, silky, blue crystalline needles and scales, which yield a beautiful blue powder. They contain 6 mol. water, which they give off at 60°, and are theu converted into a green mixture of the monobasic and tribasic salt.

Green Verdigris is manufactured at Grenoble by frequently sprinkling copper plates with vinegar in a warm room; and in Sweden by disposing copper plates in alternate layers with flan-

nols soaked in vinegar.

Calcio-cupric Acetate CaA', CuA', 8aq, obtained by heating a mixture of 1 mol. CuA', and 1 mol. Ca(OH), with 8 pts. water and sufficient acetic acid to dissolve the precipitated CuO, and evaporating the filtrate at 25°-27°, crystallises in large blue square prisms, slightly efflorescent, giving off acetic acid and falling to powder at 75° readily soluble in water. Another calcio-cuprio acctate often exists in crystallised verdigris.

Cupric Aceto-arsenite CuA',3Cu(AsO,),-Schweinfurt green, Imperial green, Mitis green, and when mixed with gypsum or heavy spar, Neuweider green, Mountain green. Used as a pigment, and prepared on the large scale by mixing arsenious acid with cupric acetate and water. 5 pts. of verdigris are made up to a thin paste, and added to a boiling solution of 4 pts. or rather more of arscnious acid in 50 pts. of water. The boiling must be well kept up, otherwise the precipitate assumes a yellow-green colour, from formation of copper arscuite; in that case acetic acid must be added, and the boiling continued a few minutes longer. The precipitate then becomes erystalline, and acquires the fine groen colour peculiar to the acetoarsenite. The salt is insoluble in water, and when boiled with water for a considerable time, becomes brownish and gives up acetic acid. Acids abstract the whole of the copper, and aqueous alkalis first separate blue cupric hy-droxide, which when boiled with the liquid is converted into cuprous oxide, an alkaline arsenate boing formed at the same time.
Didyminm Acetate DiA', 4aq. S.G. 1882.

S. V. S. 207-8.—DiA', aq. Rcd nesdles. S. G. 2·237. S.V.S. 150·6 (Clève, Bl. 2) 43, 365).
Erbium Acetate EbA', 4aq. — Isomorphous with didymium acetate (Thomsen, B. 6, 742).

Iron Acetates .- Ferrous acetate FeA' 4ad obtained by dissolving iron or ferrous snlphide in strong acetic acid, separates on concentration in small colourless silky needles, which dissolve readily in water and quickly absorb oxygen from the air.

Ferric Acetate is not known in the solid state as a salt of constant composition. The dark red solution of ferric hydroxide in acetio acid (Liquor ferri detici) contains a basic salt. The following basic ferric acctates have been distinguished: Fe.A. (OH) obtained by dissolving at 50° the ferrie hydroxide from 1 pt. Fe in 10 pts. acetic acid of 30 p.o. and evaporating at 70°. Amorphous, soluble in alcohol and water (Oudemans, J. 1858, p. 282).—Fe,A',(OH),, probably contained in the red solution formed on treating Fe2A'2(OH)Cl2 (infra) with silver oxide. Becomes syrupy in a vacuum but does not crystallise; decomposes quickly at ord. temp., forming an ochreous jelly (Scheurer-Kestner).-Fe₂A'₂(OH),2Fe₂O₃ is the ochreous deposit formed in a solution of ferrous acetate exposed to the air. Other basic salts appear to be formed in the decomposition of the normal salt by heat or otherwise. A solution of ferrie acetate, quite free from other salts, is not ppd. by boiling (Reinitzer, M. 3, 257).

Acctonitrates (Kestner, A. Ch. [3] 63, 422; 68, 472; J. 1861, 307) .- Formed by mixing solutions of ferric nitrate and acctate in various proportions, or by dissolving ferric hydroxide in various mixtures of acetic and nitric acids. Mostly very unstable, decomposed by boiling with water.—FeA's(OH)2NO3 forms deep red flattened prisms, vory soluble in water and in alcohol, insol. in ether. FeA'4(OH)NO34aq forms redrown rhombio prisms, sol. in water and alcohol, lecomposing on slight rise in temperature. The liformi-diaceto-nitrate Fe2(COH)2(OAe)2(NO3)2 s very soluble in water and alcohol, insol. in

ther, very unstable.

Acetochlorides — Fe, A', Cl, (OH) 3aq is obtained on eautionsly adding nitrie acid to a solution of FcCl, in acetic acid at 86°; also when ferrie hydroxido (1 mol.), acctic acid (1 mol.), and hydrochlorio acid (1 mol.) are digested together at 40° for two or three days. Very hard crystals, black by roflected, red by transmitted light, very soluble in water. With silver oxide they yield ferrio triacetate (K.; Schiff, A. Ch. [3] 66, 136).-Fe2A',Cl2, obtained by dissolving 1 mol. ferrie hydroxide in a mixture of HCl (2 mol.) and C2H4O2 (4 mols.), or by oxidising ferrous chloride dissolved in very strong acetic acid with nitric soid. Yellowish-red crystals, sol. in water and in alcohol, easily resolved into acetic acid and Fe₂A'₂(OH)Cl₂. Heated at 50° for twelve hours with silver nitrate, it is converted into ferric tetraceto-dinitrate:

 $\mathbf{Fe_{z}A'_{4}Cl_{2}} + 2\mathbf{AgNO_{3}} = 2\mathbf{AgCl} + \mathbf{Fe_{z}A'_{4}(NO_{3})_{z}}$ Lanthanum Acetate La A', 1 aq. — Sinall needles (Clève, Bl. [2] 21, 196).

Lead Assates, or Plumbic Acetates.—The normal salt PbA' 3aq, Sugar of lead, Sci saccharum Saturni, [75°] S.G. 2.5.—Prepared by dissolving lead oxide or oarbonate in acetic acid

(Wiohmann, J. 1853, 738).

Properties. - Conoclinio efflorescent crystals, easily coluble in water and in spirit of ordinary strength, insoluble in cold absolute alcohol, com-pletely dehydrated by prolonged boiling with absolute alcohol. An aqueous solution saturated at 15 contains 387-623g, salt in a litre, and has a density of 1 2367 (Michel a. Kraft, J. 1654, p. 296).

Density of the Aqueous Solution at 14°. (Oudemans, Fr. 7, 419; J. 1868, 29.)

Cryst.Salt p.c.	Density	Cryst.Salt p.c.	Density
1	1.0057	20	1·1399
5	1.0317 2	25	1·1808
10	1.0659	30	1·2248
15	1.1018	83	1·2525

Normal lead acetato melts at 75°, begins to give off water with a portion of its acid a little above 100°, and is completely dehydrated at 280°, Above that temperature it decomposes, giving off acetie acid, carbonic anhydride, and acetone, and leaving metallic lead very finely divided and highly combustible. The aqueous solution is partially decomposed by the carbonio acid of the air, carbonate of lead being precipitated, and a portion of acctic acid set free, which prevents further decomposition. The solution is not precipitated by ammonia in the cold, but yields crystals of lead oxide when heated with a large excess of ammonia. Normal lead acetate forms crystalline compounds with chloride and peroxide of lead (Gm. 8, 310). Bromine added to a solution of the normal salt throws down a brown precipitate of PbO2, which, if the liquid be warmed, continues to form till 2 at. Br have been added to 2 mol. PbA'2. The solution then contains lead bromide and acetic acid:

2PbA'2+2H2O+Br2=PbO2+PbBr2+4HA'

(Chapman a. Smith, C. J. 22, 185).

Basio Load Acotates. — Pb.A'. (OH). formed by repeatedly dreuching the normal salt with absolute alcohol; the rosidue crystalliscs from hot absolute alcohol in nacreous six-sided plates, easily soluble in water, sparingly in cold alcohol (Plöchl. B. 13, 1647). Pb A7, Pb O 28q. Lead-vinegar, Acetum Saturni. Prepared by dissolving litharge in the aqueous normal salt, and evaporating at a gentle heat, whereupon it crystallises in needles. According to Wittstein (A. 52, 253) the crystals contain only 1 mol. H.O. Easily soluble in water and in alcohol of 90 p.c. Reacts alkilline. Decomposed by CO. A solution of this salt mixed with alcohol forms Goulard's lotion.-PbA'2PbOaq, obtained by dissolving PbO in normal lead acctate, or by mixing a solution of the normal salt with ammonia. Crystallises in silky needles, soluble in 5.55 pts. water at 100°, insoluble in absolute alcohol (Payen, A. 25, 124; A. Ch. [4] 8, 6; Löwe, J. pr. 98, 385; J. 1866, 235).

An Aceto-chloride Cll'bA' is formed by heating recently precipitated lead chloride with normal lead acctate and acetic acid at 180°, and crystallises in monoclinic prisms. Decomposed by water into lead chloride and the more soluble salt PbA'₂ClPbA'. The analogous compounds, BrPbA' and lPbA' obtained in like manner, form small monoclinic crystals (Carius, A. 125, 87).

Sodio- and Potassio-plumbic Acetates.— PbA'2NaA'3aq. Monoclinic or sals (Rammelsberg, J. 1855, 503).—PbW2PbO4KA' is formed on padding potash-lye (S.G. 1.06) with agitation to a boiling solution of normal lead acetate (S.G. 1.25 to 1.30). Crystalline pulp, moderately soluble in water (Taddei, J. 1847-8, 548). Accto-formate (CHO2)Pb2(C2H2O2),2aq.

Needles, easily soluble in water, sparingly in alcohol (Plōchl. B. 13, 1645).

Lithium Acetates. — LiA'2sq. Rhombio prisms [o. 70°]. Dissolves in less than } pt. water at 15°; in 46 pts. alcohol of density 0.81 at 14° (Fisisch. Zeitschr. f. Physik, 4, 108). According to Rammelsberg (A. 56, 221), the crystals sontain only 1 mol. H₂O.—LiA'HA', obtained by spontaneous evaporation of a solution of the normal salt in glacial acetic acid [99°]. Rotates on water. Under somewhat different ciroumstances it crystallises in small four-sided plates, containing Aq and melting at 85° (Lescœur, Bl. 24, 516).

Magnesium Acetate MgA', 4aq. - S.G. 1.45 (Schröder). Monoclinic, slightly deliquescent, very soluble in water and in alcohol (v. Hauer, J. 1855, 501; Patrouillard, C. R. 84, 553).

Manganous Acstate MnA', 4aq. - Palo red transparent monoclinic plates, permanent in the air, soluble in 3-3.5 pts. water, also in alcohol. S.G. 1.6 (Schröder)

Acid Salt MnA'2HA'2aq (Villiers, Bl. 30, 177). Manganic Acetate MnA'32aq.—Browncrystals (Otto, A. 93, 372; Christensen, J. pr. [2] 28, 14).

Mercury Acetates.—1. Hg.A'2. Obtained by precipitation. Micaceous scales. S. 75 at 13°. -HgA'. Brilliant micaccous lamine. S. 25 at 10°, 36 at 19°, 100 at 100°. Dissolves with partial decomposition in 17.7 pts. alcohol (S.G. *811) at 19° (Gm. 8, 332) .- An aceto-sulphide, HgA', HgS, is precipitated, on passing H,S into a solution of mercuric acetate, as a white crystalline powder.—Mercuro-diammonium Acetate N.H.HgA', H.O. Rectangular plates; casily soluble in water, nearly insoluble in alcohol; smella of acetic acid, and decomposes gradually on exposure to the air (Hirzel, J. 1851, 437).

Nickel Acetate. - Apple-green prisms, soluble in 6 pts. cold water, insoluble in alcohol (Tup-

puti, A. Ch. 78, 164).

Potassium Acetates. - Normal salt KA'. Terra foliata tartari .- S. 188 at 2'; 229 at 13.9°; 492 at 62° (Osann). A boiling saturated solution contains 8 pts. salt to 1 pt. water, and boils at 169° (Berzelius). Soluble in 3 pts. cold and 2 pts. hot aleohol. Exists in anny plantiquies. White, difficult to crystallise, extremely deliquescent, insoluble in ether. Chlorine passed into its aqueous solution liberates CO, and forms a bleaching liquid. On passing an electric current through a concentrated aqueous solution of the salt, bydrogen is evolved at the negative pole, and at the positive a mixture of ethane and CO_{x} . The principal decomposition is: $2(CH_{2},CO_{2}H) = C_{1}H_{4} + 2CO_{2} + H_{2}$, methyloxido and acetate being secondary products (Kolbe, A. 69, 257). On passing CO₂-gas into a solution of the salt in alcohol of 97-100 p.c., a large quantity of potassium carbonate is thrown down, and ethyl acctate is formed (Pelouze, A. 5, 265).

Acid Potassium Acetate KA'HA' is formed when the normal acetate is evaporated with an excess of strong acetic acid, and separates in needles or lamino, or in long flattened prisms. Very deliquescent; melts at 148°, and decomposes at 200°, giving off pure AcOH. This affords an easy method of obtaining glacial acetic acid. Acid potassium acetate is also formed when the normal salt is distilled with

acids decomposes the salt thus produced. Henc when butyrio or valerio aoid is mixed with aceti acid, a separation more or less complete may b offected by half neutralising the liquid wit potash and distilling. If the acetic acid is it excess, acid potassiom acetate alone remains be hind, the whole of the butyrio or valerio aois passing over, together with the remainder of the acetic acid. If, on the contrary, the other acid is in excess, it passes over unmixed with acetiacid, and the residue consists of potassiun acetate mixed with butyrate or valerate. By re peating the process a certain number of times cither on the acid distillate or on the acid separated from the residue by distillation with sulphurio acid, complete separation may be effected Acetic acid, therefore, is an exception to the rule that when a mixture of fatty acids and thoir potassium salts is boiled the most volatile acids distil over (Liebig, A. 71, 355).-KA'2HA' [112°]. S.G. 14. Deliquescent plates (Lescœur, Bl. 22, 156). Anhydrous Potassium Diacetate or Potassium Pyroacetate K₂C₈H₁₂O₇=2KOAo.Ac₂O₇ prepared by dissolving melted KOAo in boiling acetic anhydride, forms colourless needles very soluble in water, less deliquescent than normal potassium acctate. Decomposed by heat, giving

off Ac.O (Gerhardt, A. Ch. [3] 37, 317). Rhodium Acetate RhA',21aq (Claus, J. 1860,

Rubidium Acetate RbA' .- Plates (Grandeau, J. 1863, 184).

Samarium Acetate SmA', 4aq. - S.G. 1.94. S.V.S. 205.6. Yellow crystals (Clève, Bl. [2] 43,

Silver Acetate AgA' (S. 1.02 at 14°) separates on mixing the concentrated solutions of AgNO, and NaOAo. Dissolves in hot water, and on cooling separates as naorceus ficxible laminæ. Heated with iodine it is resolved into silver iodide, mothyl acetate, hydrogen acetate, CO2, acctylene, and hydrogen (Birnbaum, A. 152, 111). When dry, it combines with NH, forming

AgA'2NH3 (Reychler, B. 17, 47).

Sodium Acetates. — NaA'3aq. [58°] (123°). S.G. 1·4. S. 26 at 6°, 42 at 37°, 59 at 48° (Osanu). S. (alcohol of S.G. 8322) 2·1 at 18°. Crystallises with 3H2O in monoclinic prisms, melting below 100°. According to Reischauer (J. 1860, 50), the crystals give off the whole of their water in a vacuum at ord. temp. fused salt in camp air quickly takes up about 7H2O, forming a supersaturated solution, whoreas the unfused salt takes up from the air only the original 3H,O. When the aqueous solution of NaA' turns mouldy, oxygon is absorbed, and small quantities of alcohol and formio acid are produced (Béchamp, Z. 6, 438).

The S.G. of solutions containing the following percentages of NaA' is givon by Franz (J. pr.

[2] 5, 297) as follows:

P.C.	s.c.	P.O.	s.G.	P.C.	s.c.
1 3 5 9	1.0058	11	1.0592	21	1·1134
	1.0174	18	1.0697	28	1·1254
	1.0292	15	1.0802	25	1·1874
	1.0390	17	1.0910	27	1·1506
	1.0488	19	1.1018	29	1·688

butyris or valeric acid; but neither of these The S.G. of a saturated solution being 1.1842.

Acid Sodium Acetates (Villiers, Bl. 29, 153; 50, 175; C. R. 85, 1284; Lescour, Bl. 22, 156). 50, 170; U. Ar. 50, 1802; Hesceut, 1... 22, 1007.

—NaA'HA'aq. Cubio (Fehling).—NaA'2HA' or
NaA'2HA'aq. Long needles. [1277]

5NaA'HA'6aq.—4NaA'HA'11aq.—

5NaA'2HA'15aq.

The three last are, perhaps, mixtures.

Strontium Acetates SrA', laq.—Below 15°

it crystallises with 4Aq in monoclinio prisms.

An aceto-nitrate NO,SRA/2H2O forms triclinic crystals (Hauer, J. 1858, 281; Zepharovich, J. 1860, 309). Villiers (Bl. 30, 176) describes the following acid acetates: SrA'_HA'2aq. 3SrA'_4HA'6aq. 2SrA'_3HA'1\faq.

Thallium Acetates .- Thallous acetate TIA'. White silky needles, easily soluble in water and in alcohol, and deliquescent (Crookes, C. J. 27, 149).

Acid Salt TlA'HA' [64°] (Lescour, Bl. 21, 516).

Basic Thallic Acetate TIA', 2TI (OH), 11aq. Colourless plates, readily resolved into acetio acid and thallio oxide.

Tin Acetates .- Tin dissolves slowly in boiling acetic acid, with evolution of hydrogen, and stannons hydroxide dissolves readily in the boiling acid, the solution when evaporated to a syrup and covered with alcohol yielding emall colourless crystals. Stannio hydroxide also dissolves in the acid, the solution when evaporated leaving a gummy mass. Stannio chloride forms a crystalline compound with glacial acetic acid.

Uraninm Acetates .- Uranous acetate. Warty groups of green needles.—Uranic acetate or Uranyl acetate UO2A'21 obtained by heating uranio nitrate till it begine to give off oxygen, dissolving the yellowish-red mass, which still contains NO₂H, in warm concentrated acetio acid, and evaporating to the crystallising point. Crystallises from strongly aoid eclutions in yellow transparent monoclinic prisms, containing UO2A'22aq, which dissolve in boiling water with separation of UO₂, but are reproduced on evaporating the solution. A weaker solution cooled below 10° deposits quadratic octahedrons of UO₂A'₂8aq, which give off 1 mol. II₂O at 200°, the rest at 275°. Double Salts.—NH₄A'UO₂A'₂8aq, NaA' UO, A', (regular tetrahedrons), KA'UO, A' aq (quadratio prisms), are obtained by adding the respective alkaline carbonates to a solution of uranio acetato till a precipitato is formed consisting of alkali-metal uranate, redissolving this in a slight excess of acetic acid, and cooling to crystallisation. The other double salts of the group are formed by boiling the carbonates with uranio acetate till all the UO, is precipitated, redissolving in acctic acid and evaporating.—BaA',2UO,A',caq. Small yellow crystalline epangles, easily soluble in water; give off the crystal-water at 275° (Wertheim, J. pr. 29, 227).—CaA',2UO,A',8aq. Sulphur-J. pr. 29, 227).—CaA',2UO,A',2Saq. Sulphuryellow rhombic crystals, easily solubo in water, permanent in the air, becoming anhydrous at 200° (Weselsky, J. pr. 75, 55).—CdA',2UO,A',2Saq. Diohroic crystals.—PbA',UO,A', 6aq. Tufts of pale yellow needles.—MgA',2UO,A', 8aq. Rectangular prisms.—NiA',2UO,A', 7aq. Emgraldeftee, rhombic crystals.—SrA',3UO,A', 6aq. Sulphy-yellow crystals.—ZnA',2UO,A', 8aq. Sulphy-yellow crystals.—ZnA',2UO,A', 8aq. Sulphy-yellow crystals.—ZnA',2UO,A', 8aq. Sulphy-yellow crystals. phur-yellow crystals, isomorphous with the nickel salt.

MnA', 100, A', 8aq. FeA', 100, A', 7aq.
TlA'2U0, A', 2aq. LiA'U0, A', 3aq.
BeA', 100, A', 2aq. AgA'U0, A', aq.
Zinc Acetate. ZnA', 8aq. [235° 257°] S.C.
1.72.—ZnA', [242°] S.G. 1.84.—Monoclinic lamine. Very soluble in water. May be, abbline as ZnA'2, especially in vacuo (Franchimont, I 12, 11). ZnA'2 may be crystallised, in anhy drous state, from HOAo (Peter a. Rochefontaine Bl. [2] 42, 573).

Yttrinm Acetate YA', 8aq(?) .- Isomorphou with the acetates of didymium and erbiun (Clève).

ALKYL ACETATES. Acetic Ethers.

Methyl Acetate C₃II₆O₂ or MeA'. M.w. 74 (55°) at 754.4 mm. (R. Schiff); (56°3°) at 760 mm (Kopp); (57°3°) (Gartenmeister); (57°5°) at 760 mm. (Elsässer, Porkin). S.G. § 9643 (G.) rou min. (Lisasser, Forkin). S.G. § '9543 (G.) 2 '9577 (E.); 15 '9398 (P.); 20 '9039 (Brühl); 26 '9039 (Brühl); 27 '9586 (P.); 27 '9825 (S.). V.D. 2-563 (for 2-564). C.E. (0^-10^-) 00133 (G.); '00136 (E.). S. 33 at 22° (J. Traube). S.V. 83-66 (S.); 83-7 (E.). #2 13654. R₂₀ 28-78 (B.). H.F.p. 96,720. H.F.v. 94,980. M.M. 3-362 at 22° (P.).

Occurrence.-In crude wood-vinegar (Weid-

mann a. Schweizer, P. 43, 593).

Preparation.-1. By distilling 2 pts. woodspirits with 1 pt. very strong acetic acid and 1 pt. strong sulphuric acid, removing the excess of wood-spirit by means of fused calcium chloride, and rectifying over sodium carbonate (Dumas a. Peligot [1835], A. Ch. [2] 58, 46). 2. By heating H2SO4 (50 c.c.) and MeOH (50 c.c.) to 140° and running in slowly a mixture of equal parts of MeOH and HOAo (Pabst, Bl. [2] 33, 350). 8. By distilling 3 pts. wood-spirit with 14.5 pts. dried lead acetate and 5 pts. strong sulphurio acid, agitating the distillato with milk of lime, treating the supernatant oil with calcium chloride, and rectifying (Kopp, A. 55, 181).

Properties. - Colourless fragrant Soluble in water; mixes in all proportions with

alcohol and ether.

Reactions .- 1. Aqueous solution only slightly decomposed by boiling .- 2. Resolved by caustio alkalis in methyl alcohol and acetic acid .- 3. When poured on pulverised soda-lime it is violently decomposed, with formation of sodium acetate and formate, and evolution of hydrogen .- 4. With sodium it reacts like ethyl acstate (q. v.), yielding as chief products sodium methylate, NaOCH, and methyl sodio-aceto-acetate, COMe.CHNa.COOMe.—5. Decomposed by strong sulphurio acid, becoming hot, giving off acetic acid, and forming methyl sulphurio acid.

Chloro-methyl Acetate CH,ClOAo (115° i.V.). S.G. 1151-1195. V.D. 3.70 (for 3.74).—Made by passing chlorine into methyl acetato at 10°. An oil. Slowly decomposed by water, quickly by alkalis, giving HCl, HOAc, and formio aldehyde:

 $CH_2Cl(OAc) + 2KOH =$ $CH_2O + H_2O + KCl + KOAo.$

With alcoholio KOAo it gives methylene acetate, CH2(OAc), v. formio aldehyde &L. Henry, B. 6, 740).

Di-chloro-methyl Acetate CHCl. OAc (145% 148°), S.G. 1.25, is formed by passing dry chlorine through methyl acetate at a gentle heat. Colourless, pungent-smelling liquid. Recomposed slowly by water, quickly by aqueous potash, acetic, and hydrochloric acids, CHCl,(OAc) + 5KOH =

CH2(OK)O+H2O+2KCl+KOAc

(Malaguti, A. 32, 47). Tricktorinated Methyl Acetate C₂H₂Cl₂O₂ (145°). Laurest, A. Ch. [2] 73, 25. Per-chlorinated Methyl Acetate

CCl.O.CO.CCl. Formed by prolonged action of Cl on methyl acetate v. tri-chloro-acetic acid.

Ethyl Acetate C,H,O₂ or EtA'.—Acetic ther. M.w. 88. (75·5'-76·5°) at 745·5 mm. (R. Schiff); (77·1°) at 760 mm. (Elsässer); (77·5°) (Gartenmeister). S.G. § 9253 (G.); 2 9239 (E.); 9007 (Brühl); 15 9072, 25 8971 (Perkin); ** 8306 (S.) V.D. 3.087 (for 3.079). S.H. 48. C.E. $(0^{\circ}-10^{\circ}) \cdot 001263$ (E.). S. 6 at $17 \cdot 5^{\circ}$. S.V. $105 \cdot 7$ (S.); $106 \cdot 1$ (G.); $106 \cdot 15$ (E.). $\mu_{\beta} 1 \cdot 3771$. $\mu_{\infty} 35 \cdot 46$ (B.). H.F.p. 114,710. H.F.v. 112,290. Formation.—(Lauragais, J. d. Scavans, 1759, 324; Thenard, Mem. d'Areueil, 1, 153; Dumas a. Bonllay, J. Ph. 14, 113; Liebig, A. 5, 84; 30,

144; Malaguti, A. Ch. [2] 20, 367; [3] 162, 58). 1. By heating alcohol with acetic acid or with an acctate and strong sulphuric acid. 2. By distilling calcium or potassium ethyl-sulphate with

glacial acetic acid (Liebig).

Preparation.—1. By distilling a mixture of 3 pts. potassium acctato, 3 pts. absolute alcohol, and 2 pts. sulphuric acid; or 10 pts. sodium acetate, 6 pts. alcohol, and 15 pts. sulphurie acid; or 16 pts. dry lead acetate, 41 pts. alcohol, and 6 pts. sulphuric acid. The acid is first mixed with the alcohol, and the liquid is poured upon the pulverised salt; the mixture is then distilled to dryncss, and the product is purified by digestion with calcium chloride and rectification.-2. Frankland a. Duppa prepare ethyl acetate by gradually pouring a mixture of 3.6 kilo. of 97-p.c. alcoliol, and 9 kilo. strong sulphnric acid, on 6 kilo. sodium acetate previously fused and dried, leaving the mixture at rest for 12 hours, then distilling and rectifying the distillate (which is free from alcohol and amounts to 6 kilo.) over fused and pulverised calcium chloride. The hest mode of anixing the alcohol and sulphuric acid is to pour the alcohol through a narrow glass tube to the bottom of the vessel containing the acid, stirring the liquid continually by means of the tube. It is best to leave the etbyl-sulphuric acid thus formed for 24 hours before pouring it on the sodium acetate. 8. A mixture of alcohol and acctic acid in molecular proportions is allowed to run into sulphuric acid at 130°, whereby ethyl-sulphuric acid is first formed, and this with the acetic acid torms ethyl acetate, which distils over, leaving the sulphurie acid to be further acted on by tho alcot of. By this process 10 g. sulphuric acid yield 232 g. ethyl acetate (Eghis, B. 6, 1177; Pabst, Bl. [2] 33, 350).

Properties .- Colourless fragrant liquid. Soluble in 10 pts. water at ord. temp.; dissolves *086 pts. of water, freely miscible with alcohol and ether.

Reactions .- 1. Burns with yellowish flame. 2. By dilute ehromic acid it is oxidised to scatic acid C₄H₄C₂ + O₂ = 2C₂H₄O₄ (Chapman a. Thorp, C. J. 19, 484).—3. Permanent in the air Thorp, C. J. 19, 484).—3. Permanent in the air when dry, but gradually decomposing when moist into alcohol and acctic acid; more quickly in chloro-ethyl alcohol and Accil at 180°. Rectified

violently by alcoholic potash, yielding formic, contact with alkalis.—4. Converted by heating apetic, and hydrochloric acids, with sulphuric acid into ethyl oxide and acetic acid; with hydrochloric acid into acetic soid and ethyl chloride .- 5. The vapour passed over zincdust at 800° 850° gives acetone Co, hydrogen and ethylene (Jalin, B. 13, 2107).—6. With lime in sealed tubes at 250°-280° it yields butyric acid as chief product, caleium acetate and ethylate as intermediate products:

2CaO + 2EtOAo = Ca(OAo), + Ca(OEt), = Ca(OH), + Ca(OH), + Ca(O, H), O,), (Lubavin, Bl. [2] 34, 679), -7. With alkaling hydroxides it yields acetic acid and cthyl alcohol; with the anhydrous oxides, acetic acid and a metallio ethylate:

 $2\text{EtOAc} + \text{Ba}(\text{OH})_2 = \text{Ba}(\text{OAc})_2 + 2\text{EtOH}$; and $2\text{EtOAc} + 2\text{BaO} = \text{Ba}(\text{OAc})_2 + \text{Ba}(\text{OEt})_2$. (Berthelot a. Flourieu, A. Ch. [3] 17, 80).—

8. With a mixture of lime-water and chloride of lime (bleaching powder), it yields ohloroform (Schlagdenhauffen, J. Ph. [3] 36, 190) .- 9. With alcoholio KUS it forms, on heating, KOAc and II.S. but no morcaptan (C. Gottig, J. pr. [2] 33, 90) .- 10. With sodium-ethylate, forms, at 130°, sodio-accto-acctio other. - 11. Ethyl acetate heated with sodium dissolves the metal, and the whole solidifies to a crystallino mass of sodium ethylate and ethyl sodio-aceto-acetate CaH, NaO2. The reaction is either

 $2(C_2H_5, O.Ac) + Na_2 = NaOC_2H_5 + C_8H_9NaO_3 + H_2;$ or $3(C_2H_3, O.Ac) + Na_4 = 3NaOC_2H_3 + C_4H_4NaO_4$. The quantity of hydrogen evolved varies considerably according to the temperature and pressure at which the reaction takes place, and the proportions of the materials used; sometimes no gas is evolved (equ. 2), and under no circumstances yet observed is the quantity of hydrogen evolved exactly equivalent to the sodium dissolved, as required by the first equation. Probably, therefore, the two reactions generally take place together (see, further, Aceto-Acetic ACID).-12. With iodine and aluminium foil ethyl acetato yields ethyl iodide and aluminium acetate, $6\text{EtOAc} + \text{Al}_2 + 3 \text{I}_3 = 6\text{EtI} + \text{Al}_2(\text{OAc})_{\bullet}$ and a similar reaction takes place with all the alkyl acetates of the series $C_nH_{2n+1}\text{OAc}$ (Gladstone a. Tribe, $C_nJ_n=0.3$, 357).—13. Ethyl acetate combines with titanic chloride in various proportions (Demarcay, Bl. [2] 20, 127; C.R. 70, 1414).

CHLORINATED ETHYL ACETATES .- Chlorine is abundantly absorbed by ethyl acctate, and acts strongly upon it, even at ordinary temperatures, replacing two or more atoms of hydrogen; the action is accelerated by heat and by direct snnshine. Seven compounds have been described as thus formed, containing 2 to 8 at. chlorine in place of hydrogen, but only three of them have been obtained of constant composition, viz.,

those containing 2, 3, and 5 at. chlorine.

Dichlorethyl Acetate C.MgCl.O. C,H,Cl,C,H,O, is the product formed when othyl acetate is kept cool and in the shade during the action of the chlorine. Transparent oil. S.G.

301 at 12° (Malaguti, A. Ch. [2] 70, 367).

Trichloro-ethyl-acetate & ICL_CHCL.OAc,
metameric with ethyl tri-CHLORO-ACETATE (g. v.) is formed by the action of chlorine at 120°, in presence of iodine, on ethylidene aceto-chloride, CH, CHCl.OAc (Kessel, B. 10, 1999).

s secus (Garzarolli-Thurnlackh, A. 210, 63). | iodides. The following table shows their boiling Faming HNO, conperts it, at 15°, into tri-chloro-asetic acid. KOH forms tri-chloro-ethyl-glycol-

lic acid, CCl_OH_O.CH_CO_H.

Octo-chlorinated Ethyl Acetate
C.Cl_O, i.e. GCl_C.Cl_O, im slowly formed on exposing the dichlorinated other, together with ohlorine, at 100° to height summer sunshins. The product, after distillation in a stream of carbon dioxide to remove excess of chlorins, forms a colourless pungent oil which remains liquid bslow 0°. S.G.1.79 at 25°. Boils, with partial decomposition, at 245°. Its vapour passed over fragments of glass heated to 400° ie cenverted into the isomeric compound chloraldshyde C,Cl,O = CCl,COCl. It is decomposed by water and moist air, and more completely by KOH, into hydrochloric and trichloracetic neids: C_4Cl_4 , $C_4Cl_3O_4 + 2H_2O = 2HCl + 2(CCl_3, CO_4l)$ (Leblanc, A. Ch. [3] 10, 197; Malaguti, ib. 15, 258).

The following chlorinated acetic ethers ars also known: the compound of ALDERYDE with AcCl, grycor chloro-acetin, and the ethyl salts

of the three CHLORO-ACETIC ACIDS.

BROMINATED ETHYL ACETATES C.H.Br.O. = CH_Br.CO.OCHBr.CH, (bromethyl bromacetate), formed on heating sthylidene acetate-chloride CHMeCl(OAc) with bromine at 100°-103°, boils under reduced pressure at 130-135°, and dissolves in boiling water, with formation of aldehydo, acetic acid, crotonaldshyds, acetal, ethyl bromids, and HBr. The crotonaldehyde and acetal are secondary products formed from acetaldehyde, produced in the first iustance as shown by the equations:

CH_Br.CO.OCHBr.CH₃ + C_H₂OH = CH_Br.COOC_H_+ + CH_2.CII Br(OH)

and CH, CHBr(OH) = HBr + CH, CHO. Cri- and Tetra-brominated Ethyl Acetates C,H,Br,O, and C,H,Br,O,, formed by the action of 1 or 2 mol. bromine at 120° and 160°, respectively, on C2H6Br2O2, and freed from absorbed HBr by heating in a stream of carbon dioxide, ars oily strongly fuming liquids, partly decomposing on distillation; decomposed also by water and alcohol, the products containing substances which reduce ammoniacal silver solution, whence it appears that both these ethers produce aldehydes. The pentabrominated compound C₄H₄Br₄O₂, probably CH₂Br₂COOCBr₂CHBr₂, formed by heating C₄H₄Br₄O₂, with 1 mol. bromine at 170°, is a liquid which scarcely fumee in the air (176°). Its product of decomposition by water does not reduce ammoniacal silver so-Intion. Heated with excess of bromine, it forms C.H.Br.O. (195°-198°) (Kessel, B. 10, 1994; 11, 1917). Other brominated acetic ethers are CH, CHBr.OAc v. ALDEHYDE, CH.Br.CH2.OAc v. GLYCOL, and the ethylic BROMO-ACETATES.

Ethyl Ortho-acetate CH₂.C(OEt),... Tricthylic acetats; (1829), S.G. 22 94, forused, together with CH₂Cl.C(OEt), by heating CH₂.Ccl, with NaOEt in a scaled tube at 110°. Fragrant liquid. Decomposed by water into alcohol and

acetic acid (Genther, J. 1870, 636).

The acetates of the higher alkyls, CaH2n+1, are analogous in their propertice and reactions to ethyl acetate, and are obtained, in like flanner, other by heating the corresponding alcohols with acetic and sulphuric acid, or by the action of silver acetate on the corresponding alkyl

۱ą.,

Iso-ceryl Acctate:		
Butyl Acetates O.H.OAc: Normal Primary: Mc.CH.OAcor OIL.PROAC SCOONDARY: Methyl ethyl carbyl acetate McEDH.OAC Tertiary: Trimethyl-oarbyl acetate CMc.OAC. Amyl Acetates O.H.OAc Normal Primary: McCH.ORO Normal Primary: McCH.ORO Secondary: Diethyl carbyl acetate Et.OH.OAC Mcthyl-Isopropyl-carbyl acetate McPrOHOAC Tertiary: Dimethyl-carbyl acetate McCOAC Heryl Acetates O.H.OAC McHyl-lardyl-carbyl acetate McCOAC McHyl-lardyl-carbyl acetate McCOAC McHyl-lardyl-carbyl acetate McCOAC McHyl-lardyl-carbyl acetate KtPrOHOAC McHyl-lardyl-carbyl acetate KtPCHOAC McHyl-lardyl-carbyl acetate KtPCHOAC McHyl-lardyl-carbyl acetate McCoH, DOAC Ethyl-carbyl acetate McCoH, DOAC McHyl-lardyl-carbyl Mc	Propyl Acetates C. H.OAo: B.P.	8.G.
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Methylecate		
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metholicate Metholication Metholication of the method of t	OII.PrOAo.	
metholicate Metholication Metholication of the method of t	or CH ProAc 0-85	96
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148-40 1370 148-40 137	acetate CMe, OAc	
Met(Cll.), OAc 137° 0.9837 130° 130° 130° 132° 0.9837 132° 0.9837 132° 0.9837 132° 0.9837 132° 0.9837 132° 0.9990 132° 125°		
1370 0-8837 1370 0-8837 1370 1320 132	March 1 140 4 10 00	63 _
1250 1250	Isoprimary:	37
1250 1250	Secondary: Diethyl - carbyl 1200 0-00	_
Methyl-propyl-carbyl acateate Mel'POHOAo Tertiary: Dimethyl: ethyl- carbyl acetate Me_BEC.OAo Hexyl Acetates O.H. OAo: Normal Primary: Mm(CH.),OAC Secondary: Methyl-butyl- carbyl acetate Mc(DH.),CHMcOAO Methyl-tert-butyl-carbyl or Pinneolyl acetate Mc(DHOAoCMe_Betyl-carbyl) acetate KEPCHOAc Heptyl Acetates: Normal from n. heptane Do. from Chanthol Methyl-iso-darbyl acetate Mm(CH.),CHMAC Methyl-iso-darbyl acetate Mc(CH.),CHMAC Methyl-iso-darbyl acetate Mc(CH.),CHMAC Methyl-iso-darbyl acetate Mc(CH.),CHMAC Bethyl-iso-dutyl-carbyl acetate Eth (C.H.),CHMAC Bethyl-iso-dutyl-carbyl acetate Mc(C.H.,CHMAC Methyl-iso-darbyl Mc(C.H.,CHMAC Mc(C.H.,C	Methyl-isopropyl-carbyl	
Tertiary : Dimethyl - ethyl-carbyl acetate Me, Etc. O.6 Heryl Acetates O.H. 100A0 : Normal Primary : Me(OH. 1, OA0 : Methyl acetate Me(OH. 1, OA0 Methyl-tert-butyl-carbyl acetate McOlloAccMe. Ethyl-propyl-carbyl acetate McOlloAccMe. Ethyl-propyl-carbyl acetate KtPrOlloAc Methyl-iso-dutyl-carbyl acetate KtPrOlloAc Methyl-iso-dutyl-carbyl acetate Me(OH. 1, OCH. OA. Ethyl-iso-dutyl-carbyl acetate Me(OH. 1, OCH. OA. OA. OA. OA. OA. OA. OA. OA. OA. OA	ncetate McPrCHOAc	
carbylacetate Me, EtC.OAc Heryl Acetates O, H ₁₀ OAc Normal Primary : Mc(OH ₂ , OAc Mc(OH ₂ , OAc	acetate MePrOHOAc 133° 0-92	92 p
Heryl Acetates O. H. 10 Ac. Normal Primary : Mc(CH,). Oac Mc(CH,). Oac Scoondary : Methyl-batyl- carbyl acetate Mc(CH,). CH Mc O. Ac Methyl-tert-butyl-arrhyl or Pinacolyl acetate Mc(CH,). CH O. Ac Heptyl Acetates Normal from n. heptane Do. from Chanthol Methyl-inolanyl-carbyl acetate Me(C, H.,). CH O. Ac Methyl-inolanyl-carbyl acetate Methyl-in	carbyl acetate Me_EtC.OAc 1250 0.89	09 "
Normal Primary : 109-50 0-8890 at 170		
Secondary: Methyl-butyl- carbyl accetate Mc(OII_s),CliMc.OAo Methyl-tert-butyl-carbyl or Pinacolyl acctate Mc.CliOAo.CMcs Ethyl-propyl-carbyl acctate ktPrCiIOAc Heptyl Acctates: Normal from n-heptane Do. from Chambiol Methyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl acctate Mc(C, H_s,Cli.OAc Ethyl-iso-butyl-carbyl Acctates: Normal (1, 10, OAc, Crystal- iline, (125°) at 15 mm. Dodccyl Acctate: C_tyl Acctate: Iso-ceryl Acctate: Iso-ceryl Acctate: Iso-ceryl Acctate:	Normal Primary:	90 at 179
carbyl accetate Mc(Oll_a)_CHM-O.Ao Methyl-tert-butyl-carbyl or Pinncolyl acctate Mc(OllOAcCMe_ Ethyl-propyl-carbyl acctate RtPrCHOAc Heptyl Acctates: Normal from n-heptane Do. Isom Chanthol Methyl-iso-dutyl-carbyl acctate Mc(C,H_a)CH.O.Ac Ethyl-iso-butyl-carbyl acctate Et (C,H_a)CH.O.Ac Ethyl-iso-butyl-carbyl acctate Et (C,H_a)CH.O.Ac Ethyl-iso-butyl-carbyl acctate Mc(C,H_a)CH.O.Ac Ethyl-iso-butyl-carbyl acctate Mc(C,H_a)CH.O.Ac Ethyl-iso-butyl-carbyl acctate Mc(C,H_a)CH.O.Ac Ethyl-iso-butyl-carbyl acctate Mc(C,H_a)CH.O.Ac Fannyl Acctates: Normal (from oil of Herneleum) Methyl-liceyl-carbyl acctate Mc(C,H_a)CH.O.Ac Ethyl-liceyl-carbyl acctate Mc(C,H_a)CH.O.Ac Ethyl-liceyl-carbyl acctate Mc(C,H_a)CH.O.Ac Ethyl-liceyl-carbyl acctate Mc(C,H_a)CH.O.Ac Solid, (151°a) Acctates: Normal C, H_a,O.Ac. Crystal- line. (125°a) at 15 mm. Dodccyl Acctate: Normal C, H_a,O.Ac. Solid, (151°a) at 15 mm. Cetyl Acctate: C, H_a,O.Ac. (31°a) (223°a) at 15 mm. Iso-ceryl Acctate: Iso-ceryl Acctate: Iso-ceryl Acctate: Iso-ceryl Acctate: Iso-ceryl Acctate:	Secondary : Methyl-butyl-	DO 80 20
of Princely acetate M.CHOAccMe. Ethyl-propyl-carbyl acetate ktrCcHOAc Heptyl Acetates: Normal from a heptane Do, from Channhol Methyl-amyl-carbyl acetate Me(C,H ₁₁ ,OH.OAc Helyl-lamyl-carbyl acetate Me(C,H ₁₁ ,OH.OAc Normal from oil of Hernelrum) Methyl-lacyl-carbyl acetate Me(C,H ₁₁ ,OH.OAc Ennyl Acetates: From Ennane in petroleum Ethyl-lacyl-carbyl acetate Me(C,H ₁₁ ,OH.OAc Normal C ₁₀ H ₁₀ OAc, Crystal- line. (125°) at 15 mm. Dodcyl Acetates Normal C ₁₀ H ₁₀ OAc, Solid. (151°) at 15 mm. Cetyl Acetates C ₁₁ H ₁₀ OAc, Need'es, [18:8°] (200°) at 15 mm. Octadecyl Acetates O ₁₁ H ₁₀ OAc, [31°] (223°) at 15 mm. Iso-ceryl Acetates:	carbyl acetate 111559_1579 0-97	78 at 0°
Ant. Of Colors and Col		
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Heptyl Acctates: 1807 1929 1920 19	acetate EtPrOliOAc 1500	
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Methyl-iso-duryl-carbyl acetate Me(G,H,)CH.OAc Ethyl-iso-duryl-carbyl acetate Me(G,H,)CH.OAc Ethyl-iso-duryl-carbyl acetate Me(G,H,)CH.OAc Normal (from oil of Herneleum) Methyl-iso-yl-carbyl acetate Me(G,H ₁)CH.OAc Ethyl-keryl-carbyl acetate Me(G,H ₁)CH.OAc Ennyl Acetates: Normal G, H, OAc Ethyl-heryl-carbyl acetate Me(G,H ₁)CH.OAc Ethyl-heryl-carbyl acetate Normal G, H, OAc Corystal- ine. (126°) at 15 mm. Dodceyl Acetates Normal C, H, OAc Cetyl Acetates C, H, OAc Cetyl Acetates C, H, OAc Corystal- ine. (126°) at 15 mm. Cetyl Acetates C, H, OAc Corystal- ine. (126°) at 15 mm. Cetyl Acetates C, H, OAc Corystal- ine. (126°) at 15 mm. Cotadecyl Acetates O, H, OAc Solid. Iso-ceryl Acetates	Normal from n-heptane . 1809	
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tate Mc(C, II, CH.O.Ac Ethyl-is-obtyl-earbyl acc- tate Et.C, H ₊ CillOAo Cotyl Acctates: Normal (from oil of Hernelcum) Mcthyl-incyl-carbyl acctate Mc(C, II, CH.O.Ac Ethyl-incyl-carbyl acctate Ethyl-incyl-carbyl acctate Normal C, oil, OAO. Crystal- line. (125') at 15 mm. Dodcyl Acctate: Normal C, oil, OAC. Solid. (181') at 15 mm. Cetyl Acctate: C, H ₊ , OAO. Solid. (181') at 15 mm. Cetyl Acctate: O, H ₊ , OAO. [31°] (223°) at 15 mm. Iso-ceryl Acctate:		
Octul Acctates: Normal (from oil of Herneleum) Methyl-hexyl-carbyl acctate) McC ₂ H ₁₁ /CH()Ao Ethyl-hexyl-carbyl acctate R(C ₂ H ₁₁)CHOAc Normal C ₁₀ H ₁₂ OAc, Crystalline, (125') at 15 mm. Dodcyl Acctates Normal C ₁₀ H ₁₂ OAc, Solid, (181') at 15 mm. Cetyl Acctates C ₁₀ H ₁₂ OAc, Needies, [18'50'] (200') at 15 mm. Cetyl Acctates O ₂ H ₁₂ OAc, Salid, (121') at 15 mm. Cotadecyl Acctates O ₂ H ₁₂ OAc, Solid, (18') at 15 mm. Iso-ceryl Acctates: O ₃ H ₁₂ OAc, Solid, (18') at 15 mm.	tate Mc(C.H.,)CH.OAc [1 207	0 at 23°
Normal (from oil of Herneleum). Methyl-hexyl-earbyl acetate Mic(C,H ₁₁)CH.OAo. Ennyl Acetales: From Ennane in petroleum. Ethyl-hexyl-earbyl acetate MC, C,H ₁₂)CH.OAc. Decyl Acetale: Normal C ₁₂ H ₁₂ OAc. Crystalline. (125°) at 15 mm. Dodcyl Acetale: Normal C ₁₂ H ₁₂ OAc. Solid. (151°) at 15 mm. Cetyl Acetale: C ₁₄ H ₁₃ OAc. Need'es. [188°] (200°) at 15 mm. Octadecyl Acetale: O ₁₂ H ₁₃ OAc. [31°] (223°) at 15 mm. Iso-ceryl Acetale:	tate Et(C,H,)CllOAo.	
Normal (from oil of Herneleum). Methyl-hexyl-earbyl acetate Mic(C,H ₁₁)CH.OAo. Ennyl Acetales: From Ennane in petroleum. Ethyl-hexyl-earbyl acetate MC, C,H ₁₂)CH.OAc. Decyl Acetale: Normal C ₁₂ H ₁₂ OAc. Crystalline. (125°) at 15 mm. Dodcyl Acetale: Normal C ₁₂ H ₁₂ OAc. Solid. (151°) at 15 mm. Cetyl Acetale: C ₁₄ H ₁₃ OAc. Need'es. [188°] (200°) at 15 mm. Octadecyl Acetale: O ₁₂ H ₁₃ OAc. [31°] (223°) at 15 mm. Iso-ceryl Acetale:	Octul Acetates:	•
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From Ennance in petroleum Ethyl-hevyt-carbyl acetate Ethyl-hevyt-carbyl acetate Ethyl-hevyt-carbyl acetate Ethyl-hevyt-carbyl acetate Ethyl-hevyt-carbyl acetate Normal C ₁₀ H ₁₀ OAc, Crystal- iline, (125°) at 15 mm. Dodccyl Acetate Normal C ₁₂ H ₂₀ OAc, Solid. (181°) at 15 mm. Cetyl Acetate : C ₁₀ H ₁₀ OAc, Need'es, [188°] (200°) at 15 mm. Octadecyl Acetate: O ₁₀ H ₁₀ OAc, [31°] (223°) at 18 mm. Iso-ceryl Acetate:	Methyl-hexyl-carbyl scetate)	•
From Ennane in petroleum Ethyl-hevyt-earbyl acctate Kt(C _a I _a , OHOAc Decyl Acctate: Normal C ₁₀ (I ₂ , OAc, Crystalline, (125°) at 15 mm. Dodccyl Acctate: Normal C ₁₀ I ₂₀ OAc, Solid, (181°) at 15 mm. Cetyl Acctate: C ₁₀ I ₂₀ OAc, Needles, [185°] (200°) at 15 mm. Octadecyl Acctate: O ₁₀ I ₂₀ OAc, [31°] (223°) at 15 mm. Iso-ceryl Acctate:	Me(C _e H _{1s})CH,OAo . }	
Decyl Acetate: Normal C ₁₀ I ₁₀ OAc. Crystal- line. (126°) at 15 mm. Dodccyl Acetate: Normal C ₁₀ I ₁₀ OAc. Solid. (151°) at 15 mm. Cetyl Acetate: C ₁₀ I ₁₀ OAc. Need'es. [18°5°] (20°) at 15 mm. Octadecyl Acetate: O ₁₀ I ₁₀ OAc. [31°] (223°) at 15 mm. Iso-ceryl Acetate:		
Decyl Acetate: Normal C ₁₀ I ₁₀ OAc. Crystal- line. (126°) at 15 mm. Dodccyl Acetate: Normal C ₁₀ I ₁₀ OAc. Solid. (151°) at 15 mm. Cetyl Acetate: C ₁₀ I ₁₀ OAc. Need'es. [18°5°] (20°) at 15 mm. Octadecyl Acetate: O ₁₀ I ₁₀ OAc. [31°] (223°) at 15 mm. Iso-ceryl Acetate:	Ethyl-hexyt-carbyl acctate	
Normal C ₁₀ (I ₂₀ ()Ac. Crystal- line. (125°) at 15 mm. **Dodceyl Acetate: Normal C ₁₂ H ₂₀ ()Ac. Solid. (151°) at 15 mm. **Cetyl Acetate: C ₁₀ H ₂₀ ()Ac. Need'es. [18·5°] (200°) at 15 mm. **Octadecyl Acetate: O ₁₀ H ₂₀ ()Ac. [31°] (223°) at 15 mm. Iso-ceryl Acetate:	Kt(C ₄ ll ₁₂)CHOAc	2 36 Om
Dodceyl Acetate: Normal C ₁₂ H ₃₀ OAc. Solid. (151°) at 15 mm. Cetyl Acetate: C ₁₆ H ₃₀ OAc. Need'es. [18:5°] (200°) at 15 mm. Octadecyl Acetate: O ₁₆ H ₃₁ OAc. [31°] (223°) at 15 mm. Iso-ceryl Acetate:		
Dodceyl Acetate: Normal C ₁₂ H ₃₀ OAc. Solid. (151°) at 15 mm. Cetyl Acetate: C ₁₆ H ₃₀ OAc. Need'es. [18:5°] (200°) at 15 mm. Octadecyl Acetate: O ₁₆ H ₃₁ OAc. [31°] (223°) at 15 mm. Iso-ceryl Acetate:	Normal C ₁₀ H ₂₁ OAc. Crystal-	
Normal C ₁₂ H ₂₀ OAc. Solid. (151°) at 15 mm. **Cetyl Acctate:* C ₁₆ H ₂₁ OAc. Need'es. [13·8°] (200°) at 15 mm. **Octadecyl Acctate:* O ₁₆ H ₂₁ OAc. [31°] (223°) at 15 mm. Iso-ceryl Acetate:		
Cetyl Acetate: C. [Han OAc. Need'es. [18-80]] (200") at 15 mm. Octadecyl Acetate: On Han OAc. [310] (2230) at 15 mm. Iso-ceryl Acetate:		
C _{1.} H _{1.0} OAo. Need'es. [18-8°] (200°) at 15 mm. Cotadecyl Acetate: O _{1.} H _{1.0} OAo. [31°] (223°) at 18 mm. Iso-ceryl Acetate:	(151°) at 15 mm.	
Octadecyl Acetate: OnHanOAc. [310] (2230) at 15 mm. Iso-ceryl Acetate:		
Octadecyl Acetate: OnHanOAc. [310] (2230) at 15 mm. Iso-ceryl Acetate:	C ₁₀ H ₃₁ OAo. Need'es. [18·5°] (200°) at 15 mm.	
O.,H.,OAO. [31°] (223°) at] 15 mm. Iso-ceryl Acetate:	7.	
Iso-ceryl Acetate:		
	15 mm.	
	Iso-ceryl Acetate:	
C ₂₇ H ₃₈ OA0 [57°]	C27H34OA0 [57°]	

AUYI ASSURES V. ALLYL ACETATE. Phonyl Acetate v. PHENOL. Benryl Acetate v. BENATA ACETATE. Methylene Di-acetate v. FORMIO ALDEHYDE. Ethylene Acetates v. GLYCOL. Polyeti vlenic Acetates v. Grycor. Ethylené Aceto-butyrate v. Glawol. Ethylene Aceto-chloride v. GLYCOL. Propylene Acetate v. PROPYLENE-OLYCOL. Butylenc Acetate v. OXY-BUTANES. Amylene Acetate v. OXY-PENTANES. Glyceryl Acetates v. GLYCERIN.

Substitution products of Acetic Acid v. BROME-ACETIC ACIDS, CHLORO-ACETIC ACIDS, IODO-ACETIO ACIDS, CYANO-ACETIO ACID, SULPHO-CYANO-ACETIC ACIDS, SULPHO-ACETIC ACID.

Other derivatives of Acetic Acid v. ACETYL BROMIDE, BROMO-ACETYL BROMIDE, CHLORO-ACETYL BROMIDE, CYANO-ACETYL BROMIDE, ACETYL CYANIDE, ACETYL CHLORIDE, ACETYL IODIDE, Di-AZO-ACETIC ACID. H. W.

ACETIC BROMIDE v. ACETYL BROMIDE. ACETIC CHLORIDE v. ACETYL CHLORIDE. ACETIC CYANIDE v. ACETYL CYANIDE. ACETIC IODIDE v. ACETYL TODIDE.

ACETIC OXIDE of ANHYDRIDE C.H.O. or Ac20.—Acetyl oxide, Acetic acid, Anhydrous acetic acid.—M.w. 102. (137.8°) at 755 mm. (Kopp.); (44.6) at 15 mm., (136.4°) at 760 mm. (Kahlbanm). S.G. 2 1.097, 152 1.799 (K.); 20 1.0816 (Brühl). V.D. 3·47 (for 3·51). μ_g 1·3953. R_∞ 35·82 (B.). H.F.p. 132,850. H.F.v. 130,820.

Formation. - 1. By the action of phosphorus triohloride or oxychloride on potassium acetate, $3KOAc + POCl_3 = K_3PO_4 + 3AeCl$, and AcCl + KOAc = KCl + AcO (Gerhardt, 1853, C. R. 34, 755, 902; A. Ch. [3] 37, 285).—2. From potassium acetato and benzoyl chloride, the first product of the reaction being acetobenzoic oxide, which, if the potassium acetate is somewhat in oxcess, and the mixture is heated to a temperature somewhat above that required for its formation, is resolved into acetic and benzoio oxides: KOAe + BzCl = KCl + AcOBz; and 2AcOBz = Ac2O + Bz2O. Similarly from potassium benzoate and acetyl chloride (Gerhardt).-3. By digesting glacial acetic acid and acetyl chloride in molecular proportions (Kanonnikoff a. Saytneff, A. 185, 192).-4. From lead or silver acetate and carbon bisulphide

 $2Pb(OAc)_2 + CS_2 = 2PbS + 2Ac_2O + CO_2$ (Broughton, Z. 1865, 306).—5. From acetal chloride and barium oxide at 100° (Gal) .- 6. In small quantity by the action of phosphoric anhydride on glacial acetic acfd (Gal; Etard, B. 9, 414).—
7. By the action of lead nitrate on acetyl chloride (Lachowicz, B. 17, 1281).

Preparation .- 1. Acetyl chloride (1 pt.) is run into sodium acctate (1 pt.) or potassium acetate (11 pt.), and the product is distilled. As, however, acetyl chloride is formed by the action of the chlorine compounds of phosphorus on acetates, it is clear that, for the preparation of the anhydride, this chloride need not be quite free from phosphoras oxychloride. It is sufficient, indeed, to add POCl, (3 rts.) directly to an excess of NaOAo (10 pts.) or KOAc (12 pts.) and distil; or to prepare a mixture of POCl, and AcCl, by the action of PCl, (7 pts.) on glacial acetic acis (2 pts.), and distil this mixture with NaOAo (20 pts.) or KOAc (24 pts.). In all these from the isomeric compound, ethylene acetate

modes of preparation it is necessary to heat the mixture strongly towards the and of the distillation, because a portion of the acetic oxide unites with the excess of metallic acetate present, forming a compound which requires a high temperature to decompose it. The acetic oxide thus obtained must be subjected to fractional distillation to free it from residual chlorides and acstic acid (Kekulé. Lehrb. 1, 570).-2. Hentschel (B. 17, 1285) prepares acetio anhydride by passing a stream of carbonyl chloride, COCl2, into fused dry sodie acetate.

Properties.—Colourless, very mobile, strongly refracting liquid, having an odour like that of

glacial acetic acid, but stronger.

Reactions .- 1. With HCl-gas acetic oxide acts strongly at 100°, forming acetic acid and chloride: Ac₂O + HCl = AcOH + AcCl (Gal, A. Ch. [3] 66, 187).—2. With chloring at 100° the products are acetyl obloride and ebloracetic acid:

(C.H₂O)₂O + Cl₂ = C₂H₄OCl + C₂H₄ClO₂ (Gal). Similarly with Br. With iodine no action at 200°, but at higher temperatures HI is given off (Gal).—3. With PCl₃, it yields acetyl chloride: $Ac_2O + PCl_3 = POCl_3 + 2AcCl. - 4$. Heated with solid aluminium chloride it forms acetyl chloride and aluminium acetate:

 $3Ae_2O + AlCl_3 = 3AcCl + Al(OAe)_2$ (Andrianowsky, J. R. 11,116).—5. With pulverised zinc chloride at 100° it yields acetic acid, acetic oxide, and a dark brown residuo having the composition C, H,O (Bauer, J. 1861, 438).—6. By heating with zinc-dust it yields acetone (Jahn, M. 1, 696).-7. Reduced by sodium amalgam it forms aldehyde, and afterwards alcohol:

 $(CH_3CO)_1O + 2H_2 = 2CH_3CHO + H_2O;$ $2CH_3CHO + 2H_2 = 2CH_3CH_2OH$ (Linnemann, A. 148, 249).-8. Heated in CO2 gas at 60° with Cl.SO2.OH, it forms an acid, C.H.SO, & Sal) .- 9. With urea, at the boilingpoint, it forms acetyl carbamide, NHAcCO.NH No reaction with oxamide (Schoitz; Marsh a. Genther, Bl. [2] 10, 460).—10. With nascent zinc-ethyl (2 mol. EtI and 1 mol. Ac.O added to zinc-sodium) it yields thethyl-ethyl ketone:

(COMe)₂O + ZnEt₂ = ZnO + 2(Mo.CO.Et). With zinc-methyl in like manner: acotone, Me.CO.Me (Saytzeff, Z. [2] 6, 104).—11. Forms erystallino compounds with NaHSO, and with NH3. The latter is formed by passing NH, into an ethereal solution of Ac.O at -26° (Loir, C. R. 88, 812).—12. Gives a mirror with ammoniacal AgNO2 (Loir).-13. Decolorises aqueous KMnO4 (Loir).-14. Converted by H.SO, at 130° into sulplio-acetic acid (Franchimont, C. R. 92, 1054).

Compounds.—A. With potassic acciate Ac O2KOAc, obtained by dissolving dry potassium acetato in Ac₂O at 100°, crystallises in needles, and is resolved into its constituents by heat (Gerhardt). B. With aldehydes.—(1.) With acetaldehyde acetio oxide forms the compounds C₂H₄OAc₂O and C₂H₄O2Ac₂O. first is obtained by heating its constituents together in molecular proportion at 180° in a sealed tube, and parified by fractional distillation, washing the portion which distils above 140° washing the portion which distributed with hot water, and drying over CaCl. It is a liquid which boils at 168°, has an allaceous odour, and is resolved by heating with KCH into acetic acid and aldehyde—distinction O.H. (OAc), whith, when similarly treated, yields glycol, C.H. (OH), (Genther, A. 106, 249). The second compound, O.H.O.2Ad,O. formed by heating paraldehyde with Ac.O. at 160°, is a liquid having a density of 1-07 at 10° (Getther, I, 1864, 329). (2) With acrolin.—The compound C.H.OAcO is obtained by heating its configuration in molecular proportion at 100° for six hours, or 1 mol. acrolein chloride with 2 mols. silver acetate at about 160°. Liquid immiscible with water, having a fishy odour and very sharp taste. S.G. 1.076 at 22°; (180°). C₂H₄O2Ao₂O, formed by heating metacrolein with Ac,O at 150°, is an oily liquid boiling at 180° (Hubber a. Genther, A. 114, 35; J. 1860, 306). (3.) With bensaldehydo.—C.H.OAo.O is formed on heating bitter-almond oil with excess of Ac.O at 150° and separates on washing the product with water and potash as an oily liquid, which solidifies to a crystalline mass melting at 44°-45° (Hübner, Z. [2] 3, 277). These compounds may be looked upon as derived from ortho-aldehydes, These compounde may bo XCH(OH)2. Similar compounds will be described in articles on the several aldehydes. C. With other oxides.—Vapour of SO, is absorbed by cooled Ac.O. forming a gummy mass soluble in water. Boric oxide dissolves slowly in Ac.O. forming a vitreous hygroscopic mass. Insoluble tartaric oxide or anhydride, C,H,O,, dissolves at 100° in acctio oxide, forming a syrup. Tho same syrupy product is formed, together with PbCl₂, by the action of AcCl on load tartrate.

Aceto-arsenious Oxide C₄H_cO₃As_cO₃ or Ac.O.O.As is formed by dissolving As_cO₃ in acetic oxide at boiling heat, as a syrupy liquid, which on cooling forms a vitreoue hygroscopic mass.

Aceto-hypochlorous Oxide AcOCI and Aceto-hypoiodous Oxide AcOI have been described as unstable compounds by Schützenberger (C. R. 52, 369; 54, 1026; J. 1862, 240), but their existence has been called in question by Aronhoim (B. 12, 26).

Aceto-silicic Oxide Si(OAc), [110] (148°)

Acto-silicic Oxide Si(OAc), [110] [148] at 6mm. From Ao,O and SiO₂ (Priedel a. Ladenburg, A. 145, 174). Decomposed by water, lent, alcohol, or NH₃, into silien and HOAcAc,O, EtOAc, and NH₂Ac, respectively. A compound, Si(OEt), (OAc) (c. 195°), is formed from Si(OEt), and Ac,O.

Aceto-benzoic Oxide CaHaO, i.e. AcOBz, from acetyl chloride and sodium benzoate, is a heavy oil. Begins to boil at 150°, and is resolved at the same time into Aco and Bz.O. By boiling with water, and more quickly with alkalis, it is converted into acetic and benzoic acids (Gerh. 3, 209). HCl converts it at low temperatures into AcCl and HOBz; at 150° BzCl and HQAc are also formed. Chlorine forms AcCl and elaboro-benzoic acid (Greene, C. N. 50, 61).

A ceto-ciessamic Oxide Ac.O.C.H.O. Obtained like the preceding, which it resembles. Oil, heavier than water, very unstable (Gerhardt, ib. 387).

Aceto-cuminic Oxide Ac.O.C., H110. Like

the preceding (Gerhardt, ib. 509).

Aceto-salicylic Oxide Ac.O.C.H.O. Solid; dissolves in aqueous sodium carbonate, with format in of sodium acetate and ealicylate (Gerardt ib. 319).

distriction of sodium acetate and ealicylate (derical to 319).

Abotic Peroxide C.H.O. or Ac.O.—Prepared a adding BaQ to a solution of acetic anhy-

dride in ether. The mixture is effected gradually, being attended with evolution of heat. The ether is distilled aff at a low temperature, and the fluid which remains is washed with water. It is a viscid liquid with pungent tasta. It decolorises indigo, oxidises mangarous by drate to peroxide, and potassic ferro- to farricyanide. It acts generally as an oxidising agent. It does not reduce CrO, or KMnO. Baryta-water is converted by it into barium peroxide and acetate. It explodes when heated (Brodie, Pr. 9, 863).

H. W.

ACET-IMIDAMIDE v. ACETAMIDINE. ACET-IMIDO-ETHYL-ETHER

CH, C

(97°). Liquid. The hydrochloride is obtained by passing dry HCl-gas into a mixture of acetonitrile and ethyl alcohol (equal mols.) diluted with \(\frac{1}{2} \) their volume of ether, cooled to 0° C. B'HCl, long trimetrio plates, decomposes at about 100° into ethyl chloride and acetamide (Pinner, B. 16, 1654).

ACETIMIDO-NAPHTHYL-AMIDE v. NAPR-THYL-ACETAMIDINE.

ACETIMIDO-TOLYL-AMIDE v. Tolyl-acetamidine.

ACETO-ACETIC ACID CH₃. CO. CH₂. CO₂H or CH₃. C(OH): CH. CO₂H.

Occurrence. In urino of diabetic patients (Geuther a. Rupstein, Fr. 14, 419; Deichmüller, A. 209, 30; Tollens, A. 209, 36; Jaksch, H. 7, 487).

Preparation.—The ethyl ether (4.5 g.) is mixed with water (80 g.) containing KOH (2.1 g.), and after 24 hours the liquid is acidified and shaken with ether (Ceresole, B. 15, 1327, 1872).

Properties.—A thick acid liquid, miscibls with water. At 100° it splits up into CO₂ and acetone. Nitrous acid gas forms CO₂ and iso nitrosoacetone.

Salts.—BaA'₂ aq. Amorphons. V.e. eol. water. Violet colour with FeCl₄.—CuA'₄2aq. Amorphous. Ethyl Aceto-acetate or Aceto-acetic Ether CI₄.—CO.CH₂. CO.Et or CI₄. C(O.H): CH.CO.Et. Di-acetic ether (Geuther, J. 1865, 302), acetone-carboxylic acid (Frankland a. Duppa, A. 138, 211 (180°) (R. Schiff, B. 19, 561); (180°8° cor.) (Geuthor); (180°6° -181°2°) at 754 mm. (Brühl); (152°5° -153°) at 330 mm. (Perkin). S.G. § 1.0256 (B.); § 1.046 (S.); § 1.70317 (P.); § 1.0235 (P.). ps. 1.4253. R_{op} 51·62 (B.). S.V. 153°34 (S.). M.M. 6·501 at 16·25 (P.).

Formation.—The formation of aceto-acetic ether by the saponification of cyano-acetone by alcoholic HCl (Matthews a. Hodgkinson, B. 15, 2679) is denied by James (4. 231, 245).

15, 2679) is denied by James (A. 231, 245).

Preparation.—Ethyl acetate (1kilo.), that this been carefully dried, is treated with sodium (100 g.) in small pieces. As soon as the first reaction abates it is heated with inverted condenser over a water bath for 2½ hours until the sodium is dissolved. Dilute (50 p. o.) acetic acid (550 g.) is then added, and where the liquid is cool, it is mixed with water (500 c.o.). The light oily layer is washed with a little water and fractioned. The yield (175 g.) is small, but much ethyl acetate (400 g.) is recovered (Conrad, A. 186, 214). Aceto acetic ether may be still further purified

by shaking with cono. aqueous NaHSO, with which it combines. Impurities may then be extracted by ether, and the compound of acetoacetlo ether with NaHSO, afterwards decomposed by K.CO. (Elion, R. 3, 246).

The farmation of acote-acetic ether may be

expressed by the equation:

2CH, CO, Et + Na, =

CH, CO, CHNa, CO, Et + NaOEt + H2, the sodio-aceto-acetic ether being afterwards dscomposed by the added acctic acid:

CH_s.CO.CHNa.CO₂Et + HOAc = CH₂.CO.CH₂.CO₂Et + NaOAe.

See also p. 21.

Properties .- A liquid with an agreeable sweet odour. Slightly soluble in water, the liquid giving a violet colour with FeCl. Unlike its ethyl and acetyl derivatives, it forms a crystalline compound with NaHSO, (indicating presence of the ketonic

oarbonyl groap, CO).

Salts.—Aceto-acetic other behaves as a mono-basic acid. This may either be ascribed to the situation of the group CII, between two CO groups, or else by having recourse to the formula CH. C(OH: CH.CO.Et, which represents a compound that might be expected, as a tertiary alcohol, to possess a phenolic character. phenol, it gives a violet colour with FeCl,

Sodio-aceto-acetic Ether CH2.CO.CHNa.CO_Et or CH2.C(ONa):CII.CO_Et. Nasdles. Produced by the action of sedium or sodic ethylate upon aceto-acetic ether in the cold.

Preparation.-Sodium (10 g.) is dissolved in absolute alcohol (100 g.); when cold, dry ether (90 g.), followed by aceto-acetic ether (56.5 g.) diluted with ether (60 c.c.), is added. If the liquid is well stirred with a little water (2 c.c.) solid sodium acetacetic ether separates (Harrow, C. J. 83, 426). The pp. is a hydrate, which becomes dry in an exsiccator (Elion, R. 3, 240)

Reactions.—(a) With iodine in othereal solution it gives di-aceto-succinic ether (q, v, l).—(b) Meated alone or with NaOEt it gives acetone, aceto-acetic ether, NaOAe, and sodic dehydracetato. - (c) With alkyl iodides it forms alkylaceto-

acctic ethers (q.v.): CH₂,CO,CHNa.CO,Et+R/k= NaI + CH, CO CHR'.CO Et.

Other iodo- bromo and chloro- compounds act similarly .- (d) But with tri-phenyl methyl pro-SIDE Ph.CBr it forms CH.CO.C(CPh.), CO.Et (Allen a. Kölliker, A. 227, 110). -(e) Chloroform, in presence of NaOEt forms oxy cviric ethen; $\mathbf{C}_{\mathbf{z}}\hat{\mathbf{H}}_{\mathbf{z}}\mathbf{Me}(\mathbf{OH})(\mathbf{CO}_{\mathbf{z}}\mathbf{H})_{\mathbf{z}}$ [1:3:4:6], the first stage probably being:

2CH, CO.CH No.CO, Et + NaOEt + CHCI, CO_Et.CHAc.CH:C(CO_Et).CO.CH_++3NaCi+HOEt

(Oppenheim a. Pfaff, B. 7, 929; 8, 884; 9, 321;

Conrad a. Guthzeit, A. 222, 249).

Other Salts.—Al(C₆H₅O₄)₂. Needles [76°]. Insol. water, v. e. sol. ether, benzenc or CS₂. May be sublimed.—Co(C₆H₉O₃)... Red pp. Soi. hot henzene or ether.—Cu(C₆H₉O₃)₂ [182]. Green needles (from aloohol). Insol. water, v. sol. benzene, either or CS2. Got by adding Cu(OAc), to a solution of aceto-acetic ether in alcohol, the calculated quantity of ammonia being also added (Conrad a. Guthzeit, B. 18, 19). -Mg(C₆H₂O₄)₂ [2±0°]. From aceto-acetic ether and 'magnesia-mixture.' Plates (from etherby shaking accto-acctic ether with HgQ.--Ni(C_aH₂O₂)₂.

Reactions .- 1. Boiled for a long time, or passed through a rad-hot tube, it forms dehy-

dracetic acid, C.H.O., and alcehol. •
2. Boilsd with alkalis it gives CO., acctone. acetio acid and alcohol, according to the resotions:

 CH_3 .CO. CH_2 . CO_2 Et + 2KOH = CH_3 .CO. CH_3 + K_2 CO₃ + HOEt; CH_2 .CO. CH_2 .CO.Et + 2KOH =2CH, CO.OK + HOEt.

Decomposed by water at 150°, or by strong acids, into CO, acetone and alcohol.

4. Action of sodium alcoholates. - (a) Heated with dry NaOEt, or with alcoholic NaOEt, ethyl acetate is got in small quantity (12 p.o. of the theoretical) (Wislicenus, A. 186, 193; Isbert, A. 234, 160).—(b) 50 g. heated with NaOEt (from 8-9 g. Na) and McOII (75 g.) at 130° gives methyl acetate (7 g.) and ethyl acetate (1 g.). Similar results are obtained by using PrOH instead of McOII (Isbert) .-- (c) At 130° with NaOPr and excess of MeOH gives methyl acetate and a little propyl acctate -- (d) Heated with alcohol at 180° it is not affected, but if a very little NaOEt be present it is completely decomposed, yielding EtOAc. Similar results are got by using PrOH and NaOPr. Resacetic Acid C1.II 22O3 is found in all these cases as a resinous body, not volatile with steam. It forms brown amorphous salts, NaA', KA', and NH,A', sol. water (Isbert, A. 234, 167).

5. Sodium amalgam forms 8-oxy-butyric acid: CH, CO.CH CO.Et + H, CIÎ,.CH(OĤ).CH,.CO,Ét.

6. Phenyl-hydrazine in the cold forms, as with all ketones, a phenyl-hydrazide:

but at 100° this loses EtOH and becomes methyloxy-quinizine (q. v.) (Knorr, B. 17, 2032). Pseudocumyl-hydrazine produces the homologous hydrazide CH₂.C(N₂HC₆H₂Me₈).CH₂.CO.Et [78]. Longyellow needles (from alcohol), or thick prisms (from ether). V. sol. hot alcohol or ether, sl. sol. cold alcohol or benzoline. Very unstable, and at 130°-140° changes to oxy-tetra-methyl-quiniziner

(Haller, B. 18, 706).

7. Hydroxylamine forms, as with other kctones, the oxin: CH, C(NOH).CH, CO,Et, \$\text{\tikitext{\texit{\text{\text{\text{\text{\text{\texi{\texi{\texi\texi{\texi{\texi{\texi{\texi{\texi{\texi{\texi{\texi{\texi{\texi{\texi{\texi{\texi{ colourless crystals, [140°], sl. sol. water, alcohol or ether (Westenberger, B. 16, 2996).

8. Furning nitric acid yields stalie seid and OXIMIDO-ACETO-ACETIC ETHER (q. v.) (Propper, A.

222, 48).

9. Sulphuryl Chloride forms mono- or dichloro-aceto-acetic ether, according to the proportions used (Allihn, B. 11, exe7): CH, CO.CH, CO, Et + SO, Cl,

CH, CO.CHCI.CO, Et + SO, + HCI; CH, CO.CH, CO, Et + 2SO, CI, = CH,.CO.CCl,.CO,Et + 2SO, + 2HCl.

10. Bremine gives mono, di-, tri-, and per benzeue). -Hg(C,H,O,), Amorphous. Formed BROMC-ACETO-ACETO ETHERS (Wedel, A. 219 Ont 11. Chlorine forms only di-CHLORO-ACETO-

ATENIO ETHER (q. v.).

12. Phosphorus pentachloride forms the ohlorides of two Ohloro-Crogonic Acids (q. v.)

13. Dry prussic acid heated with aceto-acetic ther for 8 days at 100 forms a cyanhydrin, CH, C(OH)(CN).CH, CO, Et, which is converted by boiling dilute HCl into oxy-pyrotartaric acid

by boiling dilute from the control of the control o aceto-acotio ether forms Cyano-aceto-acetio ether (q. v.) CH₃-CO.CH(CN).CO₂Et (Haller a.

Held, C. R. 95, 235).

15. Ammonia, whether dry (Precht, B. 11, 1193), aqueous or alcoholic (Duisborg, A. 212, 171), produces the imide of aceto-acetio ether.

Aceto-acetic ether imide C₈H₁₁NO₂ [34°] (213° nneor.) at 760 mm., (154°) at 154 mm. S.G. 1014. S. (cono. NH₃Aq) 1.25. Dry NH₃ is greedily absorbed by aceto-acetic ether, the compound CII, C(OH)(NH,) CH, CO, Et being doubtless at first formed. The liquid soon separates into two layers, water and the imide of acete-acetio ether. The latter is purified by distillation (Collie, A. 226, 297). Properties.—
Colourless monoclinic prisms. V. sl. sol. water, v. sol. alcohol, ether, bonzone, CS2 and CHCl3. Moisture greatly lowers its melting-point. It is CH, C(NH).CH, CO, Et or CH, C(NH,): CH.CO. Et. Reactions.—(a) Aqueous HCl splits it up into NH, and aceto-acetic ether (Duisberg).—(b) Cold dilute NaOH has no action, but on warming it gives NH, HOEt, acctone and CO, —(c) Pb(OAc), HgCl, ZuSO, or FeCl, also splits it into acctoacetic ether and NH₃, the latter throwing down the metallic hydrate. AgNO, does not give any pp.-(d) Glacial acetic acid also regenerates aceto-acetic ether on boiling.—(e) Sodium amalgam gives \$\beta\$-oxy-autyrio acid.—(f) Vitrous fumes passed into alcoholic solution forms nitroso-aceto-acetio ether. A by-product Call, NaOa forms plates [170°].—(g) Ac.O at 160° forms an acetyl dorivativo, C₆H₁₆AoNO₂ [63°] (232°), which combines with bromine, forming C₆H₁₆AcBr₂NO₂ [140°] .- (h) Paraldehyde gives di-hydro-tri-methyl-pyridine di-carboxylio acid, which is also formed from accto-acctio ether, NH, and aldohyde.—(i) Et1 at 100° forms ethyl-aceto-acetio ether and a baso (c. 290°), possibly ethoxy-di-methyl-pyridino. Condensation-products: Cl_bH₁₉NO₆ [160°]. Present in the brown resin got when C₅H₁₁NO₂ is distilled under atmospherio pressure. Insolublo in alcohol and ether. Boiled with KOHAq it forms oxy-di-mctbylpyrldime carboxylic acid.

16. Aceto-acetic other methyl-imide, CH₃.C(NMe): CH₂.CO₂Et or CH₃.C(NHMe): CH.CO₂Et,

(133°) at 50 mm., (215°) at 760 mm., is formed in like manner from aceto-acetic ether and methylamine (Kuchert, B. 18, 618). With paraldohydo and H.SO, it gives a condensation-product, C,,H,O,N, which forms trimetric crystals with blue fluorescence [86°].

17. Diethylamine forms &-di-ethyl-amidocrotonic ether, CH, C(NEt2): CH.CO.Et,

iquid (160°-168°) at 20 mm.

18. Heated with antitus (1 mol.) at 120° it yields crystalline body, C₁₆H₁₁NO₂, which melts at 81° and is probably the antildo of a et-acetic acid CH₂(NPh).CH₂(O₂H. By dissolving this

substance in cold H₂SO₄, H₂O is eliminated with formation of (15, 1) oxy-(Py. 3) methyl-quinoline (Knorr, B. 16, 2593).

19. o-Phenylene-diamine forms: C.H.(N: CMe.OH, CO,Et), 20. o Tolylene diamine gives :

CH_s.C_sH_s<NH>CMo.CH_s.CO_sEt.

(Ladenburg, B. 12, 951; Witt, B. 19, 2977) 21. With aldchydcs (Claisen, B. 12, 345):

CII, CO.CH, CO, Et + RCOH = H₂O + CH₃.CO.C(CH.R).CO₂Et.

The bodies are mixed in molecular proportions, and HCl is passed in at 0°. Or the bodies may be heated with Ac.O. Examples (Matthews, C. J. 43, 200) :- (a) Isobutyric aldehyde gives C10H16O2 (219°-222°). Oil. Smells like peppermint. Combines with bromine. (b) Valeric aldely do gives $C_1H_1O_3$ (2377–241°). S.G. $\frac{15}{2}$ 9612. Oil. Smells of strawberries. (c) Chloral gives $C_1H_2O_3$ (151°–158°) at 25 mn. S.G. $\frac{15}{2}$ 13420. From obloral, acct-acetic ether, and Ao_2O_3 at 150°. (d) Furfural gives C₁₁H₁₂O₄ [62°], (188°-189°) at 30 mm. From furfural, acet-acetic ether, and Ac.O. Easily soluble in ohloroform, acetic acid, alcohol, and benzeno. Large doubly-refracting crystals (from light petroleum and ether).

22. Aceto-acetic ether (2 mols.) condenses with aldehyde-ammonia, forming di-hydro-tri-METHYL-PYRIDINE-DI-CARBOXYLIC ETHER (q. v.):

2CH3.CO.CII2.CO.Et + CII3.CH(OH)NH2= 3II O + C II NMe (CO Et) 2.

Since the product contains three methyls and two CO.Et groups, we may assume these to be identical with the same groups in the reacting bodies. And inasmuch as the product is not acted upon by nitrous acid gas and forms an ammonium iodide with MeI, it would seem to bo a tertiary base. Nevertheless, inasmuch as metbylamine and aldehydo give a similar body, the reaction may probably be represented thus:

(Hantsoh, A. 215, 74; B. 18, 2579). Other aldehydes in presence of NH, form similar derivatives of the pyridine scries (v. METHYL-PYRIDINE). Thus cinnamic aldehyde and ammonia forms dimethyl-styryl-di-hydro pyridine di-carboxylid ether, H₂C₃NMe₂(CH:CHPh)(CO₂Et)₂, [148] (Epstein, A. 231, 3).

23. With formamide and ZuCl, acoto-acetic ether gives di-motbyl-pyridino carboxylie ether (Canzeroni a. Spica, G. 14, 449). 9 24. With acctamide and AlCl, it forms CH, C(NAo).CH, CO, Et [65]. Needles. Converted

by KOH into the amide of aceto-acetic ether. 25. Mixing with acetamidine sydrochloride and dilute NaOH, and, after standing for some days, evaporating to dryness and extracting with alcohol, yields a di-methyl-oxy-pyrimidine, C.H.N.O [190°]. Needles. V. sol. water or alcohol, al. sol. ether or benzene. It is probably

CH. CMe OH. N:0(OH)

Propionamidine forms a homologue, methylethyl-cxy-pyrimidine [130°]. Its hydrochloride forms thick prisms, C₂H₁₀N₂OHCl [c. 243°]. V. e. sol. water, v. sol. alcohol—(C,H₁₀N₂OHCl)_PtCl. [236°]. Prisms. (Pinnsr, B. 17, 2520°; 18, 2847).

26. With urea in alcoholic solution it forms A-uramido-crotonic ether (Behrend, A. 229, 5): MeC(OH): CH.CO2Et+NH2.CO.NH2=

NH2.CO.NH.CMe: CH.CO.Et + H2O. The free β -tramido-crotonic acid, when liberated from its salts, changes at once into its anhydride,

NH.CO.NH.CMe: CH.CO

27. Thio-urea (40 g.), heated with aceto-acetic ether (40 g.) slowly to 150°, gives a compound C3H, N2OS (5 g.), which may be thio-methyltracil. It may be crystallised from water. It dissolves in alkalis and is reppd. by acids. Its melting point lies above 500°. Its aqueous solution gives with AgNO, an amorphous pp. of C.H.Ag.N.SO (Nencki a. Sieber, J. pr. [2] 25, 72). If a little HCl be added to an alcoholic solution of thio-urca and aceto-acetic other, unstable necdles are formed. These are converted by alcoholic potash into potassium thio-methylnracil, C,H,KN,SO (List, B. 19, 219).

28. Aceto-acctic ether (20 g.), phenyl-urca (10 g.), and ether (6 g.) at 150° react thus:

C₆H₁₆O₂ + C₁H₂N₂O = C₁₃H₁₆N₂O₃ + II₂O.

The product is an oil which is decomposed by alcoholic potash with formation of ammonia and aniline, and by boiling conc. HCl with formation of CO., alcohol, acetone, and plienyl-carbamic ether, PhNH.CO.Et. The reactions indicate that the body C₁₃H₁₃N₂O₃ has the constitution NPhH.CO.N: CMe.CH₂CO₂Et, or perhaps

, NH CMe, CH₂, CO₂Et

(Behrend, A. 233, 1).

29. Combines directly with di-phenyl-urea, in presence of a little other at 150°, forming an oil, C., H., N.O. This body is converted by alcoholic KOH into aniline and K.CO., and by acids into phenyl-carbamic ether and aniline. The body must be

PhNH.CO. NPh.C(OH)Mc.CH2.CO2Et. Similar addition products are probably first formed in the case of other ureas, but H₂O

PhnH.CO.NH.C(OH)Me.CH, CO, Et = H2O + PhNH.CO.N: CMe.CII, CO.Et.

30. p.Di-azo-toluene Chloride, acting on an alcoholic solution of aceto-acetic ether, forms yellow needles of p-toluene-azo-aceto-acetic ether [188e] C.H., Me.N., CH(CO.CH.). CO.2Et (Richter a. Minzer, B. 17 1929; v. zzo compounds).

31. Hydrazo-bengene at 100°-150° forms

HOEt and a crystalline base, C1.H14N2O (A. Müller, B. 19, 1771).

32. Copper accto-acetic ether is converted by COCL, into an anhydride of di-acctyl-acetone di-carboxylic ether, CO(CHAo.CO,Et), The anhydride may be:

"CH, C - 0 . C.CH, CO,Et.O-CO-C.CO,Et.

[80°]. Sol. glacial HOAc, H.SO., conc. HCl, bel zene, alcohol, crather. NH converts this hod into oxy-di-methyl-pyridine-di-carboxylic ether OH, C NH - C.CH,

CO,Et.Ö - CO - C.CO,Et. (Conrad a. Gnthzeit, B. 19, 22).

33. Aceto-acetic ether, heated with CS, and PbO at 100°, forms 'thio-carbonyl-aceto-acetic ether' CH₃.CO.C(CS).CO.Et [156°-162°]. Yellov needles (from alcohol) (Norton a. Oppenheim B. 10, 703).

34. S2Cl2 converts sodium aceto-acetic ether suspended in benzene into sulphido-aceto-acetic ether S(CHAc.CO,Et), [81°] (Buchka, B. 18,

35. With succinic acid it reacts thus: $C_{a}H_{10}O_{1} + C_{4}H_{a}O_{4} = C_{18}H_{12}O_{5} + 2H_{2}O_{4}$

The product is a crystalline acid [76°], which is the acid ether of a dibasic acid C.H.O. [200°] (Fittig, B. 18, 2526).

Condensation products from aceto-acetic ether. -1. By heat: Passed through a red-hot tube it forms Dehydro-AGETIC ACID (q. v.) and other products (Perkins, jun. C. J. 47, 240).—2. By hydrochloric acid: Dry HCl at 8° forms, in four wecks, acetic ether and 'carb-aceto-acetic ether' C₈H₁₀O₃ (290°-295° uncor.). S.G. 27 1·136. This liquid is slightly decomposed on distillation. It gives no colour with aqueous Fe.Cl, (Duisberg, A. 213, 179). Carb-aceto-acetic other is also formed when accto-acctic ether is heated with acetyl chloride at 120° (Wedel, A. 219, 116).-3. By sulphuric acid:

C_eH (OH)(CO₂H).CO₂C_eH,(CO₂H)(CO₂Et)

[622]. Got by leaving a mixture of aceto-acetic ether (1 pt.) and cold conc. II,SO, (21 pts.) for fonrteeu days and then pouring into water (Hantzsch, A. 222, 4). Needles resembling asbestos. Sl. sol. cold water, m. sol. hot water and ether, v. sol. alcohoff v. c. sol. chloroform. Decomposed by heat. It is acid to litmus and, when hot, has a spicy odour. Its formation may be represented thus:

 $4C_0H_{10}O_3 = C_{18}H_{22}O_0 + 3C_2H_3OII.$ Reactions.—(a) Boiling alkalis form carbonate aud acetate, acetone, and mesityl oxide. (b) On neutralising with alcoholic potash it decomposes

 $\begin{array}{c} C_{a}H_{1}(OH)(CO_{a}H_{1})CO.O.C_{a}H_{1}(CO_{a}H)CO_{a}Et + KOH = \\ C_{a}H_{1}(CO_{a}K) & C_{O} \\ \end{array} + C_{a}H_{1}(CO_{a}Et) & C_{O} \\ + 2H_{a}^{2}O_{a} \\ \end{array}$

the potassium salt of mesitene-lactone carboxylic acid being ppd. while its ether remains in solution.

Mesitene-lactone Carboxylic Acid CH, C:C(CO,H).CMe:CH.CO.O

[155°] (iso-dehydracetic acid). Fluffy orystals (from water). Sl. sol. colinwater, v. sol. het water. Monoclinic prisms (from alcohol). May be sublimed.

Reactions .- Distilled with lime it gives mesityl exide.

Saltso-KA'laq. - NaA'. - NH.A' - BaA'. MgA'z.-CuA',2aq.-Ag,Ha',.-Ag,HA',. Strong mesitene dicarbotylio acid, OH, C(OH)C:C(CO,H).CMe:CH.CO,H,

which forms a stable copper ealt Cu,H₂A' its alkaline galts madily eplit off CO₂:

O₂H₃(OH)(CO₂Na)₂ + Na₂OH =

O₄H₃(OH)(CO₂Na) + Na₂CO₃,

forming oxy-mesitene arboxylates.

These undergo a similar decomposition: $O_eH_e(OH)(CO_2Na) + NaOH = Na_2CO_3 + C_eH_eOH$ probably forming 'mesitenyl' alcohol, which at once changes to meeityl oxide. Baryta is the best alkali to uso in these decompositions.

Mesitene Lactone CH3.0: CH.CMc: CH.CO.0

[51.5°] (245° cor.). From iso-dehydracetic acid by distillation, or by heating with H₂SO₄ at 170°. Chittering tables. Very soluble in alcohol, ether, or water, sparingly so in CS₂. Tastes bitter and epicy. Nentral to litmus. Separated by K.CO₂ from aqueous solution. Gives with Br in CS₂ a bromo-dcrivative, C.H., BrCO, [105°]. Converted even by water into oxy-meeitene carboxylic acid. Hence it is a 8-lactone.

Oxy-mesitene Carboxylie Acid C.H.(OH)(CO.H) i.c.

CH, C(OH): CH. CMe: CII. CO. H.

From its lactone (mesitene-lactone) by boiling with water or, better, with baryta. A thick liquid, soluble in water. Its salts are amorphous, e.g. BaA'₂, CaA'₂. These salts on boiling form carbonate and mesityl oxide.

Mesitene-lactone Carboxylate of Ethyl

C₂H₂(CO₂Et)CO₂ i.e. CH₂.C:C(CO₂Et).CMe:CH.CO.O.

Formed, together with its potassium salt, by the action of alcoholic KOH on the product of condensation of aceto-acctic ether, as described above. Also from its potassium salt by EtI. Also got when aceto-acetic ether, saturated with HCl. is left at 0° for a month (Polonowska, B. 19, 2402). Oil. Reactions.—(a) Br in CS₂ forms C₈H₈Br(CO₂Et)CO₂[87°]. Needles (from alcohol). (b) NH, paceed into an alcoholic solution forms satiny plates of C.H.,(CO.Et)(CO.NH.)(ONH.) melting at [104°], giving off 2NH, + H.O, and changing to the lactone. Warm water or cold alcohol also change it to the lactone. Cold HClAq liberates CoH,(CO,Et)(CO,H)(OH), which may be extracted by ether. Small plates [76°]. Boiled with water, two-thirds are changed to lactone, and when the lactone is boiled with water, onethird becomes oxy-acid. It forms salts of the form O.H.,(OH)(CO.Et)(CO.H), viz. CuA'.aq, PbA'.aq. When the acid is boiled with alkali there is formed dehydracetic acid and its decomposition products, mesityl oxide, acetone, and CO2, and the following decomposition also occurs: CH,0:0(CO,Et).CMc:CH.CO.O+3H,0=

CH, CO, H+HORt+CH, (CO, H). CMe:CH.CO, H, resulting in homomesaconic acid.

Homomesaconic Acid CH₂(CO₂H).CMe: CH.CO₂H

Ethyl effer Et.A". (240°-242°).

Theory of the Condensation.—As acctone gives mesity! Oxide O₂H₁₀ or CH₂CO.CH:C(CH₄), so accto-acctic ether, if it is CH₄CO.CH₂CO₂Et, should give the dicarboxylio ether of mesityl oxide, C.H.O(CO.Et),, or the acid, C.H.O(CO.H), Instead of this, ot gives an isomeride of the latte C_sH₁(OH)(CO₂H)₂, so that the group CO.CH must have changed to C(OH):C. Assuming that thie group prc-exists in aceto-acetic ether, the condensation would be as follows:

 $\begin{array}{l} \text{CH}_{\bullet}\text{C}(\text{OH})\text{:C}(\text{CO}_{\bullet}\text{Et})\text{H} + \text{Ho}\text{.CMe}\text{:CH}_{\bullet}\text{CO}_{\bullet}\text{Et} \\ \text{CH}_{\bullet}\text{.C}(\text{OH})\text{:C}(\text{CO}_{\bullet}\text{Et})\text{.CMe}\text{:OH}\text{.CO}_{\bullet}\text{Et} + \text{H}_{\bullet}\text{O} \\ \text{CH}_{\bullet}\text{.O}\text{:C}(\text{CO}_{\bullet}\text{Et})\text{.CMe}\text{:CH}\text{.CO}_{\bullet}\text{O} + \text{Ho}\text{Et} + \text{H}_{\bullet}\text{O}. \end{array}$

Acetyl-aceto-acetic Ether

CH., CO.CHAc.CO.Et or CH., C(OAc); OH. CO.Et. di-accto-acetic ether. (200°-205°). S.G. 15 1064 (James); 12 1·101 (Elion). From aceto-acetic ether (65 g.), ether (50 g.) and sodium (9 g.), by adding a solution of AcCl (30 g.) in ether (50 g.) in the cold (J. W. James, A. 226, 210; C. J. 47, 1).

Properties.—Pleasant smelling liquid, slightly decomposed on distillation. Miscible with alcohol, other, and benzene, hardly soluble in water, but slowly decomposed by it into acetic acid and accto-acetic ether. Fe₂Cl, gives a raspberry-red colour, removed by SO2.

Salts.-Acctyl-accto-acctie acid is a strong acid, and can expel acetic acid from its salts. -CuA', 2aq: insoluble in water; [148°].—NaA' (Elion, R. 3, 255).-NiA',2aq.

Reactions .- 1. NaOEt decomposes it into EtOAc and sodium accto-acetic ether.

Benzcyl-aceto-acetic Ether

CH₂.CO.CB₂H.CO₂Et

From sodium aceto-acetic ether and benzoyl chloride alone (Bonné, A. 187, 1), or dieselved in ether (James, A. 226, 220; C. J. 47, 10).

· Properties.—A fairly strong acid, capable of

displacing acetic acid.

Salt.—CuA'₂ (from benzene) [180'-190']. Formed by shaking the ether with agneons cupric acctate. Sl. sol. alcohol or benzene, m. sol.

o-Nitro-benzcyl-aceto-acetic Ether .- Prepared as above, using nitre-benzeyl ohloride. It is an oil. Boiled with dilnts H2SO4 it forms o-nitro-phenyl methyl ketone and also o-nitrobenzoyl-acetone in smaller quantity. • Conc. KOH forms a salt CH₂CO.C(C_eH₄NO₂)K.CO₂Et (Gevekoht, A. 221, 323).

Constitution of Aceto-acetic Ether.

Some chemists adopt Frankland'e formula for accto-acctic ether, CH₂CO.CH₂OO₂Et; others prefer Geuther's formula, CH, C(OH):CH.CO.Et: while a third party, relying upon the recults of Bacyer's researches into the constitution of the derivatives of indigo, consider that both formula are equally correct, or rather that at the moment of reaction the first may change into the second. Against Frankland's formula it is argued

(1) That the copper salt ie blue or green whereas compounde in which copper is united to carbon (e.g. copper acetylide) are red or yellow. (2) That it does not account for the existence

of acctyl and benzoyl derivatives.
(3) The formation of hydro-quinone dicarboxylio ether, a body containing two hydroxyls, by the action of eodium on di-bromo-aceto-acetic derivative C.H., CO.C.H., NHBz [170°] (Higgin, C.J. 41, 188). Dilute HCl at 120° liberates the free bases, which are described as AMIDO-RENEO-

PRENONES (q. v.).
p.BENZOYL-PHENYL-CARBAMIO ETHER
C.H., CO.C.H., NII.CO.Et [183°]. Prepared by the action of chloroformic ether on p-amido-benzophenone (Dochner a. Weiss, B. 14, 1839; A. 210, 246). Plates. Sol. hoiling alcohol, boiling acctic acid, and chloroform, insol. cold Decomposed by boiling KOH. BENZOYL PHENYL CARBAMINE

C.H..CO.C.H..NC. Iso - cyano - benzophenone From p amido-henzophenone (10 g.), chloroform (8 g.), and alcoholic KOH (Doebner, A. 210, 246). Silky needles, volatile with steam. SI. sol. lict water, v. sol. alcohol. Split up by acids into formio acid and amide-scetophenone. BENZOYL - PHENYLENE - DIAMINE

PHENYLENE-DIAMINE. BENZOYL.PHENYL.DI.ETHYL.AMINE v.

DI-ETHYL-AMINO-BENZOPHENONE.

BENZOYL - PHENYL . DI . METHYL . AMINE v. DIMETHYL-AMIDO-DENZOPHENONE.

p.DI.BENZOYL.DI.PHENYL.THIO.UREA SC(NH.C₄II₁,CO.C₄II₃)... [166°]. Prepared by the action of CS₂ on an alcoholio solution of pramido-benzophenone in presence of a little KOH (Doebner a. Weiss, B. 14, 1839). Colourless plates. Scl. chloroform, sl. sol. hot alcohol, other, benzene and CS,; insol. water.

RENZOYL, PHENYL, METHANE is BUNZOYL.

PHENYL-CARBANIC FINER (q. r.).
BENZOYL - PIPER - PROPYL - ALKEIN v. Benzoyl-Oxypropyl-piperidine

BENZOYL-PROPANE . CARBOXYLIC ACID v. PHENYL PROPEL RETORN CARBOXYLIC ACID.

a-BENZOYL-PROPIONIC ACID C, III, O, i.e. C.H. CO.CHMe.CO.H. Phenyl ethyl ketone a-carboxylic acid. From the ether and cone. H2SO, the mixture being left for 3 weeks. It ie an oil and gives a reddish-brown colour with

Fe₂Cl. Alkalis produce phenyl ethyl ketone. Ethyl ether CH, CHBz.CO.Et. (227°) at 225 mm. Formed by the action of NaOKt and Mel on benzoyl-acetic other (q. v.). Aromatic smelling oil. Fe.Cl, gives no colour in its alcoholic solution. NaOEt forms the sodium derivative CH, CNuBz.CO.Et. Phosphorus pentachloride forms C.H. CCl:CMe.CO.Et (Perkin a. Calman, C. J. 49, 156).

B-benzoyl-propionic soid C.H. CO.CH. CH. CO.H. Phenyl ethyl ketone w-carboxylic acid. [116]; [114] (Bischoff, B. 19, 95).

Formation .- 1. By the reduction of benzoyl. acrylic seid .- 2. By the action of Al₂Cl₄ (1 pts. on a mixture of succinic anhydride (1 pt.) and benzene (10 pts.). The product is chaken with water, when the acid remains dissolved in the benzene, whence it is extracted by shaking with KOH and ppg. with HCl (Burcker, Bl. [2] 35, 17; A. Ch. [5] 26, 433; Peclimann, B. 15, 889). 8. By oxidising its aldehyde .- 4. Its chloride is formed by the action of Al₂Cl₄ upon a mixture of sneeinyl obleride (1 mol.) and benzene (1 mol.) (Claus, B. 20, 1375).—5. By heating benzeyl-isosnecinic acid above its melting-point, CO₄ heing evolved (Kues a. Paal, B. 18, 8325).

Properties. -- White prisms, v. sol. hot water.

Converted by potash-fusion into benzoic and propionic acids. Reduced by sodium amalgam to y-oxy-phenyl-bntyric acid (or its lactons). C,H,.CH(OH).CH,.CH,.CO,H.

Salts.-BaA', needles. - AgA': sl. sol.

Ethyl other. EtA'. [82]; white crystals, turned rol by heat.

Phenyl-hydrazide C.H., C(N.H1'h).CH, C11, C0, H. [65°]. White silky needles, v. sol. alkalis, neids, alcohol, and benzene, sl. sol. ether.

BENZOYL-PROPIONIC ALDEHYDE

C.H., CO.CH., CIL, CHO. (245). S.G. 9 1:005; 14 998. Prepared by the netion of water upon the compound of phenyl propyl ketone with CrO.Cl... Oil, sol. ether and chloroform. It readily reduces AgNO₃₁ but does not combine with NaHSO,. Sedimn-amalgam reduces it to syrupy C,H, CH(OH).CH, CH, CH, CH, OH (c. 200°) (Burcker, A. Ch. [5] 26, 469; C. R. 94, 220). BENZOYL-PROPIONIC-CARBOXYLIO AOID

v. Phrnyl ethyl retore di-carroxylio acin.

BENZOYL-PYROCATECHIN v. DI-OXY-BENZ-

BENZOYL PYRROL v. Prinon.

Pseudo-beazoyl-pyrrol v. Pynnyn phenyk KUTHER.

BENZOYL PYRUVIC ACID C 14 H O 4 i.e.

CaH3.CO.CH2.CO.CO2H. [156]. Obtained by saponification of the ethyl other. Yellowishwhite prisms (from hot benzene). Strong acid. Evolves CO, at its melting-point.

Ethyl ether A'Et: [43]. Prepared by adding 48 pts. of acetophenone to a cooled solution of \$2 pts. of sodium in 150 pte alcohol, and then 584 pts. of exalic other. The precipitate I solium compound is washed with other, dried, dissolved in iced water, and decomposed by CO, when the benzoyl-pyruvie other crystallises out; the yield is 78 p.c. of the theoretical. It crystallises from petroleum-ether in long prisms. V. sol. all ordinary selvents. Fo₂Cl_e gives a blood-red colonration. The aqueous alcoholic solution gives with copric acetate a light-green pp. of (C₀,11₁₁O₀)₂Cu, this crystallises from hot benzene or alcohol in long green needlee. By heating with dilute aqueous NaOH the other is split up into acetophenone, oxalic acid, and alcohol. By boiling its acetic acid solution with phenyl-hydrazine it is converted into di-phenylpyrazol-carboxylic ether C, HPhN(NPh).CO, Et. By cold alcoholic NH, it is converted into acetoplienome, oxamide, and alcohol (Leyer a. Claisen, B. 20, 2181). BENZOYL-QUINALDINE v. METHYL-QUIN-

OLTL PHENYL RETONE

BENZOYL-RESORCIN v. DI-OXY-BENZOPHE-

Di-bonzoyi-resercin v. Dioxy-Phenylene pr-PRENTL DIRETORE.

BENZOYL-SUCCINIC ACID. Phenyl ethyl ketone di-carborylic acid. Ethyl ether CO_Et CHBz.CH_CO_Et. (c. 265°) at 160 mm. From sodium-benzoyl-acetic ether and chloreacetic ether (Perkin, jun., C. J. 47, 274).

Properties.—Thick colourless oil. Its also-

helio solution gives a claret colonr with Fe Cl. Cone. H.SO, forms a yellow solution turned red by heat. NaOEt forms a solid sodium compound. Boiling baryts water decomposes it

into bensote and succinic acids. ... coming units H.SO, forms bensoyl-propionic acid.

Di bensoyl-succinic acid

CO.H.CHBz.CHBz.CO.H. Obtained by dissolving the ether in alcoholic KOH and treating with H.SO. (Perkin, jun., C. J. 47, 265). The acid dissolved in alcohol gives a dark-green pp. with Fe.Cl. Conc. H.SO, forms a yollow solution, turned orimson by heat.

Ethyl ether CO.Et.CHBz.CHBz.CO.Et. [180°]. Formed by adding iodins to a solution of sodium benzoyl-acetic ether in dry ether (Perkin, jun., C. J. 47, 262). Crystals (from alcohol); sl. sol. cold alcohol, v. sol. other. Cono. ILSO, forms a colourless solution turned red, olive-green, and finally bluish-red by heat. Sodimu ethylate forms OO_Et.CNaBz.CNaBz.CO_Et. In the alcoholic solution Fa, Cl, gives a red colour. Boiling dilute anlphuric acid (1:2) forms an acid probably diphenyl-furfurane dicarboxylic acid, C, 11, O, [238°], whence Ac.O forms an anhydride C,H₁₀O₄ [255"] (Baeyer a. Perkiu, B. 17, 62).

B. Benzoyl-isosuccinio acid v. PHENYL ETHYL

KETONE W-DI-CARBOXVIJC ACID.

BENZOYL SULPHIDE (C.H., CO), S. From BzCl and potassimo thiobenzoate (Engelhardt, Latschinoff, a. Malyscheff, Z. 1868, 357). Wary prisms, insol. water, v. sol. ether. Am-monia forms henzamide and ammonium thiobenzoate. Alcoholic KOH forms KOBz and KSBz. Alcoholic KSH forms KSBz.

Benzoyl disulphide (C.H.CO) S., Mol. w, 274. [1282]. Formed from C.H.CO.SH by atmospheric exidation of its solution in CS, (Cloez, A. 115, 27), or by treatment with iodine, Fo,CL, or HNO. Also, together with Bz,S, by warming BzCl with PbS and other (Moshing, J. 118, 301). When heated above its melting-1 point it turns violet. Prises or tubles (from CS.), st. sol. boiling ether and alcohol. Insol. water, NII,Aq, and KOHAq. Alcoholic KOH forms KOBz and KSBz. Alcoholic KHS forms

BENZOYL BULPHOCYANIDE CHICOSCN. S.G. 18 1/20. From BzCl and Pb(SCN), in the sold (Mignel, A. Ch. [5] 11, 300). Trangent liquid. Decomposed by boiling water into benkamide and COS. On long standing it deposits an Isomeride (160°), which is decomposed by water at 200° into NII, benzoic acid, and H.S.

BENZOYL - TARTARIC ACID C. TARTAUG

BENZOYL-TEREPHTHALIC ACID e. BENZO-PHENONE DICARBOXYLIC ACID.

BENZOYL THIOARSENITE C., H, Ass, O, i.e. As(SB2), (179°). From BaCl and AsiS, An amunonical solution gives with HgCl₂a pp. of . Hg(SB2), (Rayman, Pd. (2) 47, 896).

BENZOYL THIO CARBAMIC ACID

C,H,NSO, Methyl other Bz.NH.CO.S.Mo. From beuzoyl sulphocyanide and methyl abohol (Miguel, A. Ch. [5] 11, 330). Slender needles (from dilute alcohol). Sl. sol, water, v. sol, alcohol. Water in large excess at 100 forms BaNH, methyl alcohol, H₂S, and CO₂. Salt,—BaNNa.CO.SMe. From the ethoreal Balt.—Banna.Co.SMe.

Con and Boundary of from alcohol, BzCl, and minutes, with gradual addition of menny, Bt. [2] 34, 399; 35, 560).

KNON (Lossiner, 1971), 1980). Long needles; v. sl. sol, water, v. sol, alcohol. Boiling, KOHAq forms KOBz, KSCN, alcohol, CO, NH, and H.S. Heated alone it gives benzonitrila. CO. and mercaptan. Salt.-BzNK.CO.SEt: ncedles.

BENZOYL THIOCYANATE v. BENZOYL SULPHOCYANIDE.

BENZOYL-THIO-UREA v. THIO-UREA.

BENZOYL-THYMOL v. THYMOL.

BENZOYL-TOLUENE-SULPHAMIDE v. Tol-UENE SULPHONIO ACID.

BENZOYL-TOLUIDE v. TOLUIDINE BENZOYL-TOLUIDINE . IMIDE - CHLORIDE

v. w-Chloro-benzylidene-toluidene. BENZOYL. TOLYLENE - DIAMINE v. Tol-YLENE-DIAMINE

BENZOYL-TROPEÏNE v. TROPIN.

BENZOYL-UREA v. URRA.

BENZOYL-URITIC ACID v. PHENYL TOLYL KETONE DI-CARBOXYLIO ACID.

a-BENZOYL-VALERIC ETHER v. Propyl-

BENZOYL-ACETIO ETHER. BENZOYL-XYLENE v. PRENYL XYLYL RE-

BENZOYL-XYLIDE v. XYLIDINE.

(a) BENZ-PINACOLINE Callado i.e.

 $\begin{array}{c|c} C'H''-C,C''H''\\ \hline C'H''-C,-C''H''\\ \hline \end{array}$ (?) [204°].

Tetra phenyl-ethylene oxide.

Formation ... 1. Together with the (B) modification by boiling a 5 p.c. alcoholic solution of benzophenone with zine and HCI (Therner a. Zincke, B. 11, 65) .- 2. Together with bsnzpiuncone by heating an ulcabelic solution of benzophenone with zinc and H.SO, (Thörner a. Zineke, B. 11, 1396). 3. By the action of zinodust on an ethercal solution of acetyl chloride (I mid.) and benzophenone (I mol.). If the acctyl chloride is used in excess the (a)-benzpisceoline first formed is converted into the (B)-benz-pinacoline (Poal, B. 17, 911).-4, By the oxidation of tetra chenyl ethylene with chromic mixture (Behr. B. 5, 277).

Properties. Needles. Almost insoluble in

cold alcohol and in cold acctic acid.

Reactions. 1. By neetyl chloride, HCl or 11.80 at is converted into the (B)-modification, 2. By heating with soda lime it gives a hydroearbon [211] which is possibly tetraphenylethylene. 3. By CrO, and acetic acid it is exidired to beincophenoue.

(B)-Benz-pinacoline (C.H.), C.CO.C. If, [1790]

(T. a. Z.) : [182°] (Zagumenny).

Formation. By boiling a concentrated solution of benzophenone in alcohol with zino and tion of tetraophenone in account with zino and fill for 20 hogrs (Thorner a, Zincke, B. 10, 1473 a, 11, 65). 2. From benzpingcome and AcCl of BzCl (Linnemann, A, 133, 28). 3. From benzpingcome and dilute H SO, or HCl at 200°. It is even slowly formed by repeatedly recrystallising benzyinacone from hot alcollol (Z.). - I. From (a)-benzyinacone from hot alcollol (Z.). - I. From (a)-benzyinacoline by heating with AcCl. HCl, or H SO,

Preparation. HClAq is added to a saturated solution of benzyinacone in HOAc until a tur-

bidity appears. The mixture is boiled 45 minutes, with gradual addition of HClAq (Zagu.

ether. This indicates the presence of hydroxyl in di-bromo-aceto-acetic ether.

(4) The action of aminonia, and especially of di-ethylamine.

In favour of Frankland's formula may be noted the compounds with NaHSO, plicnyl hydrazino, and hydroxylamine.

The action of sodium upon aceto-acetic ether would be represented by Frankland's formula,

CH, CO.CH, CO, Et+Na=CH, CO, CNaH.CO, Et+H. Ethyl iodide converts the product into ethylaceto-acetic ethor:

CH, OO.CNaH.CO.Et+EtI=CH, CO.CKtH.CO.Et+NaI These two reactions may be repeated upon the

ethyl-aceto-acetic ether :

CH_CO.CEtII.CO_Et+Na=CH_.CO.CEtNa.CO_Et+H CH_CO.CEtNa.CO_Et+EtI=CH_.CO.CEt_.CO_Et+NaI. Adopting Geuther's formula, the four equations

become:

OH_O(OH):CH_CO_Et+Na=CH_,C(ONa):CH_CO_Et+H CH_,C(ONa):CH_CO_Et+EtI=CH_,C(ORt):CH_CO_Et+NaI CH_,C(ORt):CH_CO_Et+Na=CHI_,C(ORt):CH_,CO_Et+NaI CH_,C(ORt):CH_CO_Et+RtI=CH_,C(OEt):CEt,CO_Et+NaI It will be seen that the third and fourth equations are similar to the first and second on Frankland's hypothesis, but different in kind to the first and second if Geuther's hypothesis be accepted. Such a difference is not borno out by experiment. Thus if it be held that the action of sodium upon aceto-acetic ether depends upon its affinity for oxygen, the third equation presents a difficulty. And if we suppose that, owing to some intra-molecular change, the third equation ought to be written thus:

CH, C(OEt):CH.CO, Et+Na - CH, C(ONa):CEt.CO, Et+H then by the action of acctic acid on the product we ought to gct an ether CH3.C(OH):CEt.CO2Et isomeric, not identical, with ethyl-aceto-acetic cther, CH2.C(OEt):CH.CO2Et; but the two ethers are found to be identical (James, C. J. 47, 1). Inasmuch as the change of CH₃.C(OH):CEt.CO₂Et into CH3.C(OEt):CH.CO.Et would be contrary to all analogy, it is necessary, if we adopt Geuther's formula, to assume that the modo of formation of di-cthyl-aceto-acetic ether is something very different from that of ethylaceto-acetic ether. Again Geuther's formula would make methyl-ethyl-accto-acctie ether, OH .. C(OEt): CMe. CO. Et and cthyl-niethyl-acetoacetia other, CH3.C(OMe):CEt.CO2Et isomeric, yet this does not appear to be the case (James).

It may be said that there is some improbability in the assumption required by Frankland's formula, of direct union between sodium and carbon, but such a union is known to occur in sodium acetylide and sodium ethide, and it is very probable in many cases, such as sodiomalonic ether, sodium nitro-ethane, and sodic barbiturate. In order that hydrogen attached to carbon may be displaceable by nietals, it is necessary that very powerful chlorous groups should also be attached to the carbon, such as the nitroxyl in nitro-ethanc. One carbonyl, CO, is not sufficient to produce an acid, but two are. Thus the hydrogon in the group CO.CH2.CO is displaceable by metals, as in barbituric acid,

MH.CO. CH2 NH.CO/

These considerations account for the acidity of aceto-acetic ether, if we assume Frankland's formula CH,.CO.CH,.CO.OH.

Although the existence of acetyl-aceto-acetic ether favours Geuther's hypothesis, yet the fact that this body is a strong acid is wholly opposed to that view, and is very much better explained by the formula CH₁.CO.CH(CO.CH₁).CO.OEt, since if two carbonyls can make the group CH₂. acid, a fortiori three oarbonyls can have a simi-

The formation of ethyl and di-ethyl-acetone from ethyl-aceto-acetic ether and di-ethyl-acetoacetic ether respectively cannot be explained on Geuther's hypothosis.

If, therefore, we have to choose between one formula and the other, the balance of evidence

would indicate CH, CO.CH, CO, Et.

Methyl aceto-acetate $C_0H_0O_1$, i.e. $CH_2Ac.CO_2Me$ (170° cor.) S.G. 2 1.037 (Brandes, J.Z. 3, 25). From sodium and methyl acetate. Gives a cherryred colour with Fe₂Cl₃. Boiled with acids or strong bases it gives CO2, acetone, and MeOH.

Salts. - CH3. CO.CHNa. CO.Me. Sl. sol. ether. -Cu(C₅II₂O₃)₂2aq. Separates on adding cupric acctate and baryta water to the ether as pale

green crystals, insoluble in alcohol.

Isc - butyl acete - acstate CaH, O. CH.Ac.CO.C.H., (202°-206°) S.G. 2.979; 22.932,

From iso-butyl acetate and sodium.

Ise-amyl acete-acetate $C_0H_{14}O_3$ i.e. CH_Ac.CO_C,H₁₁ (223°) S.G. $\frac{12}{2}$. 954. From iso-auyl acetate and Na (Conrad, A. 186, 228). Converted by Cl into an oily di-chloro-derivative (Conrad, A. 186, 243) and by NH, into the imide of aceto-acetato of iso-amyl [190°-195°] (Collie, A. 226, 319).

ALKYLATED ACETO-ACETIC ETHERS. Sodium aceto-acetic other is converted by alkyl iodides into mono-alkyl aceto-acetic ethers, The sodium derivatives CII,.CO.CXH.CO.Et. of thest are in like manner converted by alkyl iodides into di-alkyl-aceto-acetio ethers, CHa.CO.CXY.CO.Et. Such ethers are of great service in organic syntheses, for they are split up by weak alkalis into carbonic acid and mono- or di-alkylacetones: CH₃.CO.CXY.CO.Et + 2KOH = CH₃.CO.CXYH + HOFt + K.CO₃, and by strong potash into mono- or di-alkyl-acetic acid and acctio acid: CH_.CO.CXY.CO_Et + 2KOH = CH3.CO2K + HCXY.CO2K + HOEt.

In practice the ketonic and acetic decompositions both occur, at the same time, but the acctic decomposition increases with the concentration of the alkali (Wislicenus, A. 206, 808).

Preparation. - The alkyl-aceto-acetic ethers are prepared by dissolving the calculated quantity of sodium in 10 times its weight of absolute alcohol, cooling, adding the aceto-acetic ether and then the alkyl iodido until the liquid, which may be warmed, if necessary, is neutral to litmus. The greater part of the alcohol is then distilled off and water is added. This dissolves the NaI and the new ether rises as an oil and is fractionated (Conrad a. Limpach, A. 192, 154).

A. WITH ONE MONOVALENT RADICLE.

Methyl - aceto - acetic Acid C,H,O, CH3.CO.CHMe.CO2H. A thick liquid which splits up on warming into CO, and methylethyl ketone (Coresolc, B. 15, 1874). Its barium salt is soluble and gives a violet colour with Fe.Ol. Nitrous acid forms iso-nitroso-methyl-ethyl-ketone.

OH, CO.CHMe.CO.Me Mathy

(1771 cor.) S.G. \$ 1-020 (Brandes, Z. 1866, 458). From sodium aceto-acetate of methyl and Mel. Smells like mint. Gives a violet-red

colour with Fe,Cl., Ethyl Ethes CH, CO.CHMe.CO.Et (186.8° acor.) (Geuther, Z. 1866, 5). S. S. 21.009. Gives

a deep blue colour with Fe Cla.

Reactions .- 1. Sodium amalgam gives an oxy-valerio acid CH,CH(OH).CHMo.CO,H .- 2. Potash forms mothyl-ethyl ketone, alcohol and Potash forms mothyl-ethyl ketone, alcohol and K₂CO₃.—3. PCl₃ gives ohloro-methyl-crotonic acid [69.5°] (206°) (Rücker), ehloro-methyl-aceto-acetic ether, C₂H₁₁ClO₃ (180°) S.G. ¹³ 1·093, smelling of peppermint, and di-chloro-methyl-aceto-acetic other (210°-220°) S.G. ¹³ 1·225 (Isbert, A. 234, 188).—4. Sodium and cyanogen chloride form, 23 anomethyl-aceto-acetic ether CAOMeOy.CO₂Et (c. 93°) at 20 mm. S. G. 82 a00 Lt is a colourless liquid insol water and 22 996. It is a colourless liquid insol. water and alkalis (Hold, C. R. 95, 522; Bl. [2] 41, 330).

Acetyl derivative C.H.10, i.c. CMeAc,CO.Et. Methyl-diacetyl-acetic ether (205°-220°). From methyl-aceto-acetio ether in ethereal solution and AcCl (James, A. 226, 219, C. J. 47, 1). Sl. sol. water. Colonred raspberry red by Fe₂Cl₈. Does not pp. cupric acetate, oven on addition of

dilute NaOH.

Par France

Ethyl-aceto-acetio acid.

Methylether CH4.CO.CEtH.CO.Me (189.7° cor.) S. G. 14 995 (Brandes, Z. 1866, 457), Fe₂Ci₄ gives deep violet colour. Conc. NH, forms an oil C,II,2NO2, the imide of ethyl-acete-acetate of methyl insoluble in water, and also silky needles [83°] of a soluble amide (probably C₆H₁₁NO₂ see below) (Brandes, Z. 1866, 457).

Ethyl ether C, H, O, i.e. CACEHI.CO.Et (198° cor.) S.G. 12° 998 (G.); 14° 983 (F. D.) (Gouther, Ar. Ph. [2] 116, 97; Frankland a. Duppa, C. J. [2] 4, 396; Wislicenus, A. (86, 187).

Preparation .- Aceto-acetic ether is dissolved in benzone and four-fifths of the calculated quantity of sodium added, then EtI, and the product rectified. The aceto-acetic ether recovered is treated with the remaining fifth of the sodium. Yield 70 per cent. (Wedel, 4. 219, 100).

Properties .- An oil. Coloured blue by Fe Cla Reactions .- 1. Reduced by sodium amalgam to an oxy-hoxoio acid, CII₃.CH(OH).C1IEt.CO₂H.

2. Boiled with barifa or weak alcoholie KOH, it gives methyl propyl ketone.-3. Boiled with conc. alcoholic KOH, or heated with dry NaOEt, it gives n-butyric acid and acetic acid, or their ethers .-4. Treated with NaOEt and cyanogen chloride it forms cyano-ethyl-accto-acctic ether, CH, CO.CEtCy.CO.Et (c. 105°) at 20 mm. S.G. 22 .976. A colourless liquid with agreeable odour. Insol. in water or alkaline solutions, misciblo with alcohol or ether (Held, C. R. 98, 522, Bl. [2] 41, 330) .- 5. Bromine acting on an ethereal solution fornesomono- di- and tri- Bnomo-ETHYL-ACETO-ACETIO ETHEN (q. v.) .- 6. PCl, gives monoand di- CHLORO-ETHYL-ACETO-ACETIC ETHER, and only one ohloro-ethyl-crotonio acid (q. v.) (Isbert, A. 234, 183) .- 7. Benzoic aldehyde and HCl form some benzylidene-ethyl-acetoacetio ether or oinnamoyl-ethyl-acetic ether, Ph.CH:CH.CO.CHEt.CO.Et (210°) at 22 mm. Converted by NaOEt and Etl into cinnamo di-diethyl acetio ether .- 8. Cono. aqueous ammonia forms two amides, one soluble in water C. H. NO. the other insoluble, C.H., NO. They are formed

in equi-molecular quantities; the oily insoluble amide crystallises when cooled. On distilling the soluble amide does not pass over with steam.

Insoluble amide C.H., NO, imide of ethylaceto-acetio ether CH., C(NH). CHEt.CO_Et or CH., C(NH). CEt.CO_Et [59.5°]. Monoclinic tablets (from alcohol), smolling of peppermiut. Decomposed by water, or dilute acids, into NH, and ethyl-aceto-acetic ether (Geuther, Z. 1871,

247).

Solublo amide CoH, 1NO2 i.e. CAcHEt.CONH, [90]. Needles (from water, alcohol, or ether). May be sublimed. May be obtained from the preceding body by heating with water at 135°. On dry distillation it gives NH3, CO2 and methyl propyl ketone. The latter body is also formed by heating it with water at 200°, with boiling aqueous HCl, with CaCl., ZnCl., P2O, or PCl. (Isbert takes it to be di-ethyl ketone). Heated with dry KOH at 100° it forms butyrio and acctio

acids (Isbert, A. 234, 170).

Salts. - CH3 CO.CNaEt.CO.Et. Formed by adding sodium to a solution of ethyl-aceto-acctio ether in dry ether or benzene (3 or 4 vols.) (J. W. James, C. J. 47, 1). Also by shaking an ethercal solution of the ether with perfectly dry NaOH (Elion, R. 3, 234). It is amorphous. V. sol. ether. A little water added to its ethereal solution forms a pp. of CH3.CO.CNaEt.CO.Et aq, insol, ether or benzeue, but sol, water or alcohol. Acetic acid re-converts the sodium salt into ethyl-aceto-acetic ether (v. constitution of Aceto-ACETIC ETHER).

Ethyl accto-acctic ethor forms no copper compound. This is thought to favour the for-

mula CH3.C(OEt): CH.CO.Et.

Iso amyl ether CH3.CO.CHEt.CO.C.H11 (233°-236°) S.G. 3°., 937 gives no colour with Fc.Cl₈ (Conrad, A. 186, 228).

CH3. CO. CAcEt. CO2Et. Acetyl derivative Ethyl-di-acetyl-acetic ether (c. 230'); (144°-150°) at 50 mm. S.G. 15 1034. From CH, CO.CNaEt.CO.Et and AcCl (Elion, R. 3, 265). Liquid. Iusol. KOllAq. Gives no colour Alcoholic NH, converts it into with Fe₂Cl₆. acctamide and CH, CO.CHEt.CO.Et.

Allyl-acto-acetic C, H, O, i.e. Ether CH3.CO.CH(C3H3).CO.Et (206°) (Zeidler, A. 187, 33) (214° cor.) at 720 mm. (Perkin, C.J. 45, 540). S.G. 37., 982 (Z.); \(\frac{15}{15}\) 993; \(\frac{25}{15}\) 985 (P.). From sodium aceto-acetic ether and allyl iodide (Z.; Wolff, A. 201, 46). From aceto-acetic ether, allyl iodide, and ziuc, di-allyl-aceto-acotio other being also formed (O. Hofmann, A. 201, 77).

Reactions.—1. Fe.Cl. gives a crimson colour.
2. Boiling alcoholic KOH forms CO, and allylacetone. -3. Dry NaOEt at 150°-160° gives ethyl ncetate and allyl-acctate. - 4. Sodium amalgam forms an oxy-heptenoio acie,

CH₃.C(OH)H.CH(C₃H₃)CO₂H. Prepyl-aceto-acetio Ether C₃H O.H.O. CH, CO.CHPr.CO.Et (209°) S.G. 2 981. From aceto-acetic ether (153g.) by adding first a solution of sodinm (27g.) in dry aloghol (270g.) and then PrI (206g.) (Burton, Am. S. 385). Decomposed by aqueous KOH Into CO2, alcohol, and methyl butyl ketono.

Iso-propyl-aceto-acetio Ether C,H1.O, CH, CO.CPrH.CO.Et (201°) at 758 mm. S.G. 2880. From sodium aceto-acetic ether and isopropyl iodide (Frankland a. Dnppa, A. 145, 78)

Coloured pale reddish-violet by Fe₂Cl₆ (Demarcay, Bl. 27, 224).

130-butyl-aceto-acets Ether C_{0.}H_{1.0}O₄ i.e. Pr.OH, CHAc.CO, Et (218°) S.G. 173 951. From sodium aceto-acetic ether and iso-butyl iodide (Rohn, A. 190, 806). Decomposed by baryta giving methyl iso-amyl ketone and iso-butylacetic (hexoic) acid.

Heptyl-aceto-acstic Ether C13H21O2 i.e. CH, OO.CH(C, H₁₃).CO₂Et (272°) S.G. 177 ·9324. From sodium acoto-acetio other and hoptyl iodide (Jourdan, A. 200, 105). Colourless oil. Decomposed by dilute alkalis into methyl octyl ketono and CO2; and by cono. alkalis into acetic and n-ennoic acids.

Secondary Heptyl-aceto-acetic Ether (250°-260°). Prepared eimilarly from secondary heptyl

iodido (Venabls, B. 13, 1651).

Octyl-aceto-acetic Ether C₁₁H₂₆O₃ i. c. CH₃CO.C(C₃H₁₇)H.CO₂Et (281°) S.G. 185 1954. From octyl iodide and sodium aceto-acetic ether (Guthzeit, A. 204, 1). Decomposed by alcoholic KOH into methyl ennyl ketone and decoio

Benzyl-aceto-acstio Ether C₁₃H₁₈O₃ i.e. CH₃.CO.CH(CH₂Ph)CO₂Et (276°) S.G. $\frac{15^{\circ}5}{16^{\circ}3}$ 1·036. From sodium aceto-acetic other and benzyl chloride (Ehrlich, B. 7, 690; A. 187, 12; Conrad, B. 11,1056). Sodium amalgam gives exo-oxy-phenylvaleric acid CH, CH(OH).CH(CH, Ph).CO, Et. Alcoholio KOH forms phenyl ethyl methyl ketone.

B. WITH TWO DI-VALENT RADICLES:

Ethylene-aceto-acetio Acid. CH, CO.C(C,H,).CO,H. From the ether by saponification. Liquid. Decomposed by heat or by dilute acide into tri-methylene methyl

CH₂.CO.CH
$$\stackrel{\text{CH}_2}{\underset{\text{CH}_2}{\mid}}$$
 and CO₂.

Ethyl ether.—EtA' (193°-195°). From acetoacetic ether (26g.) by adding a solution of sodium (5g.) in alcohol followed by ethylene bromide (38g.) The liquid is boiled for eight hours, filtered, and distilled. The residue is boiled for twelve hours longer with a solution of sodium (5g.) in alcohol, evaporated, and treated with water. The ether is extracted by other and dried over K₂CO₃(W. II. Perkin, jun., C. J. 47, 934; B. 16, 2136; 19, 1247). It reacts with phenylhydrazine, forming an oil.

Ethylidene-actio-acstic Ether.

CH, CH: CAc.CO.Et (210°-212°) S.G. 15 1.023 By passing HCl into aldchyde (1 pt.) mixed with acoto-acetic ether (3 pts.) (L. Claisen a. F. H. Matthews, A. 218, 172; Claisen, B. 14, 345).

Pungent ethercal oil. Miscible with H'SO. Reactions .- 1. Hot potash decomposes it, forming aldehyde .- 2. Combines with bromine.

Tri-chlore-ethylidene-acete-acetic Ether. CCl.CH : CAc.CO.Et. S.G. 15 1.342 From chleral, aceto-acetic ether and Ac2O at 160°. (Claisen a. Matthewe, A. 218, 175). Thick oil. Decomposed by heat. Propylene-aceto-acetic Acid.

·CH, CH CAo.CO2H From the ether by saponification. Forms an amorphous silver salt, AgA'.

Ethyl-ether (210°-215°) at 720 mm. acetic ether (26g.) is heated with sodium (46g.), diesolved in dry alcohol and propylene bromide (40g.) at 100°. After two days the tubes are opened and a fresh quantity of alcoholic NaOEt (from 4 6g. eodium) is added, and the tubes hoated again at 100° (Perkin, jun., B. 17, 1448).

Tri-methylene Bromide acts on aceto. acetic ether in presence of NaOEt, but the product C₀H₁₄O₃ (V.D. 6.21) is not tri-methyleneaccto-acetic ether, for its boiling point (223°) and molecular magnetic rotation, 10.195, are both too high, and it does not react with phenylhydrazine. It is, however, the ether of a crye. tallino acid which splits up on distillatiou into CO₂ and C₆H₁₀O, and on boiling with water into CO₂ and acctyl-butyl alcohol. The acid is probably C(CO₂H):CMe

CH₂.CH₂.O (Perkin, jun., B. 16, 208, 1789; 19, 1247, 2557).

Isc-butylidsne-aceto-acetic Ether (CH₃) CH.CH:CAc.CO Et(219°-222°) From isobutyric aldehydo, aceto-acetio ether

and HCl (Claisen a. Matthows, A. 218, 174). Liquid smelling of poppermint. Iec-amylidens-aceto-acetic Ether.

(CH₁), CH.CH₂, CH:CAc.CO.Et (237°-241°) S.G. ¹⁵ 961. From valeric aldehyde, aceto-acetic other and HCl (Claison a. Matthowe, A. 218, 174).

Bsnzylidsns-acste-acstic Ether

Ph.CH: CAc.CO2Et (a-accto-cinnamic ether), [60°] (180°-182°) at 17 mm. (295°-297°) at 760 mm. From aceto-acctic ether, benzoic aldehyde and gaseous HCl at 0°. (Claisen a. Matthews, A. 218, 477) 4 or 6 sided tables (from alcohol); trimetric, a:b:c=447:1:962. Colourless oil, solidifying very slowly. V. sol. chloroform, m. sol. cold alcohol, other, glacial acctic acid or CS., v. sl. sol. benzoline. Insoluble in aqueous KOH. H2SO, forms a bright yellow solution which, on warming, becomes very dark red. On pouring this solution into water a white pp. is formed, and on adding NaOH this diesolves. forming a violet solution.

Reactions.—Bromino in ether forme a di-bromide [97°]. This forms short needles (from

benzoline).

Theory of the Process.-Benzoic aldehyde probably first combines with HCl forming Ph.CH(OII)Cl, and this reacts with accto-acetic ether thus:

Ph.CliCl(OII) + CII.Ac.CO2Et == II2O + Ph.CHCl.CHAc.CO2Et.

Two compounds of this formula may be isolated before distillation, one forme prisme [41°], the other small rhombohedra or chiclinic tables [72°] (both from henzoline). They are both unstable, giving off HCl. One of them has probably the formula Ph.CHCl.CHAc.CO2Et and decomposes into HCl and Ph.CH:CAc.CO,Et, which recombines with H& terming the other Ph.CH2.CClAc.CO2Et. On dietillation both probably give HCl and benzylidene-aceto-acetic ether.

Benzylidene-ethyl-aceto-acetic Ether

Ph.CH:CH.CO.CHEt.CO.Et (205°-220°) at 22 mm. (Cinnamoyl-ethyl-acetic ether). From benzoio aldehyde, ethyl-acetoacetic ether, and HOL. Yield small (Claisen a. Matthews, A. 218, 184).

Benzylidene-di-ethyl-aceto-acetic Ether

Ph.CH.CH.CO.OEt.CO.Et. [1010-1097]. Formation.—(1) From the above, NaOEt, and Etl.—(2) From benzoic aldehyde, di-ethyl-aceto-acetio other, and HCl (C.M.). Triclinic prisms (from benzoline). V. sol. ether or chloroform, m. sol. oold alcohol or benzoline. Dibromide [55°].

Furfural-aceto-acetic Ether

(C,H,O)"CAc.CO,Et [62.5°]. (189°) at 30 mm. From furfur-aldehyde, aceto-acetio ether, and Ac₂O at 160°. (Claisen a. Matthews, A. 218, 176.) Trimetric crystals, a: b: o = 439: 1: 465. V. sol. alcohol, glacial acetic acid, chloroform, and benzone. M. sol. ether, sl. sol. benzoline.
C. With two Monovalent Radicles.

Di-msthyl-aceto-acetic Acid

 $C_6H_{10}O_3$ i.e. CH_3 .CO.CMe₂.CO₂H. From the ether by dissolving in cold dilute (21 per cent.) aqueous KOH, setting aside for a day or two, then acidifying with H2SO4, extracting with other, evaporating the ether, and triturating with BaCo₂. The crystalline barium salt, BaA₂, is decomposed by diluto H₂SO₄ (Ceresole, B. 15, 1871). Very hygroscopic crystals, which slowly split up into CO, and methyl iso-propyl ketone. The barium salt gives a brown colour or pp. with Fe₂Cl_c. It reduces boiling silver nitrate.

Ethyl Ether

C₈H₁₁O₃ i.e. CH₃.CO.CMe₂.CO₂Et (184°) S.G. ¹²·991. From sodium methyl-acetoacetic ether and MoI (Frankland a. Duppa, A.
138, 328). Potash or baryta splits it up into
alcohol, CO₂, and methyl iso-propyl ketono.

Mathyl-ethyleasto-acetic Ether

CH₂.CO.CMeEt.CO₂Et (196° nncor.) (J.) (201° i. V.) (Wislicenus, A. 219, 308). S.G. 33. 947. From sodium ethylaceto-acetic ether and MeI (Saur, A. 188, 257); or sodium mothyl-acete-acetic ether and Etl (J. W. James, A. 226, 209; C. J. 47, 1). Oil. Fe₂Cl₆ gives a violet colour. Distilled with dry NaOEt it gives ethyl acctate and ethyl methylethyl-acctate (or valerate).

Methyl-allyl-aceto-acetic Ether

C₁₆H₁₆O₃ i.e. CH₃CO.CMc(C₃H₅)CO₂Et (c. 209°-211'). From allyl-aceto-acetic ether, MeI, and NaOEt (James, C. J. 47, 3). Pleasantsmelling oil, miscible with alcohol, other, or benzenc. Fe₂Cl₆ gives no colour. The same body may be got from methyl-aceto-acetic ether, allyl iodide, and NaOEt.

Methyl-propyl-acsto-acetic Ethsr C₁₆H₁₈O₃ i.e. CH₃.CO.CMePr.CO₂Et (214°) (L.K.); (216°) (J.). S.G. ¹² 959 (L.K.); ¹⁷ 9575 (J.F. From methyl-aceto-acetic ether, NaOEt, and PrI (Liebermann a. Kleemann, B. 17, 918) or from propyl-aceto-acetic ether, NaOEt, and MsI (E. J. Jones, A. 226, 287).

Di-ethyl-aceto-acetic acid GH, CO.C(C,H,), CO,H.
Thick colourless liquid. Sl. sol. water.

Preparation.—Di-ethyl-acet-acetic ether is left in the cold for several weeks with 10 p.c. aqusous KOH. After removing the unaltered ether, the product is acidified and extracted with sther, and the acid purified by conversion into | Sl. sol. cold water. Salt: AgA'.

the barium salt, acidifying the latter, and again

extracting with ether.

Reactions.—It decomposes very slowly in the cold, but on heating to 60° it rapidly evolves CO2 forming di-othyl-acetone. The latter body is also formed by distilling the barium salt.

Salts.—ANa; easily soluble white micro-

scopic crystals.—A'_Ba 2aq; transparent prisms, rotates on water (Ceresole, B. 16, 830).

Ethyl ether O₁₀H₁₈O₃ i.e. CH₃CO.CEt₂CO₂Et (218°). S.G. 22 974. From sodium ethyl-acetoacetic ether and EtI (Frankland a. Duppa, A. 138, 211; James, A. 226, 205). From Cl.CO, Et. Na, and Etl (Geuther a. Matthey, J. pr. [2] 6,

160). Reactions .- 1. With hot aqueous baryta it gives di-ethyl-acetone .- 2. Distilled with dry NaOEt it gives di-ethyl-acetic (hoxoic) ether, acetic acid, and sodic di-cthyl-acetato .- 3. PCl. gives mono- and di-chloro-di-ethyl-aceto-acetio ether and chloro-cthyl-crotonio ether (James, A. 231, 235) .- 4: With benzoic aldehyde and HCl gas it forms some C.H.CH:CH.CO.CEt.CO.Et, cinnamoyl-diethyl-aceticether. Crystals, [102°], (200°-205°) at 3 mm. Easily soluble in ether and chiloroform, slightly in cold alcohol and in light petroleum (Matthews, C. J. 43, 205). Bromine in chloroform forus a di-bromide, Prisms v. sol. alcohol and light petroleum. Di-allyl-aceto-acstic Ethsr

 $C_{12}H_{18}O_3$ i.e. CH_{3} : $CO.(C_3H_5)_2CO.Et$ (240°). S.G. $\frac{2}{17}$, 948. From sodium allyl-acetoacetic ether and allyl bromido (Wolff, A. 201, 45). From aceto-acetic ether, allyl iodide, and zino (O. Hofmann, A. 201, 77). Colourless oil, with faint peculiar odour. Insol. water, sol. alcohol. ether, or benzene. Boiling cone. KOHAq forms

di-allyl-acetone, or methyl heptinyl ketone, and di-allyl-acetic acid.

Di-propyl-aceto-acetio Ether C12H22O2 i.e. CH, CO.CPr. CO.Et (236°). S.G. 2 9585. From sodium propyl-aceto-acetic ether and PrI (Burton, Am. 3, 386). Alkalis split it up, giving dipropyl-aceto-acetic ether and di-propyl-acetone or methyl heptyl ketone.

Di-isobutyl-aceto-acetic Ether O₁₄H₂₆O₂ i.e. (PrCH.)₂ÖAc.CO₂Et (250°-253°). S.G. ¹⁰2947. From sodium isobutyl-aceto-acetic ether and isobutyl iodide (Mixter, B. 7, 500).

Di-n-heptyl-aceto-acetio Ether

O₂₀H₃₈O₃ i.c. CH₈CO.C(C₇H₁₅)₂CO₂Et

(332°) S.G. 17.3 891. Formed together with diheptyl-acetio other and methyl octyl ketono by heating sodium heptyl-aceto-acotic ether with heptyl iodide and dry alcohol for a long tims (Jourdan, A. 200, 112). Decomposed by dilute alkalis into CO₂ and methyl pentadecyl ketone, and by concentrated allelis into acetic and diheptyl-acetic (hexadecoic) acids.

Di-octyl-acste-acstic Ether

C22H42O3 i.e. CH3.CO.C(C6H17)2.CO2Et (264°) at 90 mm. (340°-342°) at 760 mm. From octyl-aceto-acetic ether, NaOE, and octyl iodide (Guthzeit, A. 204, 9). Decomposed by alkalis into di-octyl-acetone (methyl heptadecyl ketone) and di-octyl-acetio (heptadecoic) acid.

Henryl-methyl-aceto-acetic Acid C₁₁H₁₄O₃ i.e. CH₂,CO.CMc(CH₂Ph).CO₂H [34°] (275°). From the ether by saponification

Ethyl ether-Eth' (287°). S.G. 33 1046. Prepared by action of MeI on a mixture of benzyl-aceto-acetle acid and sodium ethylate. (Conrad, B. 11, 1055).

Benzyl ether-PhCH, A' (53°?). Mothyl hy-

dro-cinnamein. Liquid.

6.72 :-96

> Benzyl-ethyl-aceto-acetic Ether CH₃.CO.CEt(CH₂Ph).CO₂Et Colourless liquid.

Di-benzyl-aceto-acetic Ether

CH, CO.C(CH,Ph), CO.Et.

From sodium benzyl-aceto-acetic ether and benzyl chloride (Ehrlich, A. 187, 21). non-volatile liquid.

OTHER DERIVATIVES of aceto-acetic acid will be described as acetyl derivatives, e.g. Acetyl. GLUTARIC RTHER, ACETYL-SUCCINIC ETHER, &c. Sec. also Oxy-ACETO-ACETIC ETHER, OXY-DI-ETHYL-ACETO-ACETIC ETHER, OXY-DI-METHYL-ACETO-ACETIC

For analogous acids see Propionyl-Propionic ACID, VALERYL-VALERIC ACID.

ACETO-BENZOYL-BENZOIC ANHYDRIDE v. BENZOYL-BENZOIC ACETIC ANHYDRIDE.

ACETO-REOMO-ACETIC ETHER v. Bromo-ACETO-ACETIC ETHER

ACETO-BROMO-AMIDE v. ACETAMIDE. ACETO-BUTYRIC ACID v. ACETYL-BUTYRIC

ACETO-CHLORO-AMIDE v. ACETAMIDE. ACETO-CHLORHYDRIN v. GLYCERIN.

ACETO-CHLORHYDROSE: C11II 19ClO, i.e. CaH, Ao, O.Cl. Formed by treating I mol. an. hydrous glueose with 5 mol. AcCl, and purified by solution in chloroform, agitation with sodium carbonate, and evaporation. - Semifluid; sometimes crystalline. Dextro-gyrate. Bitter. Insol. in water, slightly sol. in CS, easily in alcohol, ether and chloroform. Distils in a vacuum, partly undecomposed. Gives up all its chlorino to alcoholic silver nitrate. Reduces Fehling's colution. Reconverted into glucose by heating with water (Colley, C. R., 70, 401). H. W. ACETO-CINNAMONE v. BENZYLIDENE-ACE-

ACETO-COUMARIC ACID v. COUMERIC ACID. ACETO-CURCUMIN v. CURCUMIN.

ACETO-ETHYL NITRATE C2H4O, 2C2H4NO (84°-86°) S.G. 19 1015. Formed by dry distillation of potassium ethyl-sulphate with potassium nitrate. Liquid, having a sweet taste and aromatic odour. Explodes violently when heated above its boiling point. Not miscible with water. Resolved by heating with potash-lye into aldehyde and nitric acid (Nadler, A. 116, 173). H. W.

ACETO-ETHYL-SUCCINIC ACID v. ACETYL-ETHYL-SUCCINIO ACID.

ACETO-ETHYL-TRIENONE v. ETHYL-THI-ENYL METHYL KETONE

ACETO-GLYCEROLS v. GLYCERIN.

ACETO-GUANAMINE v. GUANIDINE. ACETO-TETRA-METHYLENE v. TETRAME-

TUYLENE METHYL KETONE.

ACETO-MEDHYL-THIENONE v. METHYL-

PHIENYL METHYL KETONE. ACETONAMINES.

Di-Acetonamine

ducting tube terminating just above the liquid; the resulting mixture of acetone vapour and ammonia is passed through a tube heated to 100° and then through a condensing tube; the distillate is neutralised with sulphuric acid diluted with an equal volume of water, and, after removing the ammonium sulphate which oryetal-lises out, and distilling off unaltered acctone, the liquid is evaporated to dryness and the residue exhausted with boiling alcohol. Diacetonamine sulphate then crystallises out on cooling, and may be purified by recrystallisation from alcohol (Heintz, A. 174, 154). - 2. Acetone saturated with ammonia is left to itself for three or four weeks. finely pounded oxalic acid is then added in quantity sufficient to form an acid salt, and a quantity of water equal to that of the acetone. The resulting crystallino precipitato is easily scparated by boiling alcohol into insoluble ammonium oxalate and soluble diacetonamine oxalate. A further quantity of this last salt remains in the mother-liquor, together with salts of other bases (Sokoloff a. Latschinoff, B. 7, 1384).

Properties .- Free diacetonamine, separated from either of its salts by adding strong soda-lye and agitating with other, is a colourless liquid lighter than water, having an ammoniacal odour and strong alkaline reaction; more soluble in cold than in hot water, mixes in all proportions with alcohol and ether; oxidises and turns brown on exposure to the air; forms crystalline salts with hydrochloric, sulphuric, and oxalic acids. By distillation it is for the most part resolved into NII, and mesityl oxide CaII, O, and on the other hand is easily formed by direct combination of these bodies: $C_6H_{10}O + NH_3 = C_6H_{13}NO$.

Salts.-C.H. NOHCl crystallises from alcohol in rhombic prisms, v. sol. alcohol, resolved by dry distillation into NH Cl and Colling (Heintz, A. 175, 252)—(C. H., NO, HCl), PtCl, 2H, O crystallises from water, in which it is easily soluble (according to Sokoloff a. Latschinoff; also in dilute alcohol), in orange-yellow monoclinio prisms containing 2H.O, which they give off in a vacuum (H.); under ordinary pressure (S. and L.). The normal oxalate (C, H, NO), C, H, O, forms monoclinic tablets, very soluble in cold water, less soluble in alcohol than the acid salt. This latter CaH13NO,CaHOHH2O, forms monoclinic prisms; very soluble in hot, less in oold, water; easily in boiling alcohol, from which it separates out almost completely on cooling. The pierate C,H,3NO,C,H,3(NO,),O,H,O, forms goldyellow needles, somewhat sparingly soluble in cold water. The sulphate (C.H.3NO) H2SO, forme monoclinic crystals (from alcohol).

Reactions .-- 1. HNO, decomposes the ealte forming di-acetone alcohol and mesityl oxide: $2C_6H_{13}NO + 2HNO_3 =$

 $C_0H_{12}O_2 + C_6H_{10}O + 2N_2 + 3H_2O_4$ 2. Chromic acid mixture converts it into paraformaldehyde together with formic, acetio, and anido-iso-valerio acids NH, CMe, CH, CO, H, and a small quantity of amido-iso-butyric acid NH.,CMe,,CO.H (Heintz, A. 108, 45).—3. Solid KOH forms an anhydride, C.,H.,N.O [83°]. This is v. sol. alcohol, chloroform, or benzene, m. so?. other or light petroleum. Hot water C.H.; NO C.S. CH., CO.CH., CMc., NH.; decomposes it (Antrick, 4, 227, 381). It forms

Preparation.—1. Dry ammonia-gae is passed into a flask containing boiling acetone, the con-

namine hydrochlorale heated for ten hours at | is formed by direct combination of its constituente. 120° with aqueous hydrocyanic acid forms the hydrochlorides of diacetonamine oyanhydrin nydrochoracs of diacetonamine dyannydrin and of nitrilo-diacetonamine, together with a little amido-io-butyrio acid (Hointz, A. 189, 231; 192, 340).—6. Diacetonamino oxalate boiled with alcoholic colutions of aldehydes forms condensation products.—6. Sodium amalgam reduces di-acetonamine to a secondary amidoiso-hexyl alcohol CH3.CH(OH).CH2.CMe2NH2. CYANHYDRINS.

Di-Acetonamine cyar hydrin

C,H₁₁N₂O or Mo.C(OH)(CN),CH₂.CMe₂.NH₂. Carbylo-di-acetonamine.—Prepared as do-scribed abovo (Reaction 4).—Trimetrio prisms. V. sol. water.. Decomposed by boiling alcohol into HCN and diacetonamine. Boiling HCl eaponifies it, forming OXY-AMIDO-DEPTOIC ACID (q. v.), Me. C(OH)(CO.H). CH., CMe., NII., the greater part of which changes to its anhydride, di-oxy-tri-methyl-pyrroline,

$$MeC(OH)$$
 $CO = NH$
 $CH_2 - CMe_2$

(Heintz, A. 192, 329; Weil, A. 232, 208 v. Pyrro-LINE).

Nitrilo-di-Acetonamine C,H, N,O. The hydrochlorido ie obtained, as above stated, to-gether with its isomeride. The free base is crystalline, easily soluble in water, sparingly in ether, and absorbs CO, from the air. Distinguished from carbylediacetonamine by remaining unaltered when heated to 100°-110° with fuming

hydrochlorio acid. Resolved by boiling with nyanocanono acia. Resolved by boiling with baryta water into NH, and amido-trimethyloxybutyric acid C.H., NO, or its anhydride. The platinochloride (C.H., N.O., HCl)_PtCl, forms yellow rhombic prisms elightly soluble in water. The oxalate C.H., N.O., C.H.O., forms small crystals in solution in the control of the control tals in. sol. water, insol. alcohol (Heintz, A. 192, 542).

[27°] (200°) vinyl-di-acetonamine; oxy-tri-methyl-

tetra-hydro-pyridine.

Formation .- Together with tri-acetonamine by action of aldehyde and ammonia on acctoue. In larger quantity as oxalate, by boiling the acid oxalate of diacetonamine (10 g.) for sixty hours in a reflux apparatus with aldehyde (10 g.) and alcohol (120 g.). The oxalate is washed with hot alcohol, and the free bases separated by potash (Heintz, A. 178, 326; 189, 214; 191, 122).

Preparation.-By boiling an alcoholic solution of di-acetonamino oxalate with paralde-

hyde (E. Fischer, B. 17, 1793).

Properties. - Solidifies at -15° to rectangular or six-sided plate ong prisms. Is deliquescent. Has a ! taste, sniells like trimethylamine, but when warmed, like camphor.

Reduced by codium amalgam to its dihydride

or ethenyl-di-amtome-alcamine.

Salts.—(B'HCl),PtCl,3aq. Flat prisms.—
B'H,SO, Minute needles, v. col. water, sl. col. altokol.—B',H,C,O, Sl. sol. alcohol.—B', 3H,C,O, A platino-chloride of vinyl-di-acetonamine

and tri-acetonamine (C,H, NO.HCI + C,H, NO.HCI)P(CI, + 2H,O 100 pts. water at 14° diesolve 8.65 pts. of the

anhydrous salt (Heintz, J. 1877, 442). Pentylidene-di-acetonamine $C_{11}H_{21}NO$

Valeral-di-acetonamine; oxy-di-methyl-iso-butyl-tetra-hydro-pyridine [15°-22°]. From valerio aldehyde and alcoholio di-acetonamine oxalate (Antrick, A. 227, 367). Needles in stare (from ether). Insol. water, sol. alcohol, ether, benzene, and petroleum.

Salls.—B',H₂C₂O, Needles [190°]. V. el. sol. cold water or alcohol.—(B'HCl),PtCl,[205°]. Heptylidene-di-acotonamine C₁₃H₂₃NO or

lidene-di-acotonamine
$$C_{13}^{22}$$
 C_{13}^{24} C_{13}^{24} C_{13}^{24} C_{13}^{24} C_{13}^{24} C_{14}^{24} C_{15}^{24} C_{15}

Oxy-di-methyl-hexyt-tetra-hydro-pyridins [29.5°]. From conanth-aldehyde and alcoholic di-acetonamine oxalate (Antrick, A. 227, 370). Needles (from ether). Oxalate B'2 H2O2O4 [0.150°].
Benzylideno-di-acetonamine C12H11NO or

$$Ph.CH < CH_2 \cdot CO \\ NH.CMe_2 \\ CH_2$$

Oxy-phenyl-di-methyl-tetra-hydro-pyridine [61°] Obtained as oxalate, by boiling I pt. benzaldeliyde, 1 pt. acid diacetonamine oxalato, and 12 pts. alcohol, gradually separating as a powder which may be purified by crystallisation powder which may be purined by crystalisation from water. Colourless needles or monoclinic prisms (from ether). V. sol. alcohol and ether; sl. sol. water. Tasteless, has a faint aromatic odour. Forms normal and acid salts.—C₁₃H₁₇NOHCl. Crusts or druses of crystals.—(C₂₃H₁₇NO.HCl.). PtCl. Warty groupe of crystals. tals, or when separated from alcohol on addition of ether, elongated six-sided tablets. Slightly soluble in hot, insol. in cold alcohol. The aurochloride forms pale yellow crystals .- $C_{13}H_{17}NOHNO_3 + 2H_2O(?)$. Smallerystals, moderately soluble in cold water. - (C18H1, NO)2H2SO4-Small crystals, easily soluble in water, very slightly in absolute alcohol. $-(C_{12}H_{17}NO)_{27}C_2H_2O_4$ Microscopie crystals, nearly insoluble in alcohol, v. sl. sol. water (R. Schiff, A. 193, 62).

m-Amido-benzylidene-di-acetonamine

From the nitro-derivative by reduction with SnCl₂. Oil. Salts.—B"H₂C₂O₁ [113°], p-Amide-beuzylidene-di-acetonamine.—From

the nitro derivative by SnCl. Salt. -B"H,C,O,. o-Nitro-benzylideno-di-acetonamine

From o-nitrobenzoio aldehyde and alcoholio diacetonamino oxalate.

Salts.—B'₂ H₂C₂O₄.—B'HCl.—(B' HCl),PtCl. m-Nitro-benzylidene-di-acetonamine.

Salts.—B'HCl [208°].—(B'HCl),Pt01, [303°].
p-Mitro-bensylidene-di-acetonamine [142.5°].
Needles (from ether). Nearly insol. light petrolenm.

Salts.-B'HCl sq. [c. 206°].-(B'HCl),PiCl,

p-Ozy-benzylidene-di-acetonamine CH, . CO HO.C.H..CH NH.CMe

From di-acetonamine oxalate (5 pts.), p-oxybenzoio aldehyde (4 pts.), and alcohol (20 pts.)

Acid oxalats B'H₂C₂O₄.

Methyl derivative .CH., CO. MeO.C,H.CH NH.CMe,

From anisaldehyde and di-acctonamine oxalate.

Oxalate B', II, O, O, [210°].

Cinnamylidene-di-acetonamine

 $\mathcal{C}\mathbf{H}_2$. CO CH₂ ½ aq. [49°]. Ph.CH:CH.CH NH.CMe

From oinnamie aldehyde, diacetonamine, and boiling alcohol. Yellow needles (from alcohol). Easily soluble in ether, light petroleum, chloroform and benzene, sparingly in water.

C, H, NO, Vanillo-di-acetonamine $CH_2 \cdot CO$ C.H.(OMe)(OH)CH NH.CMe.

is obtained by boiling equal parts of vanilling and acid diacetonamine oxalate diacetonamine cohol, whereby normal. This salt forms either oxalate is three or yellowish crystalline crusts; a white water, insol. alcohol and ether. gle base is an alkaline oil, slightly soluble in water.-C1.H1.NO.HCl is casily soluble in alcohol, and precipitated therefrom by other .-(C1.H1.NO.HCl),PtCl.-C1.H1.NO.HNO. small crystals, m. sol. water, and cold alcohol, (C₁₄H₁₉NO₃)₂H₂SO₄: lamine. (C₁₄H₁₉NO₃)₂C₂H₂O₄: crystalline, v. sl. sol. water, insol. alcohol (Heintz, A. 194, 53).

ALKYL-DI-ACETONAMINES.

Methyl di-acetonamine

C,H,5NO i.c. COMe.CIL_CMe_NHMe, is formed, together with other bases, when acetone saturated with methylamine is left to itself for several weeks. The base is ppd. as acid oxalate, and purified by conversion into platinochloride.-Free methyldiacetonamine is very unstable, quickly splitting up into methylamine and mesityl oxide. The hydrochloride is deliquescent. The platinochloride (C,H, NOHCl), PtCl, crystalliscs in large light-red rhombio prisms, casily soluble in water, nearly insoluble in alcohol. The platinosochloride (C.H, NOIICI), PtCl, produced simultaneously with the platinochloride, forms dark red crystals. The aurochloride C,H12NO,HCl,AuCl3, forms short prisms, m. sol. cold, v. sol. hot, water, alcohol, and ether. The pormal oxalate (C,H1,NO),C,II,O, forms indistinct very deliquescent crystals, very soluble in absolutealcohol; the acid oxalate C, H, NOC, H,O, crystallises in small prisms, m. sol. absolute alcohol. The picrate forms yellow needles (Götschmann, A. 197, 38).

Dimethyld cetonamine C₈H₁, NO i.e. CQMc.CH₂.CMo₂.NMe₂, is formed on heating a solution of dimethyla-mine in acctone at 100°-105° in a scaled tube. Free dimethyleiacetonamine has not been obtained as it very easily splits up into dimethylamine and mesityl oxide. The platino-chloride (C.H. NOHCl), PtCl, crystallises in small tablets;

the auro-chloride in golden needles, st. sot. water; the nitrate and sulphate in long colourloss deliquescent needles v. sol. alcohol. The acid oxalate, C.H., NOC, H.O., is crystalline, v. sol. water and alcohol, nearly include in ether (Götschmann, A. 197, 27).

Ethyldiacetonaming
C₂H₁₇NO i.e. MeCO.CH₂.CMe₂.NHEt,
is obtained by heating a solution of ethylamine in acetone at 80° for six hours. CieH3. N2O2PtCle, light red hexagonal plates, insol. ether and alcohol, soluble in alcohol containing HCl. S. 1-16 at 16°. Platinosochloride: C₁₆H₃₀N₂O₂PtOl₃; dark red prisms. S. 6-62 at 21°, insoluble in cther and in alcohol. - C₂H₁, NOHCl forms hygroscopic microcrystals decomposing at 150°. C_sH₁₈NOAuCl, crystallises in large lemon-yellow S. 2.48 at 22°; easily soluble rhombie plates. in alcohol and ether; melts under water at about 70°.—The nitrate forms small needles. (C.H., NO), H.SO, forms tufts of needles.-(C,H,,NO),C,H,O,, concentric groups of hygroscopio needles.—C,H,,NOC,H,O,; needles.—The picrate C,H,,NOC,H,(NO,),OH forms short nccdle-shaped prisms v. sol. water, insol. alcohol rece cunyidiacctonamine spitts np even in the cold into ethylamine and mesityl oxide (Eppinger, A. 204, 50). The prolonged heating of cthylamine with acctone gives rise only to ethyl-diacetonamine, not to any base analogous to triacetonamine. Diethylamino does not appear to form any compound with acetone (Eppinger).

Dehydrodiacetenamine C.H., N(?) contained in the mother-liquors of the preparation of acid diacetonamino oxalate, and passes over on dis-tilling them with an alkali. The platinochloride forms slightly sol. laminæ (Heintz, A. 183, 276).

Triactionamine Collin NO. i.e. CMe₂.CH₂ CMe2.CH2

Oxy-tetra-methyl-tetra-hydro-pyridine [58°] (hydrated); [39.6°] (dry). Formation.-1. To. gether with diacetonamine, by the action of ammonia on acctone, especially at high temperatures (Heintz, A. 174, 133).—2. By prolonged boiling of acetone with a solution of diaceto- $C_8H_{13}NO + C_8H_8O = C_9H_{17}NO + H_2O$ namine: (Heintz, A. 178, 305). This, according to Heintz, is the best mode of preparing triacctonamine. It is purified by crystallisation of the oxalate. Triacetonamine separates from a solution of the normal oxalate mixed with KOH, as a hydrate C₂H₁₇NO,H₂O, which crystallises from anhydrous ether in large square tablets, and the motherliquor on further evaporation and cooling to a very low temperature yields long needle-shaped crystals of anhydrous triacctonamine. Hydrated crystals rhombic a:b:c=0.9866:0.9768:1. Triacetonamine sublimes slowly, even at ord. temp. Distils without alteration. Decomposed at 150°-200° by H₂SO₄ or P₂O₃, but does not yield definite products. Heated at 100° for 16 hours with fuming hydrochlosic acid it yields discetonamine, dehydropentacetonamine and other products. With chromic acid mixture it gives isopropyl-butyl-amine di-carboxylic said: With chromic acid mixture it C₂H₁,NO₄ i.e. CO₄H.CMe₂NH.CMe₂CH₂CO₄H (Heintz, A. 198, 69). With ethyl iodide it yields NH₂Et, NHEt₂, NEt₄I, dehydrotriacetonamine, and other products, but no ethylated Is formed together with ammonia and di-aceto-

triacetonamines (Heintz, A. 201, 100).

Salts.—B' HCl is easily soluble in alcohol, and separates therefrom on addition of ether, in prisms (BHOI) PtCl 8H O crystallises from hot water in long, dark, gold coloured needles, v. sl. sol. alcohol, insol. scher. By exposing the alcoholic solntion to sunlight, or heating tho aqueous colution for several hours, it is reduced to (B'HCl) PtCl 2H2O, which is much less soluble in water than the platine-chloride, and crystallises in dark red needles or rhombic prisms.-(CaH12NO)2H2SO4: delicate needles or prisme v. sol. in water, insol. alcohol and ether .-C₂H₁₇NC,HNO₃: rhombio crystals -a:b:c= 1.2758: 1:10251.—(C.H.,NO)2H2CrO. Small light yellow crystals converted into the acid salt by recrystallisation from hot water .-(C₂H₁,NO)₂H₂Cr₂O₇. Orange-red prisms (Heintz, A. 198, 87).—(O₂H₁,NO)₂C₂H₂O₄ forms long needles, v. sol. water, v. sl. sol. alcohol.-C.H., NO, C.H.O. Triclinic crystals, v. sol. water; resolved by boiling with alcohol or ether into the normal salt and oxalic acid (Heintz, A. 178, 326).

Triacetonamine Nitrosamine C₂H₁₈(NO)NO [73°], S.G. 1°14, is formed by heating aqueous triacetonamine hydrochloride with KNO₂ at 85°. Long needles (from alcohol). V. sol. alcohol and ether. Resolved by KOH into nitrogen, water, and phorone, also by prolonged boiling in aqueous solution. By heating with HCl or H₂SO₃, it is for the most part reconverted into triacetonamino (Heintz, A. 185, 1; 187, 233).

Tri-acetone-diamine

C_pH₂₀N₁O t.e. (NH₂.CMe₂.CH₂)₂CO. Found in small quantity amongst the products of the action of ammonia on acctone; produced more abundantly when a mixture of 1 pt. acctone, 2 pts. NH₁Aq, and 1 pt. CS₂, is left at rest for a month; 3C₂H₂O + 2NH₃ = C₂H₂₀N₂O + 2H₂O. Oily liquid soluble in water, somewhat sparingly in ether. B"2HCl forms prismatic crystals, decomposing at 200°.—B"2HCl.PtCl₄ is slightly soluble in cold, easily in hot water, insoluble in ether.—B"C₂H₂O₄; flat needles, nearly insoluble in alcohol, much more soluble in water, than the acid salt.—B"2H₂C₂O₄ aq; monoclinio prisms (Heiutz, A. 203, 336).

Dehydro-tri-acetonamine C₀H₁₅N (Tetramethyl-di-hydro-pyridine?) (158°) (II.); (163°)

(C. S.).

Occurs as oxalate, together will tri-acctonamine, in the mother liquor got in preparing diacetonamine oxalate (q. v.), and may be separated thorefrom by distillation with potash (Heiutz, A.

174, 166; 183, 276).

Preparation.—Acetono (20g.), acetamide (8g.), and ZnCl₂ (30g.), are heated for 6 hours at 140° (Canzeroni a. Spica, G. 14, 341). Another base (240°) is a by-product in this reaction. It appears to be Cl₃H₂₇N. Its plating-chloride forms dodecahedra.

Properties .- Cily liquid which readily oxi-

dises, becoming brown.

Salts.—(B'HCl)₂PtCl₄. Rhombohedra (from water). V. sl. sol. cold water, insol. alcohol.— B HAuCl₄ [127°]. Long yellow prisms (from dilnts alcohol). Insol. water.

Dahydro-penta-acetonamine $C_{14}H_{12}N = 5C_1H_1C + NH_1 - 5H_2C$.

Is formed together with ammonia and di-acetonamine by heating tri-acetonamine with fuming HCl at 130°, the hydrochloride then separating as a crystalline powder, sparingly soluble in water. The base separated therefrom by potash is an oily liquid (Heintz, A. 181, 70). H. W.

is an oily liquid (Heintz, A. 181, 70). H. W. ACETO-NAPHTHYL-THIAMIDE v. a-Naph-

ACETONE C.H.C i.e. CH.CO.CH.

Di-methyl Ketone.

M. w. 58 (55·6°-55·9° cor.) (Perkin, C. J. 45, 478); (56°) (Dumas; R. Schiff); (56·3°) at 760 mm. (Kopp, Regnault, Zander); (56·53° cor.) (Thorpe, C. J. 37, 212). S.G. 2° 814; 132 799 (Kopp, A. 64, 214); 3° 8186 (T.); $\frac{1}{4}$ 3° 7965; $\frac{2}{3}$ 3° 7867 (P.); $\frac{1}{6}$ 3° 8125 (Z.); $\frac{29}{4}$ 4° 7920 (Brühl); $\frac{4}{4}$ 4° 7506 (R. Schiff, A. 220, 103). V.D. 200 (Dumas). C.E. (0°-10°) 00138 (T.). S.V. 77·08 (S.); 77·3 (Z.); 76·78 (T.). H. F. p. 65,000 (Berthelot); 58,710 (Thousen). H. F. v. 57,260 (Th.). μ_{B} 1° 3639. R_{∞} 25·55 (Brühl). M. M. 3·514 at 15·2° (P.).—Occurs in the urine, blood, and brain of diabetics (Markownikoff, B. 8, 1683; Peters, Kaulich, Betz, J. 1861, 805).

Formation.—1. By the dry distillation of acetates: e.g. (MeCO.O), Ba = Ma, CO + BaCO,.—2. From zinc-methyl and acetyl chloride;

(a) MeCOCl + ZnMc, = MeCClMe.OZnMe, (b) MeCClMe.OZnMe + H,O =

Mc_CO + IICl + ZnO + CH₄ (Freund, A. 118, 1).—3. By treating bromo or chloro-propylene with aqueous hypochlorous acid and mercuric oxide, whereby chloracetone is formed:

 $2C_3H_5Cl + 2HOCl + HgO = HgCl_2 + H_2O + 2(C1l_2Cl.CO.CH_3),$

and reducing this compound to acetone with zinc and HCl (Linnemanu, Bl. [2] 6, 216).—

4. By treating the isomeric compound, propylene oxide, with sodium-amalgam, and dehydrogeniang the resulting isopropyl alcohol with chromic mixture, C₂H₆O + H₂= (CH₃)₂CH.OH, and

(CH₃)₂CII.OH + O = H₂O + (CH₃)₂CO (Linnemann, A. 140, 178). Berthelot (C. R. 68, 334), effects the oxidation with aqueous solution of mercuric bromide (Kutscheroff, B. 14, 1541), or chloride (B. 17, 15), on allylene.—6. By passing aldehyde vapour over red-hot lime (Schlocnilch, Z. 5, 336).—7. Together with propionic aldehyde, by heating a dilute aqueous solution of propylene glycol at 180°-190° (Eltekoff, J. 11, 409).—8. By heating propylene bromide with water at 170°-180°:

C₃H₆Br₂+H₂O=2HBr+C₄H₆O (Linnemann, A. 161, 58).—9. By heating a-a-dichloro-propane CMe₂Cl₂ with eilver acetate and alcohol in sealed tubes at 100°: CMe₂Cl₂+2AgCO₂Me=2AgCl+2COMe₂+CC₂.—10. Together with a bromine compound (probably CHMo₂Br) by the action of zinc and dilute sulphuric acid on the product C₂H₂Cl₂Br₂O, formed by the action of bromise on dichlorhydrin (Lange, B. 6, 98).—11. By distilling with water the product formed, with evolution of HCl, on dissolving chloro-propylene Me₂CCl:CH₂ in sulpharic acid (Oppenheim, A. Suppl. 6, 365).—12. Together with mesitylene, on distilling with water as olution of allylene in sulphuric acid (Schrohe, B. 8, 367).—13. Together with other

products, by the action of lime on glycerin (Tawilderow, B. 12, 1487).—14. Together with isohutyric aldehyde, by oxidation of ico-butyl alcohol.—15. By oxidising with chromic acid the hexylene obtained by the action of alcoholic potash on di-methyl-isopropyl-capbinyl iodide (Pawlow, Bl. [2] 29, 875).—16. By the action of nascent zino-methyl on acetic oxide (Saytzeff, Z. [2] 7, 104): (COMe), O + ZnMe, = ZnO + 2COMe, —17. Together with other products, by the action of zinc-eodium on a mixture of methyl iodide and acetic oxide (S.).—18. By the dry distillation of wood: occurs therefore in crude wood-spirit; also of sugar, gum, or staroll, with 8 pts. lime (Fremy, A. Ch. 59, 7).—19. By dry distillation of citric acid, and in the oxidation of that acid hy potassium permanganate, or by MnO₂ and dilute sulphuric acid (Péan de St. Gilles, A. Ch. [3] 55, 374).

Preparation.-1. By dry distillation of barium or calcium acetate. The barium salt decomposes at a moderate heat, and when dry and pure yielde pure colourless acetone. The calcium salt requires a higher temporature and yields a distillate contaminated with an empyreumatic oil (dumasin) and other products. -2. By distilling in an iron retert or quicksilver bottle, a mixture of lead acetate (2 pts.) and quick lime (1 pt.), rectifying over calcium chloride, and finally distilling over the water-bath. The product may be purified from wood-spirit by distillation over calcium chloride, or better by combining the acctone with sodium hydrogen snlphite, and decomposing the resulting compound by an acid or alkali; also by converting the methyl alcohol into an ether (oxalic or benzoic). Crude acetone may also be purified by treating it with potassium permanganate, which does not attack pure acctone at ordinary temperatures.

Properties. — Limpid, very mobile liquid having a epirituous and elightly empyreumatic odour and biting taste. Very inflammable; hurns with a white smokeless flame, mixes in all proportions with water, alcohol, and ether. Dissolves camphor, tats, and resins. Separated from aqueous solution by CaCl, and by KOH (difference from alcohol). Even if boiling between 56° and 58° it is liable to contain methylacetal, CH₂,CH(OCH₂)₂; this can be detected by heating with cone. HCl, for it then gives off MeCl. Acetone reacts with hydroxylamine and with phenyl-hydrazine (v. Acetonix, Acetonic Theyl-hydrazine). It does not restore the dneed of a solution of a rosaniline salt that forms dateached by SO₂ (Schiff).

C.H., NO.H.C., 1. An alcoholic liquid supposed cold, v. sol. hot, is mixed with an equal volume germal exalate (C.p. of benzoic aldehyde and finct very deliqueseen H are added. After some absolute alcohol; the acic 4i-benzylidene-acctone crystallises in small prisalve in H_SO, giving alcohol. The picrate form. Ponder, A. 223, (Götschmann, A. 197, 38). ueous KOH and

Dimethyldhectonamine 3, A solution C₁H₁,NO i.e. CQMc.CH₂.CMe₂, previously is formed on heating a solution of A black mine in acctone at 100°-105° in a sears, but Free dimethyldiacctonamine has not des this tained as it very easily splits np into dimee to amine and mesityl oxide. The platino-chlo-be (C₂H₂NOHCl)₂PtOl₃ cryetallises in small tables

Reactions.-1. Acetone-varour passed through. red-hot tube deposits carbon and yields sooalled dumaein, also naphthalene, CO, CH, and H (Barbieri a. Roux, C.R. 102, 1559).—2. By nascent hydrogen (sodium-amalgam and water) acetone is converted into isopropyl alcohol: Me.CO.Me + H₂ = Mc.CHOH.Me (Friedel, O.R. 55, 58).—8. Chlorine-gas passed into acetone displaces 1 or 2 ats. H, forming C,H,ClO and C3H,Cl2O, but does not remove the whole of the hydrogen, even in sunshine. Grabowski (B. 8, 1438), by passing chlorine into pure acetone, assisting the action by heat towards the end, obtained in addition to dichloracctone, two bodies C₃H₂Cl₃O and C₆H₂Cl₃O. The former is a liquid insoluble in water (186°). S.G. 1·330 at 29°. V.D. 6·60 (calc. 6·56). Decomposed by etrong potash-lye, with separation of chloroform. The second body, C₆H,Cl₈O, is also liquid (206°–208°). S.G. 1·326 at 26°. V.D. 7·55 (calc. 7·0). Completely decomposed by strong potash-lye or sulphuric acid. Perhapstrichlorotrimesityloxide. When acetone is treated with excess of chlorine, and the product first with KOH and then with HCl, isapoglucic acid CoH10Os is produced. With alcoholic potash, on the other hand, a body C_sH₁₀O₅ (?) is formed, together with an acid whose lead-salt has the composition Pb(C, H,O,), (Mulder, J. 1868, 494) .- 4. Chlorine, in presence of alkalis, converts acctone into chloroform: $C_2H_0O + 6Cl_2 + H_2O = 2CHCl_2 + CO_2 + 6HCl_3$ Bromino acts in like manner, producing bromoform, and iodino forms iodoform.—5. When acctone saturated with HCl-gas is mixed, after 8-14 days, with water, a heavy brownish oil separates, consisting mainly of compounds of and phorone, $C_9H_{10}O(=2C_2H_0O - H_0O)$ and phorone, $C_9H_{10}O(=3C_2H_0O - 2H_0O)$. The mesityl compound $C_9H_{10}O(H_0O)$ heated with KCN and then with KOH, yields the K-salt of a modular interval of the compound $C_9H_{10}O(H_0O)$ nobasic acid CaH, NO, (r. Mesitonic Acid), thus: $C_8H_{12}OCl_2 + 2KCN = 2KCl + C_8H_{12}O(CN)_2$; and $C_8H_{12}O(CN)_2 + KOH + H_2O = NH_3 + KC_8H_{12}NO_3$. The phorone compound, similarly treated, yields a neutral azetised body crystallising in shining plates and subliming at about 300° (Maxwell Simpson, Pr. 16, 364). According to Pinner (B. 14, 1070) the neutral body is a nitrile $C_{11}H_{10}O_2N_2$, formed according to the equation $3C_2H_2O + 2HCN = H_2O + C_{11}H_{10}O_2N_2$; it crystallises in plates melting above 320°. Heated with aqueous hydrochloric acid it gives phoronic acid $C_{11}H_{18}O_{2}$ $[q. v.]: C_{9}H_{18}O_{2}(CN)_{2} + 4H_{2}O = 2NH_{2} + H_{2}O + C_{9}H_{18}O(CO_{2}H)_{2} - 6$. By distillation with strong sulphuric acid, acetone yields mesitylene, C,H₁₂ = 3C,H,O - 3H₂O; but when mixed with H.SO, in a cooled vessel it forms mesitylsulphonic acid CaHa.SOaH, which, when heated with potash, yields mesityl oxide (Hlasiwetz, J. 1856, 487).—7. With PCl, actione yields chloropropylene C,H,Cl and di-chloro-propane C3HaCl2 (Friedel A. 112, 236).-8. With bromine acctone unites directly, forming C.H.OBr., a viscid, very unstable liquid, heavier than water (Linnemann, A. 125, 307). According to E. J. Mulder, however (J. pr. 91, 47), it gives rise to eubetitution-producte.—9. With HI acctone yields iodopropylene, C.H.I; with PI, a could and two liquid iodides (Harnitz-Harnitzky, Z. 1863, 416). According to Berthelot (Bl. [2] 7, 69), acctone freated with HI yields propane.

19, With todine chleride acetone yields C.H.I.O | like manner from acetone and p-netro-benzalde-(Maxwell Simpson, Laboratory, p. 79) .- 11. Electrolysis of a mixture of acetone and dilnte sulphurio soid produces acetic, formic, and carbonic acids (Arredel, J. 1852, 838).—12. By chromic acid mixture it is oxidised to acctic and carbonic acids.-13. Acctone heated with ammonia yields a mixture of three bases, the composition and mode of formation of which are indicated by the following formula:-

Diacotonamine $C_aH_{1a}NO=2C_3H_aO+NH_3-H_3O$ Triacotonamine . . . $C_aH_{1a}NO=3C_aH_aO+NH_3-2H_3O$ Dehydrotriacotonamine . $C_aH_{1a}N=C_bH_{1a}NO-H_aO$

With methylamine, in like manner, acctone yields methyldiacetonamine C,H1,NO and other bases. With dimethylamine only dimethyldi-acetonamino CaH, NO. -14. With hydroxylamine, acetone forms ACETOXIM [q.v.] Mc.C:NOH, which crystallises in prisms [60°], (135°).—15. Sodium strongly attacks acetone, with formation of orystallised pinacone hydrate C,H,2O 7H2O Na₂O + C₄H₁₂O, and 3C₂H₁₂O + C₂H₁₂O + Na₂ = Na₄O + C₄H₁₂O, and 3C₂H₁₂O - 2H₁O = C₃H₁₄O (Städeler, A. 111, 277).—16. Heated with aniline hydrochloride at 180° it forms di-methyl-culture (Feeder & Bishw B. 11, 2017). •quinoline (Engler a. Riehm. B. 18, 2245, 3296). -17. Caustic alkalis, e.g. KOH and CaO, exert a dehydrating action on acetone and form condensation-products varying in composition, according to the proportion of water abstracted, viz.:

Vaponr of acctone passed over strongly heated KOH or potash-lime is resolved into methane and carbonic acid, $C_3H_cO + 2KOH = K_cCO_3 + 2CH_1$. At a lower temperature the chief products are acctic acid, formic acid, and hydrogen, $C_3H_4O + 2KOH + H_2O = KC_2H_3O_2 + KCHO_2 + 3H_3$ (Dumas a. Stas, A. Ch. [2] 73, 149; Persoz, Rev. Scient. 1, 51).—18. Acctong heated with ZnCl. yields have methyl-benzene C₈Me₈ (W. 11 Greene, C. R. 87, 931).-19. Gently heated with AlCl., it yields mesityl oxide, phorone, and other products (Louise, C. R. 95, 602).—20. Dry PtCl, dissolves in acctone, and the solution when evaporated leaves a brown resinous mass containing a yellow crystalline substance, $C_cH_{10}OPtCl_1$ (?), called Acechloride of Platinum, or Chloroplatinite of Mesityl (Zeise, A. 33, 29) .- 21. On adding HCl to a mixture of acetone, with potassium cyanide and sulpho-cyanide, the compound C₃H,O₂NS is obtained. This compound heated with HCl is resolved into CO₂₂ NH₄, and a-oxy-iso-butyric acid. With silver nitrate it yields C₅H_aAgO NS (Urech, B. 0, 1113).—229 By action of alkalis or of HClgas on a mixture of 1 mol. acetone and 2 mol. benzaldehyde, nibenzylinene acetone (q. v.) PhCH;CH.CO.CH:CHPh is obtained (Claison a. Claparede, B. 14, 849). By the action of alkalis on a solution of o-nitro-benzaldehyde in acetone, methyl o-nitro-8-oxy-8-phenyl-ethyl ketone [68°] is formed according to the equation

NO.C.H.CHO+CO(CH.).= NO.C.H.CH(OH).CH.COCH

hyde (Baeyer a. Becker, B. 16, 1968) .- 23. With furfuraldehyde, acetone forms a compound crystallising in long white needles [87°] (J. G. Schmidt, B. 14, 574) -v. FURFURYLIDENE-ACETONE. 24. With purrol in presence of HCl it forms $C_{1}, H_{1}, N_{2}, (291^{\circ})$ (Baeyer, B. 19, 2184). Combinations. -1. With Bisulphites.

Formed by direct combination. C.H. (OH) SO, NH. erystallises in lamine (Städeler, A. 111, 307) C₂H_q(OH)SO₃Na.—Lamine, moderately soluble in water, less in alcohol. Gives off acetone when boiled with aqueous sodium carbonate (Limpricht, A. 93, 238)—C₃H₆(OH)SO₂K (L.). 2. With Meronric Oxide 2C₃H₆O 3HgO. Formed by mixing acctone with mercuric chloride and weak potash-lye, dialysing the filtered liquid, and precipitating the liquid remaining in the dialyser with acetic acid. - Gelatinous precipitate which becomes resinous on drying. Its solution gelatinises when heated or whon merely left at rest (Emerson Reynolds, Pr. 19, 431). Formed also by dissolving HgO in acetone

Acetone-boric Acid, C.H.O(BHO), [50°]. Formed together with (a) and (B) acetone-fluoboric acid, and hydrocarbons, on saturating acctone with boron fluoride and distilling the product. (a) Acetonefluoboric acid, C₄H₆O 3HFB₄O (120°-123°); the isomeric (8) modification [36°] (90°-92°) forms shining white lamine. three compounds fume in the air, burn with green flame, and are quickly decomposed by water, yielding boric acid and acotone hydro-

(Kutscheroff, B. 17, 20).

fluoric acid (Landolf, C. R. 89, 173).

Acetone-hydrofluoric Acid C₃H₄OHF (55°) obtained by fractional distillation from the product of the action of water on acctone-fluoboric acid. An inflammable liquid with pleasant ethereal odour (Landolf, C. R. 96, 580). Another compound, C₃H_sO 2HF (12°) is gaseons at ordinary temperatures.

Acstone-sulphonic Acid CH3.CO.CH, SO3H. Formed as K-salt by treating dichloracetone (118°) with a strong solution of potassium snl-phite: C₃H₂Cl₂O + K₂SO₃ + H₂O = K₂SO₄ + HCl + C₃I₁ClO, and

 $C_3H_3ClO + K_2SO_3 = KCl + C_2H_3O.SO_3K$ The K-salt may be extracted from the product by boiling alcohol, and separates therefrom in white lamine. Very soluble in water, not decomposed by boiling with dilute acids. Boiled with strong potash-lyc, it yields sulphito and perhaps an acetone alcohol (Bender, Z. 1870, 162; B. 4, 517). Salts. -KA' Plates (from alcohol V. e. sol. water—BaA'₂ aq. Plates.—PbA'₂ aq.

[140°]—CuA', 14aq. Greenish plates.
Acstone-phosphorous Acid C,H₀O.PO.H.
Remains on distilling accions with I and P. (C.H.O.PO.) Ba is amorphous, soluble in water,

Acetone - cyanhydrin CH, C(CN) (OH) . CH, (Oxyisobutyronitrile). Formed by the action of aqueous HCN (20 p.c.) on acctonc, the by the action of nascent HCN on acctone diluted with ether.

It is very unstable, for even on evaporation Its solution it changes into di-acetone-cyanhydrin with evolution of HCN (Tiemann a. Friodlander, B. 14, 1970). Alcoholio NH, Bacyer a. Drewsen, B. 15, 2856).—The corre-at 60° converts it into a amido-iso-butyronitrile tponding para-compound [58°] is obtained in CH, C(ON)(NH₂), CH, whence HCl forms a amidoiso-butyric seid. Alcoholic HCl forms the imido-ether Me,C(OH)C(OEt):NH (Pinner, B.

Discotone cyanhydrin CMe, (CN).O.CMe, (OH), is prepared by adding 1 mol. HCl (gaseons or aqueous) to 1 mol. KCN immersed in acetone. dissolving the product in other, and evaporating (Urech, A. 164, 259). Thick shining anhydrous prisms, easily soluble in water, alcohol, and ether. Melts at 135°-152° and sublimes below at ord. temp. by HCl into No. acctone, and acoxy-iso-butyric acid.

Substitution Products v. 5365

CHLORO-ACETONE, CHLORO-DROMO-ACE 25 ACETONE, CYANO-ACETONE, THIO-ACETO.

Meta-acetone. - This name was Fremy (A. Ch. [2] 59, 6) to an oil ot among the products of the distillation of . starch, or gum, with quicklime. He ascribed it the formula C6H10O and boiling-point 84°. Gottlieb (A. 52, 128) converted it by chromic mixture into propionic acid (called therefore Metacetonic acid). Benedikt (A. 162, 303) found V.D. 3.53 instead of 3.59, and stated that it did not combine with NaHSO₃. Meta-acctone has also been examined by Favre (A. Ch. [3] 11, 80), Cahours (C. R. 30, 319), who describes it as present in crude wood spirit, Lies-Bodart (J. 1856, 455), and Schwartz (J. 1850, 533). Nevertheless Pinner (B. 15, 586; 16, 1729) considers metacetone to be a very complicated mixture.

Para-acetone v. PINACONE.

ACETONE-ALCAMINES. - These are products derived from the acctonamines by reduction, their CO being converted into CH(O11).

Diacetone - alcamine C₆H₁₃.NO NH2.CMe2.CH2.CH(OH).Me (175°). Formed by reduction of diacetonamine by gradually adding sodium-amalgam to its solution in alcohol diluted with aqueous ammonia. Liquid, having a faint ammoniacal odour, miseible in all proportions with water. Absorbs CO2 from the air; fumes with HCl. (C6H15NOHCl),PtCl, forms orange-red triclinic crystals, easily soluble in hot water.

Ethylidene-diacetone-alcamine

H_C-CH(OH)-CH2 C₈H₁₇NO i.e. MeHC - NH - CMe2

Oxy-tri-methyl-hexa-hydro-pyridine [123°].
Colourless crystalline colid. Easily soluble in water and alcohol, sparingly in other, and benforme. Formed by reduction of othylidene-C, H, NO, Mine with codium-amalgam in slightly cold, v. sol, hot, solution. The hydro-chlorido pormal oxalate (calles, the eulphate large flat inct very deliquescen 7, 1794).

absolute alcohol; the acid, C.H., NO crystallises in small pnsat: alcohol. The picrate form. I [128-5] (Götschmann, A. 197, 38). uello: 3, inc. Formed by reduction by reduction of the picrate o

Dimethyld cetonamine 3, inc. Formed C.H., NO i.e. COMc.CH. CMd. pr. by reducis formed on heating a solution of Algam in mine in acctone at 100°-105° in a cears, B. 17 Free dimethyldiacetonamine has not his this tained as it very easily splits up into dime to amine and mesityl oxide. The platino-chi. be (C.H. NOHCl), PtCl, crystallises in small table | 69), dich on saponification produces citric soid.

forms rhombic crystals (Heintz, A. 188, 200,

817). Methyl-tri-acetone-alcamine CleH2NO [74°] or, when hydrated, [60°]. Formed from tri-acetone-sleaming by MeI and MepH at 100° (E. water). Strongly alkaline. H. W. ACETYL-CARBINOL.

ACETONE-AMMONIA v. AOETONAMINES. ACETONE-BENZIL C₁,H₁₀O₅ i.e. Ph.CO.CPh(OH).CH₂, CO.CH₃ [78°].

Preparation.-Benzil is shaken with excess of puro acctone and a little conc. KOH, and the crystals obtained are dissolved in ether (free rom alcohol), which ie allowed to ovaporate. $_{14}H_{16}O_{2} + C_{3}H_{6}O = C_{17}H_{16}O_{3}$ (Japp a. Miller, C. 47, 21).

Properties .- Colourless square prisms. Sol. her or alcohol. Resolved by heat into its conituents.

Reactions.-1. Chromic mixture gives benzoic and acetic acids.—2. Dry NH₂ gives acetone-benzilimide (q. v.).—3. Alcoholic hydroxylamine gives C₁₇H₁₈O₂(NOH), [146]; m. sol. benzene, sl. sol. ether. This body is not affected by further treatment with hydroxylamine.

Dehydro-acetone-benzil C1:H1:O2 i.e.

CH₂ CO [149°]. Ph.CO.CPh

Preparation .- Benzil is chaken with excess of pure acetono and excess of cone. KOH (J. a. M.) $C_{14}H_{10}O_2 + C_3H_8O = C_{17}H_{14}O_2 + H_2O$. Properties.—Colourless priems.

Reactions. - 1. Converted by bromine in chloroform to a bromo derivative, $\hat{C}_1, H_1 BrO_2$ [172°]; slender needles (from glacial HOAc).—2. Chromic acid in glacial 110Ac forms an acid, C16H16O2, [152°]; needles. Salts, Aga'—BaA', 2acf. Dehydro-acetone-di-benzil C₃₁H₂₁O₄ [195°]

Formation .- 1. From acetone-benzil and dilute alcoholic KOII .- 2. From acetone, excess of benzil, and a little conc. KOII.

 $2C_{14}H_{16}O_2 + C_3H_6O = C_{31}11_{24}O_4 + H_2O_4$ Properties. - Colourless crystals (from benzene); sl. sol. boiling alcohol. Crystallises from alcohol with one molecule EtOH.

ACETONE-BENZILIMIDE C1,111,NO2 [176°]. From acetone-benzil and dry NH₃. Flat plates (from alcohol). Heated with HCl and oxalic acid, gives a red gum (J. a. M.).

ACETONE-BORIC ACID v. ACETONE. ACETONE - BROMIDE = DI - BROMO - PROPANE

ACETONE - BROMOFORM C.H.OBr. Me₂C(OH).CBr₂ [175°], or, when hydrated [167°]. From bromoform (5g.), acetone (30g.), and sodalime (8g.) (Willgerodt a. A. Müller, C. C. 1884,

ACETONE CARBOXYLIC ACID = ACETO-

ACETIC ACID (q. v.).

Acetone di-carboxylic acid C,H,O, i.e. CO₂H.CH₂.CO.CH₂CO₂H [o. 130°]. Formed by heating citric acid with H₂SO₄. Colourless needles. Split up into CO, and acctone by heat, by boiling water, or by warm acids or al-It contains methylenic hydrogen displacable by Na. Fe Cl, gives a violet colour. It reacts with phenyl-hydrazine (Pechmann, B. 1,17, 2542). It forms a compound with HCN,

MaNO, converts it into di-oximido-acetone (Pechrann a. Wahsarg, B. 19, 2465).

The sthyl sther is an oil which can give rise to salts by exchanging its methylenic hydrogen for sodium or oopper. Reactions. -1.
By successive treatment with sodium and an alkyl iodids (RI) the following sthers may be got: CO₂Et.CH₂.CO.CHR.CO₂Et,

· OO,Et.OHR.CO.CHR.CO.Et, CO,Et.CHR.CO,OR,.CO.Et,

and finally CO, Et. CR, CO.CR, CO, Et. The acids obtained by saponifying these bodies are split up by heat into CO2 and alkyl-acetonss (Dünschmann a. v. Pechmann, B. 18, 2289).— 2. Acetone di-carboxylic sther (100 g.) is converted by heating with Na (21 g.) into di-oxyphsnyl-acetic di-carboxylio ether

O_sH(OH)₂(CO₂Et)₂OH₂CO₂Et (Cornslius a. Pechmann, B. 19, 1446).—3. Ammonia produces \$ oxy-\$-amido-glutaramic ether, CO₂Et.CH₂.C(OH) (NH₂).CH₂.CO.NH₂ [86°] (v. Pschmann a. Stokes, B. 18, 2290; 19, 2694). ACETONE CHLORIDE v. di-CHLORO-PROPANE.

ACETONE CHLOROFORM C,H,OCl, i.e. Me2O(OH).CCl .. Oxy - iso - butyro - tri - chloride. [96°] or when hydrated, + \aq,[81°] (167° uncor.).
• Prepared by adding solid KOH to a cold mixture of acetons and chloroform. It is a crystalline solid, resembling camphor in appearance and

smsll. Rotatss on water. V. sol. alcohol, other, ohloroform, acetone, or glacial HOAc, insol. water. Volatile with steam. Converted by water at 180° into α-oxy-iso-butyric acid (Willgerodt, B. 14, 2451; 15, 2505; 16, 1585). ACETONE CYANHYDRIN v. ACETONE.

ACETONE-HYDROQUINONE C.H.,O. From acetone and hydroquinone (Habermann, M. 5,

ACETONE HYDROXYLAMIDE v. ACETOXIM. ACETONE-PHENANTHRAQUINONE

O, H, O, [90]. Formation.—From phenanthraquinone by heating with a large excess of acctone at 200°. The product is washed with NaHSO, and extracted with ether (Japp a. Streatfeild, C. J. 41, 274).

Preparation.—Phenanthraquinone (50 g.) is

shaken in a glass with acetone (60 g.) and cone. NH, Aq(40 c.c.). Acetonc phenanthraquinonimide is formed and filtered off; after washing with ether, it is made into a cream with water and stirred into a solution of oxalie acid (90 g.) in water (800 c.o.) at 25°. The substance dissolves, but minute heedles of acctone-pheuanthraquiuone soon ssparats (Japp a. Miller, C. J. 47, 18).

Properties .- Largo thin blades (from other). Insol. water, v. sol. other, acetone or alcohol.

Reactions.-1. By heat, by boiling water, or by boiling alcohol, it is resolved into acstone and phsnanthraquinono: C₁,H₁₄O₂ = C₁₄H₂O₂ + C₂H₃O₄.

2. Zino dust and glacial HOAc form C₁,H₁₀O₄ [121°]. This is extracted by other and crystallissd from alcohol. It forms long slender needles, v. s. sol. ether or chloroform, v. sol. boiling alcohol, v. sl. sol. cold alcohol. Sublimes in feathsry crystals. It accolorises bromins.—3. A few drops of dilute aqueous KOH added to an alcoholic solution forms minute crystals of scetone-di-phenanthraquinone (q. v.).—4. Cors. KOH (S.G. 1.27) addsd to a solution of acetonephenanthraquinone in acstone forms a crystal-line mass of di-acetons phenanthraquinons.— VOL L

5. Ammonia passed into an ethereal solution of acetone-phenanthraquinone forms crystals of acetone-phenanthraquiffonimide.

Acetone - di - phenanthraquinone [190°]. Formsd by adding a little dilute KOH to an alcoholic solution of acetone-phenanthraquinons (J. a. M.) * 2C₁₇H₁₀O₃ = C₃₁H₂₂O₅ + C₅H₆O₅ Colourless crystals (from banzens).

Di - acetone - phsnanthraquinone C.H.O.

Preparation.—Purs acctone (43 g.) is shaken with finely powdered phenanthraquinous (50 g.) and a little (2 c.o.) conc. KOHAq (S.G. 1-27). After 12 hrs. the resulting solid caks is washed with ether and crystallised from acctons (J.a.M.).

Properties.—Short oblique prisms. Sparingly soluble in the usual menstrus. Sol. scetone or benzeno. Decomposed by boiling glacial HOAc or amyl alcohol. Decomposed on multing into acctone and phenanthraquinene.

Reactions.—Boiling Ao O converts it into de-

hydro di-acctons phonanthraquinone.

Dehydro-di-acstons-phenanthraquinone $_{0}\mathrm{H_{18}O_{3}}$ [179°-181°]. Colourless pointed prisms (from benzens). Formed as abovs. Its constitution is perhaps:

Dshydro-acstone-phenanthraquincae C₁₇H₁₂O₂ [195°]. Formed in small quantity, together with di-acetone-phenanthraquinone when excess of KOHAq (S.G. 1.27) acts upon a mixture of acetone and phenanthraquinone. It is present in the ethercal washings of the diacstonc-phenanthraquinone (J. a. M.).

Groups of minute needles (from benzene): m. sol. hot benzene, hot alcohol, or sthsr.

Acetone-phenanthraquinonimids O, H15 NO2 [130°]. Formation.—By passing NH, into an ctlicrcal solution of acetone-phenanthraquinone.

Preparation .- Phenanthraquinone (50 g.). acetone (60 g.), and conc. NH, Aq (40 c.c.), shaken together form a white crystalline powder which is washed with other and crystallised from acetone containing a little NHaAq:

 $C_{11}H_{0}O_{2} + C_{3}H_{0}O + NH_{3} = O_{17}H_{15}NO_{2} + H_{2}Q$ (J. a. S.). Colourless rhomboidal lamine.

Reactions.-1. Ac O decomposes it, forming phenanthraquinono. -2. Cold conc. HOl dissolves it, but the solution soon dsposits a dark blue substance. The diluted filtrate deposits colourless needles of acetone-phenanthraquinone:

 $C_{17}H_{15}NO_2 + H_2O = C_{17}H_{14}O_3 + NH_3$ By using cono. aqueons oxalic acid as a solvent the formation of the blue compound may be avoided, and after dilution, the needles separate as before.

The following constitutional formulæ are suggested by Japp and Miller to explain the properties of the preceding bodiss:

C.H..C(OH).CH..CO.Me for C, F.O. C.H.CO C_sH₄.C(OH).CH₂.CO.Ms C₀H₁O.NH for C₁H₁₅NO₃
O₄H₄O(OH).CH₂CO.Me
O₄H₄.C(OH).CH₂CO.Me ACETONE-PHENYL-HYDRAZIDE:

(CH_e)₂C:N.NHPh (165°) at 91 mm. Oil. Prepared by mixing acetone with phenyl-hydrazine. It dissolves in cold aqueons acids and on warming the solution it is resolved into its constituents (Reisenegger, B. 16, 662).
DI-ACETONE-PHQSPHINIC ACID

 $C_4H_{12}PO_4$ aq, i.e. $(CH_2)_2CH.CHAc.PO(OH)_2$ aq, or, less probably $(CH_1,CO,CH_2)_2PH(OH)_2$ aq. Diacetonyl-phosphinic acid, iso-propyl-acetonylphosphonic acid, w-acetyl-iso-butane w-phosphonic acid [64°]. Formed by the action of water npon di-acetone-phosphorous ohloride (infra). (Michaelis, B. 17, 1273; 18, 902):

 $C_6H_{10}O_2PCl + 21I_2O = C_6H_{13}PO_4 + HCl$ Slender needles. V. c. sol. water or alcohol, v.

sol. ether. Strong dibasic acid.

Salts .- NII, HA". V. sol. water, sl. sol. alcohol. Crystals. -(NH.) 11A" 2 2aq. Insol. alcohol. —BaH,A", 2aq. Needles; v. sol. water, sl. sol. alcohol.—BaA" 6aq; m. sol. hot water; trimetric tables, a:b:c=.785:1:2.525. PbA".-PbA" 1 PbO.-MgA" 6aq; ppd. by alcohol from aqueous solution in glittering plates. -KHA"; deliquescent gum: v. sol. alcohol.-KH₂A"₂; slender needles, v. sol. water, sl. sol. alcohol.—Ag₂A".

Reactions. - 1. HNO, forms a tribasic crystalline acid C.H.PO, possibly &-carboxy-propanephosphonio acid: CH₃.CH(CO₂H).CH₂.1 O(OI1)₂.

The salts Ag_AA", and Ba_AA", are crystalline.

Oxim.—C₈H₁₈(NOH)PO₂ [170°). Colourless crystals; v. sol. water or alcohol; dibasic acid. Colourless

Di-Acotone-phenyl-phosphinic Acid C₆H₁₂(C₂H₁)PO₂, probably (CH₂)₂CH.CH(CO.CH₂).PO(C₆H₂)OH. [86°]. Prepared by adding P.O. to a mixture of acctone and phosphenyl chloride, and treating the product with water

(1) $2C_3H_6O + C_6H_3PCl_2 - H_2O = C_6H_{10}(C_6H_5)POCl_2$ (2) $C_0H_{10}(C_0H_5)POCl_2 + 2H_2O =$

Long colourless plates (+H₂O). Sol. not water, sl. sol. cold water and ether, v. e. sol. alcohol. The anhydrous acid forms a glassy mass, v. sol. ether. A'Ag; crystals, v. sol. water. (Michaelis, B. 19, 1009.)

Di-Acetone-p-tolyl-phosphinic Acid

C_sH₁₂(C,H₂)P₀, probably (CH₅)_sCH.CH(CO.CH₃).PO(C,H₃)OH. [103°]. Obtained by adding P2Os to a mixture of acctone and p-tolyl-phosphorous chloride, and treating the product with water. Glistening plates; sol. hot water, v. sol. alcohol and ether. slender glistening soluble needles. (Michaelis, B. 19, 1012.)

DI-ACETONE-PHOSPHORIC-TRI-CHLORIDE

 $(CH_3)_2:C \rightarrow O$ C.H,O.PCL i.S. CH,.CO.C11 - PCl, [115°]. Formed by passing chlorine into a solntion of di-acetone-phosphorous chloride in petroleum-ether (Michaelis, B. 18, 901). Colourless crystale; sl. sol. petroleum-ether. DLACETONE-PHOSPHORIC-CHLORO-BROMIDE

(CH₂)₂C = O C.H.O.PCIBr. i.e. CH₂.CO.CH - PCIBr₂ [1420]. Formed by addition of bromfae to a solution of di-acetone-phosphorous chloride in light petroleum (Michaelis, B. 18, 900). Colourless erustals. Sl. sol, light petroleum. It is decomposed by water into mesityl oxide, phosphoric soid, HOl, and HBr.

ACETONE-PHOSPHOROUS ACID v. ACETONE. DI-ACETQNE-PHOSPHOROUS CHLORIDE $(CH_{i})_{i}C = O$ CoH10O2POFice.

CH..CO.CH. PCI (154°) at 100 mm.; (235°) at 745 mm. S.G. (liquid) 17 1 209. Prepared by slowly adding Al.Cl. (8 pts.) to a cooled mixture of PCL (50 pts.) and 21 times its volums of acetone; yield: 5 pts. It is decomposed by water into di-acetone-phosphinic acid C, H, PO, and HCl. It readily combines with 1 mol. of chlorine or bromine (Michaelis, B. 17, 1273; 18, 898).

ACETONE SULPHONIC ACID v. ACETONE. ACETONIC ACID v. a-Oxy-iso-butyrio Acid. DI-ACETONIC ALCOHOL v. ACETYL-BUTYL ALCOHOL

ACETONINE C₉H₁₈N₂. This base described by Städeler (A. 111, 277), Hlasiwetz (A. 76, 294), and Mulder (A. 168, 228), was found by Heints (A. 201, 102) to be a mixture of di-acetonamine, tri-acctonamine, and tri-acctone-di-amine.

ACETONINES. Bases obtained by dehydrating acetone-alcamines by conc. H.SO.

Ethylidene-di-acetonine C₈H₁,N i.e. CH:CH.CH₂ CH₂CH:CH or

MeCH.NH.CMo2 MeCH.NH.CMe. Tri-methyl-tetra-hydro-pyridine. (132°-137°.)
Formed from cthylidene-di-acctone-alcamine (1 pt.) and conc. H2SO4 (3 pts.) by heating for 11 hours at 100°.

Colourless oil with alkaline reaction. Volatile with steam. Sl. sol. water, more soluble in cold than in hot water. Miscible with alcohol, ether, or chloroform. Heated with 111, it yields iodotri-methyl-piperidino, C,H,Me,IN [60°],

Salts .- B'H1; slender, sparingly soluble needles .- B'HBr; small soluble trimetrio pyramids (Fischer, B. 17, 1795).

Benzylidene-di-acetonine C, H, N i.e. $\mathbf{H}_{2}\mathbf{C} - \mathbf{C}\mathbf{H} = \mathbf{C}\mathbf{H}$ $H\ddot{C} = CH - CH$ or

PhHC-NH-Olc PhHC-NH-CMe2 Phenyl-di-methyl-tetra-hydro-pyridine. Formed by the action of strong H2SO4 on benzylidenedi-acctone-alcamine.

Distils undecomposed. Volatile with steam. V. sol. alcohol or ether, v. sl. sol. water.

Salts. - *B'HBr: colourless tables or needles, sl. sol. cold water. B'H1; sparingly soluble needles.— B'HAuCl. (Fischer, B. 17, 1797.) Tri-acetonine

H₂C.CH:CH C,H,N i.e. (146°) at 540 mm. Me.C.NH.CMe.

Tetra-methyl-tetra-hydro-pyridine. Prepared by heating tri-acetone-alcamine (1 pt.) with strong H.SO. (3 pts.) on a water-bath for an hour, pouring into water, neutralising the acid, and distilling the base over with steam

(Fischer, R. 16, 1604).

Properties.—Mobile fluid, smelling like piperidine. It combines with water forming a hydrate ridinc. It combines with water to the erystallising in long white codles, which give erystallising in long white codles, which give latile with steam. Poisonous. By heating with M it gives iodo-tetra-methyl-piperidine.c

Salts.—B'HBr; large white prisms, sl. sol. cold water.—B'HCl; v. sol. water or alcohol.— B'HAuCl, ; yellow needles.

Nitrosamine O, H, N(NO): yellowish tables; v. sol. alcohol, ether, and benzene, nearly insoluble in water; volatile with steam; strong camphor-like smell

Methyl derivative C.H. NMed colourless oil very volatile with steam, sparingly soluble in water (Fischer, B. 17, 1789).

ACETO-NITRANILIDE v. NITRO-ANILINE.

C2H3N i.e. CIL1.C: N. ACETONITRILE Methyl cyanide, methyl isocyanide. M.w. 41. (81.6°) at 760 mm. (Vincent a. Delaehanal, Bl. 33, 405); (81.3°) (R. Schiff, B. 19, 567). S. G. 2 865; 13 789 (V. a. D.). S. V. 5723 (S.). H. F. p. - 15,680. II. F. v. - 16,260 (Thomsen). μ_0 1·3458 (V. a. D.). R_{∞} 18·00 (Kauonnikoff, J. pr. [2] 31, 361). V.D. 145 (for 142).

Occurrence. - In crude benzenc (V. a. D.). Formation .- I. Dry KMeSO, is distilled with dry KCN and the distillate rectified over CaCl2 (Frankland a. Kolbe, C. S. Mem. 3, 386; A. 65, 288).—2. From Mo.SO, and KCN, the product being distilled over HgO and then over P.O. (Dumas, Malaguti a. Leblanc, C. R. 25, 474).-3. By distilling NH,OAc with P.O. (Dumas, C. R. 35, 383).-4. By distilling acctamide with P.O. (Buckton a. Holmann, C. J. 9, 212) .- 5. By distilling acetamido (5 mols.) with P₂S₃ (1 mol.), washing the product with NaOHAq and digesting with PbO (Henry, A. 152, 149). -6. From acetamide by action of PCl, (Wallach, A. 181, 21).

Preparation. -1. By boiling acetamide (500 g.) for a week with a little glacial acetic acid. the water produced being constantly allowed to distil off. The theoretical yield is got (Dodistil off.

marçay, Bl. [2] 33, 456).

Properties.—Colourless liquid with a pleasant ethereal odour; burns with a reddish-bordered flame. Miscible with water, but separated by salts from the solution. Mixes with alcohol. The presence of a little alcohol lowers its boilingpoint several degrees (D.).

Reactions .- I. Hot aqueous KOH acts thus : CH, CN + H2O + KOH = CH3CO2K + NH3.

2. Chromic and nitric acids have no action. 3. Heated with Na, it forms Cyanmethine (q. v.) and NaCN .- 4. Glacial HOAc at 200° forms diacetamide: $\mathrm{CH_{2}CN}+\mathrm{CH_{3}CO}.\mathrm{OH}=(\mathrm{CH_{2}CO}).\mathrm{NH}$ (Gautier, A. 150, 189).—5. Ac.O, forms triacetamide: $\mathrm{CH_{2}CN}+(\mathrm{CH_{2}CO}).\mathrm{O}-(\mathrm{CH_{2}CO}).\mathrm{A}.$ 6. Combines with dry HBr, HI, and (with difficulty) with HCl (Gautier, A. 112, 291) .- 7. Bromine forms the hydrobromido of the nitrile of Bromo-acetic acid (q, v.), CII, Br.CH:NBr [65°].

Combinations .- C.H. N2HBr, or CH, CH, NBr. [47°-56°] crystals; may be sublimed, -C.H. NPCl. (72°): dissociated above its boiling point (Hencke, A. 106, 281).-C.II. NSbCl., formed with great riso of temperature; white crystals which may be sublimed (IL). • C.H., NAUCl; brownish yellow powder (H.). • (C.H., N). TiCl; white crystallino orusts; may be sublimed (H.) .- (C.H.N) SnCl4: sublimes in arborescent formations (H.) .-C.H. N21Ig(CN): white vitreous mass; decomposes ovon over H.SQ. (Hesso, A. 110, 202), CH., C(NH.): NOH, formed by the union of aceto-nitrile with hydroxylamine v. ETHENYL-AMID-OXIM.

ACETONURAMIC ACID C.H., N.O. 2.6.

NH, CO.NH.CMe, COOH. a Uramido iso but fric acid, di-methyl-hydantoic acid. Obtained, as barium salt (C.H., N.O.), Ba(OH),, by prolonged boiling of a solution of di-methyl-hydantoin with

baryta-water. The acid itself appears to be very unstable (Urech, A. 164, 255). A more stable acid of the same composition is obtained by evaporating the mixed solutions of the sulphate of amido-isobutyric acid and potassium oyanato. It forms crystals, melting, with loss of water, at 160°, modorately soluble in hot water and aloohol. Decomposed by prolonged heating at 130°-140° into water and di-mothyl-hydantoin. C, H, AgN, O, crystallisos in needles (Urech, A. H.W.

ACETONYL-ACETO-ACETIC ETHER C.H.O. i.e. CH, CO.CH, CHAc.CO.Et, a-B-di-acetyl pro-

pionic ether.

From aceto-acetic ether and ohloro-acetone (Weltner, B. 17, 67). Liquid. Warm cono. HCl changes it to pyrotritaric ether C,H,O,Et. Water at 160° produces some acetonyl-acetone.

ACETONYL-ACETONE C.H 1002 i.e. CH, CO.CH, CH, CO.CH, Di-methylethylene di-

ketone. (188° uncor.)

Formation. -(1) By heating pyrotritario acid (di-methyl-furfurane-curboxylic acid) with water at 150°-160°; yield nearly theoretical. (2) By heating acetonyl-aceto-acetic ether with water at about 160°; small yield.

Properties. - Mobile liquid of peculiar smell. Miscible with water, alcohol, and ether, insol.

conc. KOHAq, or K.CO₄Aq.

Reactions. -P.S., when heated with it, forms thioxeno C.H.S.-Heating with alcoholio NH, gives di-methyl-pyrrol (Paal, B. 18, 2251); amines beliave similarly (Paul a. Schneider, B. 19, 3156).

Di-oxim CH, C(NOH). CH, CH, C(NOH). CH, [135°]. White glistening plates, v. sol. hot water, alcohol, or other, v. sl. sol. benzone.

Di-phenyl-di-hydrazide C.H. (CMe:N2HPh)2 [120]: plates, v. sol. alcohol, ether, or benzene,

nearly insol. light petroleum (Paal, B. 18, 58).
ACETONYL-CARBAMATE C₃H,NO₃ [76°]. Formed by boiling acetonyl thiocarbamate (infra) with lead acctate or silver oxide. Crystallises from water in prisms, may be distilled; dissolves in water, alcohol, and ether, Decomposed by heating with strong hydrochloric acid or baryta-water, yielding CO₂, NH₂, and α-oxy-iso-butyric acid. The salts C₂H₂AgNO₂, and AgNO, 2C, H, NO, are crystalline (Urech, B. 11, 467; 13, 485).

ACETONYL-PHOSPHINIC ACID C₃H₁PO₃ i.e.

CH, CO.CH, P(OH), Residue left after distilling acctone with I and P (v. Acctone-phosphorous acid under ACETONE). Salt. -Ba(C.H.PO.)2.

Di-acetonyl phosphinio Aoid v. Dr-AOETONE-PHOSPHINIO ACID

Di-acetonyl-phosphorous Chloride v. Di-ace-

TONE-PHOSPHOROUS CHLORIDE. ACETONYL-QUINOLINE C12H11NO CH:CH

CuH4 N :C-CH2.CO.CH2 Quinolyl-acctone. [76°]. Prepared by reducing o nitro cinnamoyl-acetono in alcoholio solution with SnCl. Long yellow noedles. Distils without decomposition. Sparingly volatile with Bycs wool and silk yellow. Heated with strong HCl at 170° it gives (Py. 3)-methyl-quinoline (Fischer a, Kuzel, B. 16, 163).

ACETONYL-THIO-CARBAMATE (so called) Thiacetonuramic acid. [152°]. C.H.NSO.

Marie Committee of the second

Formed by treating acetone with a mixture of potassinm cyanide and enlphooyanide, and HCl: C₂H₂O + CNH + CNSH + H₂O = NH₂ + C₄H₂NSO₂. Long needlee; easily eublimable; very coluble in ether; less easily in cold water. Resolved by heating with HČl in a sealed tube at 120° into CO, H,S, NH, and a-oxy-ico-butyric soid. The silver ealt C.H. AgNSO, ie very sparingly soluble (Urech. B. 6. 1117).

H. W.

scinble (Urech, B. 6, 1117). H. W. ACETONYL-UREA v. di-METHYL HYDANTOÏN. ACETO-PHENINE v. ACETOPHENONE, Reaction 6

ACETO-PHENONE C,H,O i.c. C,H,.CO.CH, Phenyl methyl ketone, Acetyl-benzene. 120. [20·5°]. (202° eor.). S. G. 15 1·032.

Formation .- 1. By distilling calcium benzoate with calcium acetate (Friedel, A. 108 122) .- 2. From BzCl and ZnMe, (Popoff, A. 161, 296).—3. By action of KOHAq on benzoyl-accto-acctic ether.—4. From phenylacetylene by shaking with diluted (75 p.e.) H₂SO₄ (Friedel a. Balsohn, Bl. [2] 35, 54): Ph.C:CII+II.O = Ph.CO.CH₃.—5. From bromo-etyrene and H.SO₄: small yield.—6. Bromostyrene heated with a large excess of water for 12 houre at 180° yields 66 per cent. (Friedel a. Balsohn, Bl. [2] 32, 613).—7. From ethylbenzene and chromic acid in acetic acid (F. a. B.) .- 8. From di-bromo-phenyl-propionic acid, CH.Br.CBrI'h.CO.H by boiling water (Fittig a. Wurster, A. 195, 160).

Preparation .- From benzene (10 pts.), neetyl chloride (1 pt.), and AlCl3 (2 pts.) (Richter).

Properties .- Large plates. Does not combine with NaHSO2 but, like other ketones, it reacts with hydroxylamine, phenyl-hydrazine, and HCN

Reactions.-1. Chromic-mixture oxidises it to benzoic and carbonic acids (Popoff).-2. Sodiumamalgam reduces it to phenyl methyl carbinol, CH, CHPh.OH, and acetophenone-pinacone. 3. With HI and P at 140° it gives di-phenyl-dimethyl-ethane, CicHis, and a compound CisHisO (Graebe, B. 7, 1626; v. Acetophenone-pinaco-LINE).—4. Chlorine produces chloro-acctophenone Ph.CO.CH, Cl [59°] (245°) and di-chloro-acctophenone Ph.CO.CHCl, (250°. 255°) v. Chloro-ACETOPHENONE. -5. Bromine in CS. produces bromo-acetophenone, Ph.CO.CH Br [50°] v. Bro-MO-ACETOPHENONE. - 6. Ammonia in presence of P₂O₃ forms 'acetophenine' C₂H₁₇N together with methans. Accorphenine crystallises from alcohol in slender needles, which may be sublimed. It is a weak base; its hydrochloride crystallises in platee, decomposed by water into HCl and the baco. Finning HNO, forms tri-nitro-acetophenine C23H14(NO2)3N; slender needles (from ether). Acetopheniue is probably tri-phenyl-pyridine:

CPh:CH. [≿]СРЬ.СП СРЬ

(Engler a. Rielun, B. 19, 40) .- 7. When taken internally it reappears in the urine as hippurio acid, having, doubtless, been previously oxidised as in Reaction 1 (M. Nencki, J. pr. 123, 288).

Besides the derivativee described below, see aleo: Amido-Acetophenone, Bromo-acetophenone. BROMO NITRO-ACETOPHENONE, IODO-ACETOPHENONE, I)1-METHYL-AMIDO-ACETOPHENONE, NITRO-ACETO-PHENONE, THIO-ACETOPHENONE, ACETOPHEN-OXIM.

Acetophenone - Aceto - Acetic C₁₂H₁₂O₄ i.e. CH₃.CO.CH(CO₂H).CH₂CO.C₄H₃ Acetophenone-acetone-carboxylicacid[130°-140°]. Small colourless crystals. Obtained by saponifica-tion of the ether which is prepared by the action of ω-bromo-acetophenone on sodio-aceto-acetic ether. It is very anstable. On warming with absolute alcohol it evolvee CO, and yields acetophenone - aeetone CH, CO. CH, CH, CO.C, H, (Paal, B. 16, 2865). Acetophenono-aceto-acetio ether is reduced by sodium amalgam to a oily lactone, CH3.CH.CH2.CH(CHMeOH).CO.O, col.

aqueous KOH or Ba(OII), but insol. aqueous K.CO. (Weltner, B. 17, 69). Amines convert acetophenone aceto-acetic ether into derivatives of pyrrol (Paal a. Selmeider, B. 19, 3156).

Dehydro - acet ophenone - aceto - acetic C12H10O2 [114°]. From acetophenone-aceto-acetic ether C_sH_s.CO.CH_s.CH(CO_sEt).CO.CH_s by heating with alcoholio KOH. Large cryetals (from benzene mixed with benzeline). From dilute alcohol it separatee in hydrated needles [115°-120°]. By boiling with HCl it is converted into phonyl-methyl-furfuranc-earboxylio aoid

HC - C(CO2H) PhC-O-CMe.

Salts.-KA'. Long eilky needlee (from alcohol). NH,A'.

Oxim C12H12N2O3: [172°]; glietoning white plates; sparingly solublo in water, easily in alcohol, ether, benzene, aqueous acids, and alkalie.

Phenyl-hydrazide $C_{12}II_{16}N_2O_2$: emall needlee (Paul, B. 17, 916, 2761).

ACETOPHENONE - ACETONE C11H12O2 i.e. CII .. CO.CH2.CH2.CO.C4II3 (acctyl-benzoyl-ethane or ethylene methyl phenyl di-ketone). Prepared by heating acetophenone-aceto-acetic acid (q.v.)with absolute alcohol. Yellowish heavy oil. Sl. sol. water, quite insol. alkalis. Cannot be distilled (Paal, B. 16, 2868).

Reactions.-1. P2O5 removes II.O forming CH--CH phenyl-methyl-furfurane,

MeČ.O.ČPh, 2. Heated with P2S3 it gives, similarly, phenyl-

3. Heated with alcoholic NII, it gives, eimi-CH -CH larly, phenyl-methyl-pyrrol, MeC.NH.CPh

(Paal, B. 18, 367).

Oxim C₁₁H₁₃NO₂. [123°]. Formed by action of hydroxylamine. Long white needles, soluble in acids or alkalie.

Phenyl-hydrazide C1,H1,N2O [c. 105°]. White prisms, got by adding phenyl-hydrazine slowly to a solution of the ketone in ether (3 vole.). V. sol. ether or bonzene, nearly insoluble in light petroleum (Paal, B. 17, 2763)

C1, II16N2 [155°]. Formed by mixing the ketone with phenyl-hydrazine (cf. Knorr, B.18, 305). Dehydro-acetophenon-acetone [83°]. Thie body ie formed together with the isomoric phenyl-methyl-furfurant by the action of Ac,O and other dehydrating agents upon acetophenone-acetone. Cannot be dietilled, even with etsam. It combines with bromine, and

gives with phenyl-hydrazine the sams compound,

C,H,N, [155°], that the acetophenone-acetone itself gives. Hence its constitution must be something like CH, CO.CH, C:C.C, H

ACETOPHENONE ALCOHOL. C.H.O. i.e. C.H., CO.CH., OH v. Benzoyl-cardinol. An isomerio body, C.H. (OH). CO.CH., is described as CXY-ACETOPHENONE.

ACETOPHENONE - ANILIDE PHENYL-10.

AMIDO-ACETOPHENONE

ACETOPHENONE-BENZIL C22H18O3 [102°]. Acetophenons and powdered benzil in equivalent proportions are shaken with an excess of cone. KOH (S. G. 1.27). After a few days a solid oake is formed, which is washed with water and then treated with ether. This leaves dehydroacetophenone-benzil undissolved, and on evaporation deposits oblique prisms of acetophenone-benzil, which should be recrystallised from alcohol. It is v. sol. ether or hot alcohol, sl. sol. cold alcohol. Above its molting-point it gives off acetophenone. Its constitution is probably Ph.CO.CPh(OH).CH2.CO.Ph (Japp a. Millsr, Č. J. 47, 34).

Dshydro-acetophenone bonzil C22H16O2 [129°]. Formation.—See above.

Preparation .- Equivalent quantities of acetophenone are shaken with excess of cono. KOH (S.G. 1.27) and kept liquid for some hours by the

application of sufficient heat. The product is treated as described above, but ether extracts hardly anything. The residue insoluble in ether is orystallised from alcohol (J. a. M.)

 $\begin{array}{c} C_{11}H_{10}O_2+C_{11}H_0O=C_{22}H_{16}O_2+H_2O,\\ Properties.-Tufts of flat needles (from alcohol). V. sl. sol. ether or cold alcohol, v. sol. \end{array}$

boiling alcohol.

Reactions .- Bromine added to its solution in chloroform unites forming large reddish crystals which are apparently the tetrabromido C₂H₁₆O₂Br₄ [110°-115°]. Becomes dark at 70°, and pale again at 80°. The bromine is given off in a fsw weeks over lime.

Constitution. - Dehydro-acetophenone-benzil differs from dehydro-acotone benzil not only in forming a bromido but also in having a very much lower melting-point than would be expected if they were of analogous structure. Japp a. Miller assign to dehydro-acctone-benzil the formula Ph.CO.CPh CH CO, and the unsatu-

rated formula Ph.CO.CPh:CH.CO.Ph to dehydroacetophsnone-bonzil. The latter formula can. however, account only for a di- and not for a tetra-bronide (C. J. 47, 37).

ACBTOPHÈNONE CARBOXYLIC ACIDS. Acetophsnone w-Carboxylio Acid C,H,O, C.H. CO.CH 2.CO.H v. BENZOYL-ACETIC ACID.

Acetophenone o-Carboxylio Acid [1:2] CO.H.C.H. CO.CH. o-Acetyl-benzoic

Formation .- 1. Together with CO2, from acetophsnons di-carboxylio acid by heating, long boiling with water, or by potash-fusion.—2. By heating phthalyl-acetic acid with water at 200°:

CO.H.C.H.,CO.CH. + CO. (Gabriel a. Michael, B. 10, 1554).—8. From methylene-phthalide by warming with aqueous KOH (Gabriel, B. 17, 2524);

$$\mathbf{C}_{\delta}\mathbf{H}_{\delta} \underbrace{\mathbf{C} = \mathbf{C}\mathbf{H}_{\delta}}_{\mathbf{C}\mathbf{O},\mathbf{O}} + \mathbf{H}_{2}\mathbf{Q} = \mathbf{C}\mathbf{O}_{2}\mathbf{H}.\mathbf{C}_{\delta}\mathbf{H}_{\delta}.\mathbf{C}\mathbf{O}.\mathbf{C}\mathbf{H}_{2}.$$

Properties.—Broad orystals, with sweet taste. Reactions .- 1. Bromine and glacial acotic acid at 100° convert it into bromo-methylens-

C = CHBrphthalide C.H. 0.00

2. Cone. H_uSO_4 forms, in the cold, two bodies, $C_{18}H_{12}O_4$ [216°] and $C_{18}H_{11}O_5$ [0.194°]. The latter body is a monobasic acid, discetophenono carboxylic acid, and splits up into CO, and the former body when it is heated above its molting-point (W. Roser, B. 17, 2620; Gabriel, B. 17, 2665). -3. Aleoholie NH3 for fourteen hours at 100° forms a base, C15H17N3O2 [204°-210°] It crystallises in long needles, insol. water or alcohol and gives a nitroso-derivative [246°] (Gabriel, B. 18, 1258).—4. Ac.O and NaOAc form Ac.C. H. CO.Ao [71°]; needles, insol. alkalis (Gabriel, B. 14, 921).

Phenyl-hydrazide

CO2H.C.H. CMc(N2HPh) Small prisms or large tables, v. e. sol. alcohol (Roser, B. 18, 804).

Oxim .- The anhydride of this body,

C.H. < CMe; N

[159°] is formed by the action of hydroxylamine (base) on acetophenone-o-carboxylic ether, or of hydroxylamine hydrochlorido upon acetophenone di-carboxylic other. It is also got, together with CO, when the oxim of acetophenone di-car-boxylic acid is heated. It crystallises in colourless needles (Gabriel, B. 16, 1993).

Acstophenone p-Carboxylio Acid C.H.O. [1:4] CO.H.C.H., CO.CH., [200°]. Formed, together with terephthalic acid, by warming exooxy-isopropyl-benzoio acid Ms2C(OH).C.H4.CO2H with chronic mixture (R. Meyer, B. 12, 1071. A. 219, 259). The process is similar to that by which tri-methyl-carbinol is converted into acctone. Needles (from water). May be sublimed. V. sl. sol. cold water, sl. sol. hot water, alcohol or ether.

Salts. BaA'2 aq. CuA'aq. PbA'21 aq.

Methyl ether. - MeA', [92°]. Small needles. Acstophenone ο-ω-di-carboxylic acid

C10H2O3aq i.e. CO.H.C4H2CO.CH2CO2H aq. Benzoyl-acct-carboxylic acid. [90°]. Formed by dissolving phthalyl-acetic acid (q. v.) in cold aqueous NaOH and ppg. by HCl (Gabriel a. Michael, B. 10, 1553). It behaves, therefore, as if phthalyl-acetic acid were its anhydride. Broad needles (from water). On molting, it splits up into H2O, CO2 and acetophenone o-carboxylic acid (q. v.).

Salt: AgA': granular pp.

Phenyl-hydrazine, in alcoholic solution in presence of HOAc, forms the anhydride of the C—CH_s. C_{p} H phenyl-hydrazide: $C_{s}H_{s}C_{p}$ N₂Ph

It is soluble in NaOHAq and is reppd. by HCl. [160°] giving off CO₂. It forms salts, e.g.: (Cl. H₁₁N₂O₂)₂Ca3aq (W. Roser, B. 18, 808).

Hydroxylamine forms, in like manner, not

the oxim but its anhydride:

со.е.≫и со.е.≫и

[c. 150°]. This is a mono-basic soid, and splits up, when heated, into CO, and the anhydride of the oxim of acetophenone-o-carboxylio acid (above).

ACETOPHENONE CHLORIDE v. DI CHLORO-

ETHYL-BENZENE.

A1 9 1. 1. 16 16 18

ACETOPHENONE CYANHYDRIN C.H.NO, Ph.C(OH)(CN). Me a-oxy-a-phenyl-propionitrile, a-oxy-hydro-atropo-nitrile. Formed by mixing acetophenone with KCN, and adding fuming HCl (Spiegel, B. 14, 235). A brown oil.

Reactions.—1. KOHAq gives atrolactic acid, Ph.CMe(OH).CO₂H.—2. HCl at 130° gives chloro-hydro-atropic acid : Ph.CII(CH2Cl).CO.H (Spiegel, B. 14, 1352).—3. Ammonia forms Ph.C(NH₂)(CN).Me (Tiemann a. Köhler, B. 14, 1980).

ACETOPHENONE-DI-METHYL-ANILINE v. DI-METHYL-AMIDO-BENZYL PHENYL KETONE.

ACETOPHENONE DI - METHYL - HYDRA. ZIDE C₁₀H₁₄N₂, i.e. PhCMe:N₂Me₂(165°) at 190 mm. Formed from acetophenone and di-methylhydrazine at 100° (Riesenegger, B. 16, 663).

ACETOPHENONE NITRANILIDE v. NITRO-

PHENYL-AMIDO-ACETOPHENONE.

PHENYL-HYDRAZIDE ACETOPHENONE C1.H11N2 i.e. CPhMe:N.PhH [105°]. Formed by shaking acetophenone suspended in water with a solution of phenyl-hydrazine hydrochloride and sodium acetate (Fischer, B. 17, 576). Also by allowing a cone, alcoholic solution of phenylhydrazine and acetophenone to stand for a day (Riesenegger, B. 16, 661), or by heating the oxim with phenyl-hydrazine (Just, B. 19, 1206). Slender white needles or plates. V. sol. ether, sl. sol. water or cold alcohol.

ACETOPHENONE-(\beta)-PINACOLINE C16 H16O, i.e. Ph₂CMe.CO.CH₃ (?). [41°]. (310° uncorr.). Prepared by the action of zine and HCl on an alcoholie solution of acetophenoue. Rhombic prisms or short pillars. Soluble in CaHe, ether, acetio acid, hot alcohol, &c. By heating with soda-lime it gives HOAc and di-phenyl-methylmethane, Ph₂CII.CII₂. On reduction with III and P, it gives a hydrocarbon C₁₆H₁₈ [128°], which is apparently identical with the hydrocarbon formed by the action of Na on bromo-cthylbenzene: PhCHMe.CHMePh. CrO3 oxidises it to di-phenyl-propionie acid, CH3CPli_CO.H. It An isomeric acetophenone-pinacoline [70°] in ph43° i. V.) is formed when acetophenone is base. Fuyith HI and P at 140° (Gracbe, B. 7, C.3H, (NO, forms plates or tables (from alcohol). Acetopheninecked by AcCl; HI reduces it to the

C₁₀IL₁₈ (Thörner a. Zincke, B. 11

N QNE-PINACONE (Engler a. Richm, L. C.H. - C(OH) - CH. purie acid, having, dout is action of sodium or acid again leaction is actophenom in 123, 288). - prisms. V. sol.

Besides the derivatives descrit is split up on also: Amido-Acetophenone, Bromo-Ahenyl-methyl-BROMO NITRO-ACETOPHENONE, IODO-ACETVET it into DI-METHYL-AMIDO-ACETOPHENONE, NITTIGE.CO.CH. PHENONE, THIO-ACETOPHENONE, ACETOPHEN-

ACETOPHEN-OXIMO, H. NO. i.e. Ph. C. Me: NOH [59°]. Phenyl methyl kelovim. Formed by mixing alcoholio solutions of acetophenone and hydroxylamine; after 24 hours, the alcohol is distilled off, and the product Crystallised from water. It forms colourless silky needles. Volatile with steam; socuble in hot water, alcohol, ether, benzene, ehloroform, or benzoline. Soluble in acids and in alkalis.

ACETO-PROPIONIC ACID v. ACETYL-PRO-

PIONIC ACID

ACETO-SINAPIC ACID v. SINAPIO AOID. ACETO-SUCCINIC ACID v. AGETYL-SUCCINIO

ACID. ACETO-THIENONE v. THIENYL METHYL RETONE

ACETO-THIO-TOLUIDIDE v. THIO-ACETYL-TOLUIDINE.

ACETO-VALERIC ACID v. ACETYL-VALERIO

ACETOXIM C3H,NO, i.e. Me,C:NOH Dimethyl-ketoxim, acctone hydroxylamide [60°] (135° i. V.) at 730 mm. Prepared by leaving an aqueous solution of acctone mixed with hydroxylamine hydrochloride, neutralised with NaOH, to stand for 24 hours; and extracting with ether (V. Meyer a. Janny, B. 15, 1324).

Properties.—Colourless prisms; extremely volatile and smelling like chloral. Very soluble in water, alcohol, ether, or benzoline. Neutral to litmus. Ether extracts it from a neutral, but not from an acid or alkaline, solution. It is readily decomposed by boiling acids (even acetic) into acetone and hydroxylamine. Acid reducing agents have a like effect, but zinc dust and NaOH does not affect it.

B'HCl, white powder [c. 100°], very unstable, formed by passing HCl gas into a dry ethereal solution of acetogim. -CaH.NCNaOEt, crystalline scales, got by adding NaOEt to au

ethereal solution.

derivative Me₂C:N(OBz), [42°], Benzoy tsmall colourless tables, very soluble in alcohol and ether, slightly in water; formed by the action of benzoyl enloride on acetoxim.

Benzyl other Me.C:N(OC,H.), (c. 190° oily fluid, soluble in alcohol and ether, insoluble in water; formed by the action of benzyl chloride and sodium ethylate on acetoxim; on warming with aqueous ItCl, it is split up into benzyl-hydroxylamine (II,N.OC.H.) and acetone (danny, B. 16, 170).

ACETOX1M1C ACID C,H,N,O,, CH .C(NOH).CH(NOH). Nitroso-acetoxim, Dinitroso-propane [153°]. Formation .- (1) By the action of hydroxylamiuc on u-di-chlor-acetone (CH_.CO.CHCl_). - (2) By the action of hydroxylamine on nitroso-acctone (CH, CO.CH(NOH)) (Meyer a. Janny, B. 15, 1165). Small prisms. Soluble in alcohol, other, and hot water. Its alkaline solutions are colouriess.

ACETOXYL. Kolbe's Now used to denote C.II.3O2. Kolbe's name for ACETYL

ACET-TOLUIDE v. Acetyl-TOLUIDINE. ACET-TOLYL-IMID-TOLYL-AMIDE v. Tolyn-

ACETAMIDINE C.H,NO. ACETURIC ACID CH. (NHAe). CO.H (acetyl-glycocoll, acetyl-glycine, or acetamido-acetic acid). [2060]. S. (at (acetyl-glycocoll. 15°) 2.7.

Formation: 1. By heating glycocoll with

ia.O. 2. By heating glycocoll-silver with acetyl-hloride (Kraut a. Hartmann, A. 138, 99).

Long colourless orystals, readily soluble in tot water and in alcohol, inecluble in ether, hieroform and benzene.

Reactions.—Givee a red coloration with Fe₂Ol₄.

Reactions.—Offees rea coloration with regard-leadily saponified by boiling with acids or alkalis. Salts.—A'NH, aq: soluble needles or largo ables.—A'Ag: soluble plates.—A'₂Ba5aq: easily cluble needles.—A'₂Cu4½aq: blue trimetric risms, easily soluble in water and in alcohol.— L'HHClx: needles, decomposed by water.

Methyl ether .- A'Me, [59°], (254°) at 712 mm., ong colourless tables, easily soluble in water,

dcohol, and benzene, sparingly in ether.

Ethyl ether.—A'Et, [48°], (260°) at 712 mm.,

rimetric plates. Amide CH2(NHAc).CO.NH,-[137°], large colourless tables, soluble in water and alcohol, insoluble in ether (Curtius, B. 17, 1663).

ACETUREÏDE v. Acetyl-UREA.

ACET-XYLIDE v. Acetyl-Xylidine.

ACETYL C.H.O, CO.CII, COMe or Ac. The radicle of acetio acid, &c. The name Acetyl was formerly applied to the radicle C.H.. The prefix eacet-often indicates the radicle CH.. C; as in acctamidine; sometimes it is merely a contraction for acetyl, as in acet-xylide. The acetyl derivatives obtained by displacing H in OH or in N11 or in NH are described under the compounds from which they are derived by this displacement.

DI-ACETYL C₄H₂O₂ i.e. CH₃.CO.CO.CH₃. The exim, CH₄.C(NOH).C(NOH).CH₃, of this hypothetical body, called also di-methyl-glyoxim methyl-ethyl-acetoximic acid, is formed by adding hydroxylamine hydrochloride to an aqueous solution of methyl oximido-cthyl ketone, CH, CO.C(NOH).CH, Glittering needles (Schramm, B. 16, 180).

DI-ACETYL-ACETONE DI-CARBOXYLIC ACID v. Aceto-Acetic ether, Reaction 32.

ACETYL-ACETOPHENONE v. BENZOYL-ACE-

ACETYL-ACRYLIC ACED v. TETRIC ACID. DI-a,-a2-ACETYL-ADIPIC ACID C16H14O, i.e.

CO2H.CHAc.CH2.CH2.CHAc.CO2H. Di-ethyl-ether.-Et,A". Formed as a byproduct (20 p.o.) of the action of ethylene bromide upon sodio-aceto-acotio ether, and found in tho residue after distilling with steam. It is a thick colcurless oil; its alcoholio solution gives a dark reddish violet coloration with Fe,Cl,

Reactions.-1. It gives a tolorably stable disodio-derivative which, on treatment with iodine, yields the di-ethylic ether of di-acetyl-totramethylene di-oarboxylio-acid,

CH,CAo.CO,H ; an acid which orystallises (with CH.CAo.CO.H. 2aq in pearly scales [210°].—2. By conc. NH. di-acetyl-adipio ethor is converted into the ketonoimide, C_{1.}H_{2.}N₂O₄ [177°].—3. Phenyl-hydrazine forms the phenyl-hydrazide [145°], which readily splits off alcohol giving ethylene-di-methyl-dioxy-di-quinizine,,

4. By distillation, or on solution in conc. H2SO4, it loses H.O. giving an ether C. H.O. which probably has the constitution

CH CO. CH(CO,Et).CH,

The corresponding acft, [189°], forms a phenylhydrazide, C₁₀H₁₂O₄(N.PhH) [192°] (Perkm a. Obrembsky, B. 19, 2051).

ACETYL-AMIDO COMPOUNDS v. AMIDO COMPOUNDS.

ACETYL-BENZOIC ACID v. ACETOPHENONE CARBOXYLIO ACID.

ACETYL-BENZOYL-ETHANE v. ACETOPHE-

NONE-ACETONE. ACETYL-BENZOYL-ETHANE CARBOXYLIC

ACID v. ACETOPHENONE-ACETO-ACETIO ACID. ACETYL-BENZYL-SUCCINIC ETHER

C₁,II₂O₃i.e.CO₂Et.CAc(CH₂Ph) CII₂,CO₂Et(310°) S. G. $\frac{1}{10}$ 1088. Prepared by the action of benzyl chloride on a mixture of sodium ethylate and acetyl-succinic ether (Conrad, B. 11, 1058).

ACETYL BROMIDE C.H. OBr i.e. CH. CO.Br. Acetic bromide (81°). Formed by treating acetic acid with PBr, (Ritter, A. 95, 209). Prepared by gradually adding 240 g. bromine to a mixture of 90 g. glacial acetic acid and 33 g. amorphous phosphorns, and distilling when the action is complete (Gal, A. 129, 537). Hanriot (A. Ch. [5] 17, 83) uses 1 pt. phosphorus, 15 acetic acid and 40 bromine. Colourlese Colourlese fuming liquid. Heated with bromine at 100° in scaled tube, it yields bromacetyl-bromide C.H.BrO.Br, together with more highly brominated compounds, which may be separated by fractional distillation (Gal). On the action of bromino on C2H3OBr, see also Urech (B. 13, 1720; J. 1880, 386). H. W.

ACETYL-BUTANE-PHOSPHONIC ACID v. di-Acrtone-phosphinic acid.

ACETYL-BUTYL ALCOHOL C.H.,2O. Di-acetonic alcohol Cll3.CO.CH. CMe2.OH. (164°). S.G. 25 .931.

Preparation.-Acid oxalate of di-acetonamine ($\bar{1}$ pt.) is dissolved in water (3 pts.) and cooled to 5°, when it deposits some of the salt; solid KNO, (2 pts.) is slowly added, and the mixture kept cool for some days and then heated to 50° or 60°; the oily layer (mesityl oxide) is removed partly by distillation, partly by a tapfunnel; and the aqueous solution, neutralised with K,CO, is shaken with ether (Hcintz, A. 169, 114; 178, 342).

Properties .- Syrup, misoible with water, alcohol, or other, gives off hydrogen when treated with Na.

C,H,2O2 γ-Acetyl-n-butyl Alcohol CH3.CO.CH2.CH2.CH2.CH2OH. Methyl 8-oxy-nbutyl ketone. (155°) at 718 mm. S.G. 2 1 0148.
Formation.— 1. From brome-propyl-aceto-

acetic ether (50 g.) by boiling for an hour with water (50 g.) and HCl (20 g. of S.G. 1·18) (Lipp, B. 18, 3280).—2. From so-called tetra-methylene methyl ketone carboxylic acid by boiling with water (Perkin, jun., B. 19, 2557).

Properties.—Liquid with campbor-like emell v. sol. water, alcohol, and other; scarcely volatile with steam. It does not reduce Tehling's solution or ammoniacal AgNOs. Chromic mixture oxidises it to 8-acetyl-n-butyrio acid. Sodium amalgam reduces it to w-5-di-oxy-hexans.

Anhydride O.H.O is. CH. CH: CMe OH, CH. Oil. Formed by distilling the alcohol or the following acid.

Tetra-methylene methyl ketone carboxylic seid, C,H12O2, appears to be a carboxylio acid formed from the anhydride of acetyl-butyl alcohol: CH₂ C(CO₂H):CMc O.

Its ethyl ether C₂H₁₁O₃, (223°), M.M. 10·195, is formed by the action of trimeth clene bromide on

aceto-acetic ether (v. p. 24).

ACETYL-BUTYL BROMIDE C, H, BrO i.e. CH₂.CO.CH₂.CH₂.CH₂.CH₃.Br. (215°) at 718 mm. From the preceding acid, $C_1H_{16}O_3$, or from acetylbutyl alcohol by the action of HBr. Also formed by heating bromo-propyl-accto-acctic other with dilute acids. It is a colourless oil, v. sol. alcohol or ether, v. sl. sol. water; boiling water converts it into the alcohol (Lipp, B. 18, 3281; Perkin, B. 19, 2557)

ACETYL-BUTYRIC ACIDS C, H1,0O,

a-Acetyl-n-butyric acid CH₃.CH_{...}CHAc.CO_{...}H v. Ethyl-aceto-acetic acid under Aceto-Acetic Acid.

β-Acetyl-n-but yricacid CH, CHAc.CH, CO.H. [c.-12°]. (242°). Formed, together with its ether, by boiling a-acetyl-a-methyl-succinic ether, CO_Et.CMeAc.CH2.CO2Et, with HCl (Bischoff, A. 206, 331).

Very hygroscopio liquid. V. sol. water, alcohol, or ether. Oxidises in air. Hot dilute

HNO, forms pyrotartaric acid.

-ZnA', (at 100°): nodules (from Salts. — ZnA', (at 100°); nodules (from alcohol). The salts of the alkalis and alkaline earths are syrupy, the lead salt may be got as a vitreoue mass.

Ether.-EtA' (204° 205°).

~Acetyl-n-butyricacidCII,Ac.CII,,CH,,CO.H [13°]. (c. 275° i. V.). From sodium aceto-acetic ether and 8-iodopropionic cther (Fittig a. Wolff, A. 216, 127). Thick liquid. V. sol. water, alcohol, or ether. Solutions are acid and decompose Na, CO,. Forms a crystalline compound with water, CII3.C(OH)2.CH2.CH2.CH2.CO2H [35°-36°] which forms inonoclinic prisms,

a:b:c=.769:1:.885 $\beta=75^{\circ}$ 20'. Over H2SO4 it loses H2O, becoming liquid.

 $Salts. - Ca(C_sH_9O_s)_zaq. - Pb(C_sH_9O_s)_zaq.$ ZnA'2.-- AgA'.

Reactions .- Sodium amalgam reduces it to δ -Oxy-hexoic acid (q. v.).

a-Acetyl-iso-butyric acid (CH3)2CAc.CO.II v. di-Methyl-aceto-acetic acid under Aceto-acetic

B-Acetyl-iso-butyric acid CH.Ac.CMeH.CO.II (248°). Formed, together with its ether and CO, by boiling a acetyl-8-methyl-succinio ether, CO, Et. CHAc. CHMe. CO, Et, with IICl (Bischoff, A. 206, 319). It is a liquid. V. sol. water, alco-

hol, or ether. Turns brown in air. Dilute HNO3 torms pyrotartaric acid. The salts are amorphous. The silver salt deposits silver on warm-

ing its solution.

Ether.-EtA' (206°-208°). Oil. ACETYL-TRI-CARBALLYLIC ETHER

O14H22O2 i.e. CO2Et.CH2 CAc(CO2Et).CH2 CO2Et. From coloro acetic ether and sodium acetyl-succinio ether, CO₂Et.CH₂.CAcNa.CO₂Et (Miehle, A. 190, 323). It boils with much decomposition at 280°–300°. Boiling baryta water or cone. alcoholic KOH split it up completely into alcohols

acetic, and tricarballylic, acids.
ACETYL-CARBINOL C.H. O. i.e.
CH., CO.CH.OH. Pyruvyl alcohol, Oxy-acetone, Acetal.

Formation.—Cone. H., SO, dissolves \$\textit{\beta}\$-chlore allyl alcohol, CH,:CCl.CH,OH, giving off HCl the solution is diluted and distilled (Henry

Bl. 39,526).

Ethyl ether.—CH, CO.CH, OEt. (128°)
S.G. 12 92. Formed by heating proparall ether CII: C.CH, OEt, with water and HgBr, (Henry C. R. 93, 421). Colourless liquid with peculial odour and burning taste.

Acetyl derivative C3H3O(OAc). Colourless fluid. (172°) S.G. 11 1.053. Solublo in water. Prepared by heating potassium acotate with chloracetone. Also from propargyl acetate, water, and HgBr2. The alcohol has not been got by its sanonification. Readily reduces ammoniacal silver nitrate or Fehling's solution, the chief product of the exidation being lactic acid.

Benzoyl derivative C₃H₃O(OBz). needles. [24°]. Soluble in hot water, easily in alcohol and ether. Prepared by heating potassium benzoato with ohloracetone. Zincke, B. 13, 637.)

ACETYL CHLORIDE C.H.O.Cl, i.e. Ac.Cl. Acetic chloride. M.w. 78.5. (50.9° cor.) (Thorpe, Acetic chiornee, ht.w. 100 (507 500), [Ann. Brühl, A. 203, 14). S. G. 2 11377 (T.); 32 11051. C. E. (0°-10°) 001391; (0°-50°) 001504 $(0^{5}-10^{9})$ 001391; $(0^{5}-50^{9})$ 00150474'05 (T.). μ_{8} 1'3954. R_{∞} 26'82 (B.). H.F.p. 63,300 (Berthelot).

Formation.—1. From POCl₂ and potassio ace-tate: 3KOAc + POCl₂ = K₂PO₄ + 3AcCl (Gorhardt, A. Ch. [3] 37, 294).-2. Contained in the more volatile portions of the product of the action of chlorine on aldehydo (Wartz, A. Ch. [3] 49, 58). 3. By distilling glacial acetic acid with PCL: HOAc + PCl_s = AcCl + ClH + POCl_s (Ritter, A. 95, 209).

Preparation .- By distilling glacial acetic acid (61g.) with phosphorus tribiloride (93g.) (Béchamp, J. 1856, 427). The follows (Thorpe, C. J. 37, 186): The reaction is as

3HOAc + 2PCl, = 3AcCl + 3HCl + P.O. If more HOAc be used Ac2O is also formed. The action of PClsis therefore precisely like that of PCl, amounting to a displacement of O by Cl.; the molecule IICl.Ac, which might be expected to be formed, cannot hold together on account of the monovalent character of chlorine, and so splits up at once into HCl and ClAc. Under precisely similar conditions, alcohol, HOEt, gives HCl and ClEt.

Properties.—Colourless, fuming, mobile, and strongly refracting liquid. Its vapour strongly attacks the eyes and respiratory organs.

Reactions.-1. Violently acted on by water, with formation of HCl and acctic acid.-2. With ammonia it yields acctaniide AcCl + NH. HCl + AcNII. and with aniline in like manner. acetanilide, AcNHPh.-3. Distilled with potassium acetate or benzoate, it forms acetic or aceto-benzoic oxide: KOAc + AcCl = KCl + Ac2O; and KOBz + AcCl = KCl + AcOBz. Similarly with salts of other acids .- 4. With potassium hydrosulphide it yields acetic hydrosulphide or thioacctio acid, and with potentium wonosulphide it forms acetic sulphide or thioacetic anhydride, AcCl + KSII = KCl + AcSH; and 2AoCl + K₄S = 2KO+Ac,S (Jacquemin a. Vosselmann, A. R. 49, 371).—5. With potassium nitrite it gives of nitrosyl-chloride, and towards the end of the reaction NO, and on heating the residue to 150°,

acetic anhydride distils over: AcCl + KNO₂= NOCI+KOAo, and AcCI+KOAo=KCI+Ao₂O
[Armstrong, O. J. 26, 683].—6. Silver nitrate
sats: 2AcCI+ANO₂=AgCI+NO₂+CI+Ao₂O.
Similarly with other nitrats: Hg(NO₃)₂ Pb(NO_{s)2}, and KNO_s are attacked immediately; Ba(NO₂)₂ is not affected; OL(NO₂)₂ readily. With KNO₃, chlorine is first evolved, NO₂ only towards the end (Armstrong) .- 7. With succinic acid it yields acetic acid and succinic anhydride,

> $C_2H_4(COOH)_2 + CH_3.CO.Cl =$ HCI+CH,.COOH+C,H,(CO),O;

and it reacts in like manner with other dibasic dihydrio acids; viz., isodibromosuccinic, plithalie, diphonic, and camphoric acids; isosuccinic, ordinary dibromosuccinie, fumaric, and terophthalic acids are not attacked; cublimed anhydrous oxalic acid is rosolved into H.O.CO, and CO₂; benzoio acid yields benzoic chloride and acetio acid (Anschütz, B. 10, 325, 1881).-8. With titanic chloride, acctyl chloride forms the compound TiCl, AcCl (Bertrand, Bl. [2] 33, 403). Large transparent octahedral crystals [25°-30°] sol. in CS2.-9. Aluminium chloride reacts according to the equation $Al_*Cl_u + 4(CH_*, CO)Cl =$ •4HCl+2(CH_*, CO.CH; CO), Al_*Cl_d (Winogradoff, Bl. [2] 34, 325). The product is a solid, decomposed by water into CO2 and acctone .-10. PCl, at 190° forms chloro-acetyl chloride CH, Cl.CO.Cl (Samosadsky, Z. 1870, 105), and tri-chloro-acetyl chloride, CCl, CO.Cl (Hübner, A. 120, 330).-11. Zinc produces a brown mass whence alcohol extracts 'ac etylide,' C, H1004, which may be ppd. by water. Red plates (from chloroform). Sol. ether, alcohol, HClAq, fuming IINO₃₁ or Ac₂O. Combines with bromine; does not reduce Fehling's colntion (Tommasi a. Quesneville, C. R., 76, 496).—12. Acts upon benzene, in presence of AlCl., with formation of acetophonone (q. v.) .- 13. Acts similarly upon thiophene, or its mono-haloid derivatives, displacing, in presence of AlCl, H by Ac. But in di-bromo- or di-iodo-thiophene it displaces, in presence of AlCl, Br or I by Ae, c.g.:

 $C_4H_2SBr_2 + ClAo = C_4H_2SBrAo + ClBr$

(Gattermann a. Römer, B. 19, 688). H. W. Use in Organic Investigations.—Acctyl chloride evolves HCl when it is heated with any substance containing the radicles hydroxyl, amidogen, or imidogen. Henco, if a substance does not evolve HCl when so treated, it may be assumed to be free from these radicles. If the hydroxyl be alcoholic, i.e. attached to an atom of oarbon that is not attached to any more oxygen, it will be converted into acetoxyl (AcO); and, if the substance contains no nitrogen, the number of acetoxyls it contains after this treatment gives the number of alcoholic hydroxyls the body contains. Before making the experiment, all carboxyls should be etherified, since the group CO.H is attacked by AcCl (e. Reaction 7), while the group CO2Et is not attacked. Tho number of acetyl groups that have entered may in many cases be determined by boiling with standard alkali and subsequent titration (Schiff). Acetyl chloride converts NH₂ into NHAc, but hardly ever into NAc₂. It converts NH into NAc. It has no action upon tertiary amines, hence it oan be used in the diagnosis of bases. Acetyl chloride does not set upon hydrogen directly hydrogen and gaseous or vaporous carbon com-

nnited to oarbon, except in presence of AlCl, or some similar agent.

ACETYL CYANDE C.H.ON or Pyruvo-nitrile. M. w. 89. (93°). V.D. 2.4.

Preparation .- When acetyl ohloride and silver cyanide are heated together in a sealed tube at 100°, and the product is distilled, a colourless liquid passes over at 80°-90°, and afterwards a compound having a much higher boiling-point. The first yields acetyl eyanide on rectification.

Properties .- Oil, lighter than water, which gradually dissolves it, forming HCN and HOAe, converted by HCl first into CH₃.CO.CONH₂, and subsequently into pyruvic acid (Hübner, A. 120, 230; 123, 271; see also Fileti, G. 5, 391; J. 1875, 510).

Di-acetyl-di-oyanide C.H.O.N. [69°] (210° V.D. 4.57 (for 4.77). Formed from acetyl eyanido by heating it with KOH, or even by keeping it for some time in a closed vessel.

Preparation .- Powdered KCN (32 pts.) is boiled with acctic anhydride (50 pts.), diluted with benzeno (200 pts.): yield is 25 p.c. of the theoretical (Kleeman, B. 18, 256). Glistening tables, sl. sol. hot water, v. sol. alcohol, ether, or benzene. Di-acetyl-di-cyanido, like acetyl cyanide, is converted by boiling with water, H₂SO₄, or KOH, into HCN and AcOH. Heated with

AgNO₄ it yields AgCN.

ACETYLENE C.H.₂ or CH.CH Ethine,
Ethinene. M.w. 26. Physical Properties of
liquid acetylene: S.G. 2 451; 12 420; 24 381.

C.E. (-7° to 36°) 00489. Vapour-pressure:
16,340 mm. at 0°; 24,900 mm. at 13.5°. Critical Point 37° (G. Ansdell, Pr. 29, 209). Properties of gaseous acetylene: V.D. 91. S. 1 at 18°; S. (CS2 or isopentane) 1; S. (CCl, or turpentine oil) 2; S. (amyl alcohol) 3½; S. (benzenc) 4; S. (glacial acetic acid or abs. alcohol) 6 (Berthelot, A. Ch. [4] 9, 425). H.F.p. -47,770. H.F.v.

-47,770 (Th.); -64,000 (Berthelot).

Occurrence.—In coal-gas (Boettger, A. 109,

351).

Formation.—1. Synthetically by passing hydrogen gas over charcoal heated to whiteness in the electric are (Berthelot, C. R. 54, 640); the hydrogen may be passed through holes drilled through the centre of carbon points discharging powerful sparks (Dowar, Pr. 29, 188). -2. By exposing marsh-gas or coal-gas to a strong heat, or to the spark of a powerful induction coil: $2CH_1 = C_2H_2 + 3H_2$ (Berthelot, C. R. 54, 515). Part of the C.H., is, however, polymerised during the process, being converted partly into benzene C, He, partly into black tarry hydrocarbons (Berthelot, Bl. [2] 11, 142). The vapours of many other organic compounds, as chiylene, alcohol, ether, acctone, amyl afcohol, and benzenc, likewise yilld acctylene when induction sparks are passed through them (De Wilde, Bl. [2] 6, 267).—3. By the incomplete combustion of hydrocarbons and other organic bodies—abundantly, for example, in a Bunsen lamp, when the flame strikes down and burns within the chimney—also in the incomplete oxidation of organic compounds at ordinary temperatures, as in the voltaio circuit, e.g. in the electrolysis of a solution of potassium aconitate or succinate (Berthelot, Bl. [2] 9, 103).— 4. By the incomplete combustion of mixtures of

pounds not containing hydrogen, e.g. CO, CS, CM.—5. By passing a mixture of methane and carbon monoxide through a red-hot tube; CH.,+CO=H.,O+C.H.,—6. Together with H. CH, and free carbon, by passing the vapour of methyl chloride (Berthelot), or of ethylene chloride (De Wilde), or of pentane from American petroloum (Vol.1, Bl. 4, 302), through a redhot tube .- 7. Together with benzene, by passing etyrene vapour through a red-hot tube: C_sH_s=C₂H₂+C_cH₄ (Berthelot, J. 1866, 511).— 8. By passing chloroform vapour over red-hot copper: 2CHCl₃ + Cu₆ = 3Cu₂Cl₂ + C₂H₂ (Berthelot), or by treating chloroform with potassiumamalgam (Kletzinsky, Z. 1866, 127), or with sodium (Fittig, ibid.). - 9. From iodoform by the action of finely divided silver either alone or mixed with finely divided copper: 2CHI₃+3Ag₂=6AgI+C₂H₃. Also by the action of finely divided zinc or of the zinc-copper couple on iodoform in presence of water (P. Cazeneuve, C. R. 97, 1371; Bl. (2) 41, 156). - 10. By passing a mixture of CO and HCl over red-hot magneeium silicide (Berthelot). - 11. By the action of alcoholic potash on bromethylene: $C_2H_2Br + KOH = KBr + H_2O + C_2H_2$ (Sawitsch, C. R. 52, 157). -12. Dy the action of water on calcium carbide (produced by strongly heating an alloy of Zn and Ca with charcoal): CaC2 + H2O - CaO · C2H2 (Wöhler, A. 124, 220). 13. Formed, together with succinic acid, by the electrolysis of sodium funnarate or malcate; $\mathbf{C_4H_2Na_2O_4} + \mathbf{H_2O} = \mathbf{C_2H_2} + \mathbf{2CO_2} + \mathbf{Na_2O} + \mathbf{H_2}$ (Kekulé, J. 1864, 389). -14. By heating is this onie acid with potash; C.H.O.SO, K+KOH = C, H, + K, SO, + 2H, O, -15. Formed in small quantity by heating cupric acctato (1 pt.) with water (200 pts.) in a closed flask at 100° (Tommasi, *Bl*. [2] 38, 257).

Preparation .- 1. Air is burned in a cylinder full of coal-gas, and a portion of the products of combustion are sucked (by an air-pump) first through a metallic condenser to cool them, and then through several bottles containing an ammoniaeal solution of cuprous chloride. A red pp., C.Cn.aq, is formed; this is collected, washed by decantation, and warmed with adheous HCl, when it is decomposed with evolution of acetylene gas: $C_aCu_aH_aO + 2HCl = C_aH_a + 2CuCl + H_aO$ (Jungfleisch, G.~R.~90, 264; J.~Ph.~[5]~1, 307).— 2. Ethylene bromide is slowly dropped into a strong alcoholic solution of potashat boiling heat, and the evolved gas is passed through a second eimilar boiling solution to remove bromethylene (Miasnikoff, A. 118, 330; Sawitsch, A. 119, 184; Sabanejeff, A. 178, 111). To remove the last traces of bromethylene, Zeisel (A. 191, 372) recommends passing the gas over moderately heated soda-line. The gas may also be purified, as in the first method, by passing it through an ammoniacal solution of cuprous chloride.

Properties.—Colourless gas, having a disagreeable odopt. According to Zeisel, when prepared from the copper compound as described above, it is contaminated with vinyl chloride. Acetylene is liquified by a pressure of 83 atmospheree at 18°, forming a mobile, highly refractive liquid, lighter than water. Liquid acetylene dissolves parallins and many fate (Cailletet, C. R. 85, 831).

Iteactions .- 1. Decomposed by the induction-

spark with separation of carbon, and partly converted into a liquid and a solid polyacetylene, the latter incoluble in the ordinary solvents.—
2. Slowly passed through a portel in tube heated to bright reduced it is almost wholly recoived into C and H, together with small quantities of ethylene and of the containing naphthalene. Acctylene is also recolved into C and H by exploding a percussion-cap in it (Berthelot, C. R. 93, 613). Heated to dull redness in a bent glass tube standing over mercury it is gradually polymerised, forming: a very volatile liquid probably C_sH_s , benzene C_sH_s , styrene C_sH_s (135°-160°), a liquid mixture (210°-250°) of naphthalene $C_{10}H_s$ and probably naphthalene hydride CioHio, & mixture of strongly fluorescent oils distilling at 250° 340°, retene distilling at 360° (Berthelot, C. R. 62, 905) .- 3. Mixed with excess of hydrogen over mercury, and in contact with platimm. black, acetylene is converted into ethane: C.H. + 211 = C.H.; by alkaline reducing agente, into ethylene, C.H., e.g. by the action of zino and aqueous ammonia on its copper compound (Berthelot).-4. With oxidising agents. Converted by KMnO, into oxalic acid, C.H.O., with formic and carbonic acids as secondary products (Berthelot, . C.R.74, 35). - 5. Slowly absorbed by an ammoniacal cupric solution, and for the most part oxidised, a carbonaceous substance being at the same time deposited, together with a small quantity of the compound C.Cu.ll.O (Berthelot, A. Ch. [4] 9, 122) .- 6. Passed with phosgene, COCl, through a red-hot tube, it is polymerised to beuzene (Berthelot, Bl. [2] 13, 9).-7. With chlorine, either pure or mixed with other gases, acetylene sometimes detonates, yielding HCl and free carbon. Frequently, however, C.H.Cl. is formed with explasion; or this compound is formed at first, and then the mixture suddenly explodes (Berthelot, Bl. [2] 5, 191). -8. Acetylene passed into bromine under water forms C.H.Br. (v. tetra-Baomo-ETHANE) and a non-volatile solid, a polymerido of C.HBr3 .- 9. Acetylene passed over iodine moistened with alcohol forms C.H.I. (Sabanejeff, A. 178, 109, v. di-Iodo-ETHYLENE).-10. Acetylene passed into a solution of ICl in HCl forms C.H.ICl (Plimpton, C. J. 41, 392, v. Chiloro-lodo-ETHYLENE). -11. With nitrogen. When a cerics of strong induction sparks is paesed through a mixture of acetylene and nitrogen, hydrocyanio acid is formed, Cally + N2 = 2HCN. Carbon and hydrogen are at the same time separated, but this may be prevented by diluting the gascoue mixture with 10 vol. H (Berthelot, C. R. 77. 1011).-12. Passed with vapour of hydrocyanic acid through a red-hot tube acetylene yielde a small quantity of picoline C.H.N, and probably homologues thereof (Ramsay, Ph. M. [5] 4, 241). 13. Strongly heated with butylene and amylene it forms C.H(C,H,) and C.H(C,H,) (Prunier, A. Ch. [5], 17, 5). -14. Successive troatment with H.SO, and water forms come etable culphonic acid (Zeiscl, A. 191, 366).—15. Converted into aldohyde by an aqueoue solution of mercuric oromide, even in the cold (Kurscheroff, B. 14, 1540):

 $CH_1CH + H_2O = CH_2:CH.OH = CH_1.CHO$.

16. Sec1, absorbs acetylene forming C₂H₂SeCi, which, on heating, splits up into SbCi, and C₂H₂Cl₂.—17. Cono. HBrAq at 100° forms a little bromo-ethylene (vinyl bromide).—18. Cone. HIE.

forms some lodo-ethylene and ethylidene iodide. 19. Passed through boiling sulphur, it forms some thiophene (V. Meyer, B. 16, 2176).

METALLIC REINVATIVES.—Sodium acety-lene C.HNa, is formed, with colution of hydro-gen and small quantities of cthylene and ethane, when sodium is gently heated in acetylene. At a dull red heat disodium-acetylene C.Na, is formed. - Potassium decomposes acetylene in like manner, but with greater violence; when melted in the gas it takes fire and is converted into C2K2, which is also formed when K is heated to dull redness on othylene-gas. All these compounds are decomposed by water with explosive violence and reproduction of acetylene (Berthelot, A. 139, 150).

Calcium-acetylene C.Ca is formed by strongly heating an alloy of zino and calcium with charcoal. Decomposed by water into

Ca(OH), and acetylene (Wöhler, A. 121, 220).
Copper-acetylene C,Cu,H,O. It may be looked upon as C.Cu.aq or as HC:C.Cu.Cu.OH; in the latter case it may be called cuproso-vingl hydroxide. Berthelot (A. 138, 315) considers it to be cuproso-vinyl oxide (C.HCn.).O. It constitutes the red precipitate formed on passing acetylene or coal-gas into an ammoniacal solution of enprous chloride. In the dry state it explodes when struck or when heated to 100 '-120°, leaving a velvety black powder containing copper and charcoal. Takes fire in contact with chlorine, bronnine, or finely divided iodine. Its formation affords a very delicate test for acetylene, the prescuce of 0.005 mg, of that compound being thus recognisable. The formula above given for it is due to Blochmann (A. 173, 174). According to Berthelot (Bl. [2] 5, 191) when acetylene is passed into a conc. solution of cuprous chloride in KCl a yollow crystalline pp. of cupreso-vinyl chloride C. HCu. Cl is formed, corresponding bromides and iodides being formed in a similar way.

Silver acetylene C.Ag.H.O, is formed on passing acetylene into an ammoniacal solution of silver nitrate, as a white or yellowish precipitate which, when dry, explodes even more easily than the copper-compound. The above formula, due to Blochmann, is that of argentovinyl hydroxide, CHIC.Ag.Ag.OH; Berthelot on the other hand regards the compound as the corresponding oxide (C.HAg.)O. The formula of Blochmann and Berthelot require 83-7 and 86.7 p.c. Ag respectively; Miasnikoff (A. 118, 832) finds 88 p.e. Ag in the pp., a result that has been confirmed by Plimpton, and agrees with the formula C.H.2Ag.O. Acetylene completely pps. the silver even from a nentral solution of AgNO3; the pp. contains variable quantities of AgNO, (Plimpton). The chloride C11: C.Ag.Ag.Cl, is prepared by passing acetylene into an annuoniscal solution of silver chloride (Berthelot). By agitating silver acetylene with a solution of iodine in other, till the colour of the liquid disappears and then evaporating, yellow offen-sive-smelling crystals are formed, the vapour of which strongly attaches the eyes (Berend, A. 135, 257); Baoyer (B. 18, 2275) has shown that they

are di-iode-acetyleno: C₁Ag₂ + 2I₂ = C₂I₂ + 2AgI.

Gold and Mercury Compounds.—In an ammoniacal solution of anrous thiosulphase, acc tylene forms a yellow highly explosive precipitate, and in an alkaline solution of potassio- succinic acid with alcoholic KOH (4 mols.)

mercuric iodide a yellow pp.: O.HHgI,H.O. which explodes slightly when heated and yields acetylene when treated with acids (Bassett, C. N., 19, 28). H. W

Theoretical considerations .- The explosive character of acetylene is undoubtedly connected with the fact that its formation from C and H is attended with disappearance of heat (Berthelot). It has been suggested by Baeyer (B. 18, 2277) that this disappearance of heat may be due to the production of a strained condition owing to the alteration in the direction of the attraction between the two carbon atoms.

To represent his views in a mechanical model, he supposes four steel wires fixed to a ball and radiating from it in the direction of the angles of an inscribed tetrahedron. Such a ball represents a free atom of earbon; union of such atoms is represented by a wire of one ball being attached to, and in a straight line with, a wire of another ball. If two such balls be taken and three of the wires from one ball be fastoned to three of the wires from another ball and then bent in such a way that all six wires are parallel, then the arrangement is in a strained condition. for the wires will readily fly apart, representing the explosion of acetylene. The anglo between two adjacent wires in one of the balls just described is 109° 28', which is very near the angle of a pentagou (108'); hence if five balls be placed at the angles of a peutagen, very little bending will be required to make a wire from each ball in a straight line with a wire from the next. The angles of a hexagon, of a square, and of an equilateral triangle, differ by 10½, 19½, and 49½ respectively from 109½; these numbers ought therefore to indicate the relative stability of rings containing 6, 4, and 3 carbon atoms. As a matter of fact, closed rings of carbon atoms usually contain five or six atoms, while rings containing three, four, or seven atoms are almost

Haloid derivatives of acetyleue v. Bromo-Chloro-, and Iodo-, ACETYLENE, -ETRYLENE, and ETHANE.

DIACETYLENE HC:C.C:CH. Gas of peculiar smell resembling dipropargyl. Formed by heating diacetyleno-di-carboxylie acid with ammoniacal Cu.Cl. solution. With ammoniacal Cu.Cl. it gives a violet-red pp., with ammoniacal AgNO₃ a very explosive yellow pp. By the action of a solution of iodine upon the silver-compound di-nodo-di-acetylene is formed. (Baeyer, B. 18, 2272.

ACETYLENE DI-BROMIDE v. DI-BROMO-ETHYLENE.

ACETYLENE TETRA-BROMIDE v. Tetra-BROMO-ETHANE

ACETYLENE DI-BROMIDE DI-CARBOXY LIC ACID v. DI-BROMO-FUMARIC ACID.

ACETYLENE BROMO-IODIDE v. BROMO-IODO-ETHYLENE. ACETYLENE CARBOXYLIC ACIDS

CH:C.CO.H. Acetylene mono-carboxylio acid v. Propiolio

Acetylene-di-carboxylic acid C,II,O,. i.e. CO,H.C:C.CO,H. Formed by treating di-bromo- or iso-di-brome-

100° (Bandrowski, B. 10, 838). The yield is 75 per cent. of the theoretical (Baeyer, B. 18, 677).
Separates from water in afflorescent orystals,

these contain aq., which they lose over H,SO,, and then crystallise from other in thick foursided tables. The hydrated acid is v. e. sol. water, alcohol or ether, but the dry acid is less soluble. The acid decomposes when melted.

Salts.—Na₂C₄O₄ 3 aq.: slender needles. KHC₄O₄: small crystals, sl. sol. water.— ZnC₄O₄1; aq. — PbC₄O₄aq. — CuC₄O₄3aq.: blue plates, el. sol. cold water (Bandrowski, B. 12,

2212).

Reactions .-- 1. The acid and its acid salts are converted, by heating with water, into propiolio acid: CO2H.C;C.CO2H = CO2H.C;CH + CO2. -2. Sodium amalgam reduces it to succinio aoid .- 3. Bromine combines forming di-bromofumario acid. -4. HCl, HBr, or H1 combine readily forming chloro-, bromo-, or iodo-, fumaric acids.

Methyl ether Me₂A". (197°). Colourless liquid (Bandrowski, B. 15, 2694).

Acetylene tetra-carboxylic acid, so called, v.

ETHANE TETRA-CARBOXYLIC ACID.

Di-acetylene di-carboxylic acid CeH2O4aq. i.e. CO2H.C:C.C:C.CO2H aq. Prepared by the action of a concentrated aqueous solution of potassio ferrioyanido upon a cooled magma of the ouprous compound of sodium propiolato:

or ether, v. sl. sol. benzene or benzoline. Turns brown at 100° and explodes very violently at s. 177°. Explodes also on percussion. Turned purple by light. Gives a brownish-red pp. with ammoniacal cuprous ehloride.

Reactions.-1. Sodium-amalgam reduces it to

hydro-muconic acid,

CO2H.CH:CH.CH2.CH2.CO2H,

and adipio acid, CO2H.CH2.CH2.CH2.CH2.CO2H; some propionio is formed at the same time. Zinc dust and HCl also roduce it to adipie acid.

Ether.—Et₂A" (184°) at 200 mm. An oil. Reduced by zinc dust and HCl to ethyl propiolate (Baeyer, B. 18, 678, 2269).

Tetra-acetylene di-carboxylio acid O10H2O4 i.e. CO2H.CiC.CiC.CiC.CiC.CO2H.

Preparation.-An aqueous solution of the acid sodium salt of di-acetylene di-carboxylie acid is heated on the water-bath. Sodium di-acetylene mono-carboxylate, H.C:C.C:C.CO2Na, is then formed, with evolution of CO2; the cuprous compound of this salt, Cu.C:C.C:C.CO2Na (?), is then prepared and this is oxidised by potassio ferricyanide:

 $2CuCiC_1C_1C_2C_2Na + O_a = 2CuO + (CiC_1C_1C_2C_2Na)_a$

(Baeyer, B. 18, 2271). Tetra-acetylone di-carboxylic acid may be reduced to sebacic acid, со,н.сн,си,сн,сн,си,оп,он,сн,си,со,н, by sodium-amalgam.

Icdo-acetylene carboxylic acid v. Icdo-Pro-

ACETYLENE DI-CHLORIDE v. DI-CHLORO-ETHYLENE.

ACETYLENE TETRA-CHLORIDE v. TETRA-CHLORO-ETHANE.

ACETYLENE CHLORO-BROMIDE v. CHLORO-BROMO-ETHYLENE

ACETYLENE CHLORO-IODIDE v. CHLORO-IODO-NTHYLENE.

ACETYLENE HYDROCHLORIDE v. CHLORO-ETHYLENE

ACETYLENE DI-HYDROCHLORIDE v. di-

CHLORO-ETHANE.
ACETYLENE GODIDE v. di-Iodo-ethylene. ACETYLENE NAPHTHALENE ACENAPHTHYLENE.

ACETYLENE-UREA O.H.N.O. i.e.

NH.CH.NH. CO. Glycoluril. NH.CH.NH/

S. 094 at 17°. Formed as white needles when cone. HCl is added to a solution of glyoxal (1 pt.) and urea (2 pts.) and water (3 pts.). If the filtrate is ovaporated, it deposits a yellow modifi-cation or impure form (Schiff, A. 189, 157; Böttinger, B. 11, 1787). Also formed by heating a mixture of tri-chlore-lactic acid, urea, and a little water at 100° (Pinner, B. 17, 1997). Formed when allantoin is reduced with (1 p. o.) sodium-amalgam (Reineck, A. 131, 119; Widman, B. 19,

Properties .- White glistening prisms, sol. hot water. Gives a white flocculent pp. with Hg (NO3)2: Split up by boiling baryta water into urca and hydantoic acid—C,H,Ag,N,O,.
ACETYL-ETHYL-PROPIONIC ACID v. ACE-

TYL-VALERIC ACID.

a-ACETYL - B - ETHYL . SUCCINIC ETHER C1211200 i.e. CO Et CHAC.CHEL.CO Et (263°) S. G. Tr. 1 vul. From aceto-acetio ether (68 g.), alcohol (120 g.), sodium (12 g.) and a-bromo-butyric ether (102 g.) (L. T. Thorne, C. J. 39, 336; S. Young, C. J. 43, 172).

Reactions.—1. Decomposed by potash (2 pts.)

and water (1 pt.) into acetic and ethyl-succinio acid.—2. Decomposed by potash (1 pt.) and water (20 pts.) into salts of CO. and \(\theta\)-acetyl-aethyl-propionic acid (v. Acetyl-valenc acid) .-3. With NaOEtand Melitgives a-acetyl-a-methyl- β -ethyl-succinic ether (q, v)—4. The other (3 pts.) boiled with conc. HCl (2 pts.) and water (4 pts.) for 2 days forms β-acetyl-α-ethyl-propionic acid, which may be extracted by other, and a crystalline acid, Ketolactonic acid (q. v.) which remains in the water. Ethyl-succinic acid is also formed .- 5. When heated, it partly splits up into alcohol and ketolactonic acid.

a.ACETYL.a.ETHYL.SUCCINIC ETHER C12H20O5 i.e. CO2Et.CAcEt.CH2.CO2Et (264°). From sodium acetyl-succinio ether and EtI (Huggenberg, A. 192, 146). Cono. alcoholio KOH converts it into cthyl-succinic acid.

DI-ACETYL-FUMARIC ETHER i.e. CO_Et.CAc:CAc.CO_Et [96°]. Formed the action of iodine (1 mol.) upon di-sodio-diacctyl-succinic ether (1 mol.) susponded in ether: CO_Et.CNaAc.CNaAc.CO_Et + I_2 =

CO.Et.CAc:CAo.CO.Et+Chal.

Long silky needles (Just, B. 18, 2636). a ACETYL-GLUTARIC ETHER C, H18O, CO2Et.CHAc.CH2.CH2CO2Et (272°) S.G. 1.0505. From aceto-acetic ether, \$\beta\$-iodo-propionio ether, benzene, and sodium (Wisligenus a. Limpach, A. 192, 180). Conc. alcoholio KOH splits it into acetic and glutaric acids. Boiling HCl forms CO2 and γ-acetyl-butyrio aoid.

B-Metyl-glutaric acid CHAc(CH2.CO2H), [109°]. Formed by heating a-carboxy-\$-acetylglutaric soid (from chlore-acetyl-propionic ether

and sodio-malonic ether) (Conrad a. Guthzeit, B. | by action of a-brome-propionic ether on sodium 19, 44).—AgA'. Di-acetyl-glutario

other C18H200 CO_Et.CHAo.CHAo.CH.CO_Et or CO_Et.CHAo.CH_CO_CHE CH_CO_Et. (c. 245) at 140 mm. From brome acetyl-pro-

pionicether, CH, CO.CHB.CH, CO, Et and sodioaceto-acetic ether (Knorr, B. 19, 47). Ammonia in HOAc converts it into di-methyl-pyrryl-acetic carboxylic ether

CMe:C.CH.,CO.Et CMo:C.CO,Et

ACETYL-GLYCOCOLL v. ACETURIC ACID.

ACETYLIDE v. ACETYL CHLORIDE, Reaction 11. ACETYL 10DIDE C.II.OI or AcI (108°) (G.); (I05°) (C.). S. G. 12 1.98 (O.). Prepared by the action of P and I upon acetic anhydride (Guthrie, P. M. [4] (1857) 14, 183) or on KOAe (Cahours (1857) C. R. 44, 1253). Also by heating acetyl chloride with Cal 3 aq at 75°. Tho water of crystallication has little saponifying effect (Spindler, A. 231, 272).

Properties .- Liquid. Fumes in air; pungent emell; cour taste. Water quickly forms Ill and HOAc; alcohol forms EtOAc. Decomposed by zine or sodium at 15°, and by mercury in sun-

ehine, forming HgI (G.). II. W. ACETYL-MALONIC ETHER C, II, IO, i.e. CII, CO.CII(CO, Et), (239°-215°) S.G. 24 1.080. From aceto-acetio cther, alcoholic NaOEt and CICO, Et. (Ehrlich, B.7, 892; Conrad, A. 214, 35). Aqueous NaOH decomposes it into acetone, CO., alcohol, and acetic acid.

ACETYL-METHYL-ACETO-ACETIC ETHER v. Aceto-acetic acid.

ACETYL-TETRA-METHYLENE v. TETRA-METHYLENE METHYL KETONE

ACETYL-TETRA-METHYLENE CARBOXY-LIC ACID, eo called, v. ACETYL-BUTYL ALCOHOL. ACETYL-TRI-METHYLENE v. tri-METHYL-

ENE METHYL KETONE a-ACETYL-a-METHYL-8-ETHYL - SUCCINIC

ETHER C13II.20, i.e. CH3.CO.CMe(CO.Et).CHEt.CO.Et. From α-acetyl-β-ethyl-succinic ether, NaOEt and MeI (S. Young, C. J. 43, 178). Boiled with dilute HCl it forms a y-oxy-octoic acid (q. v.) and methyl-ethyl-succinic acid.

ACETYL-METHYL-TRI-METHYLENE CAR-BOXYLIC ACID v. propylene-aceto-acetic acid under Ackro-Ackric Acid.

α-ACETYL-α-METHYL-GLUTARIC ACID

 $O_{12}H_{2a}O_{5}i.e.CO_{2}Et.CMeAc.CH_{2}CH_{2}.CO_{2}Et.(281°);$ S.G. 20 1043. From \$-iodo-propionic cther and sodium methyl-aceto-acetic other (Wislicenus a. Limpach, A. 192, 133). With conc. alcoholic KOH if gives acetic and a-methyl-glutaric acids.

ACETYL-METHYL-PYROTARTARIC ETHER v. Acetyl-di-Methyl-succinic ether.

a-ACETYL-a-METHYL-SUCCINIC ETHER C₁₁H₁₈O₃ i.e. CO_Et.CMcAc.CH₂CO_Et (e. 263°) S.G. 1 067. From sodium acetyl-succinic ether and MeI (Kressner, A. 192, 135). Decomposed by conc. alcoholic KAH with formation of acetic and pyro-tartario acide. Baryta-water or HCl produce CO, and β acetyl-butyric acid (q. v.).

• α. Acetyl-β-methyl-succinic ether CO_EL.CMeH.CA.cH.CO_Et (c. 268°)(C.); \$\oldsymbol{\phi}\$. 263°) (Gottetein, \$A\$. 216, 31); (c. 227°)eat 165 mm. (Bischoff, \$A\$. 206, 320). S.G. \$\oldsymbol{\text{T}}{\text{T}}\$. \$\oldsymbol{\text{T}}\$. \$\oldsymbol{\text{T}}\$. Tormed

aceto-acetic ether (Conrad, A. 188, 226). Decomposed by conc. KOH into alcohol, pyrotartario acid, acetic acid, CO, and β-acetyl-iso-

methyl-euccinic ether, codium, and McI (Hardtmuth, A. 192, 142). Boiling cone. alcoholic KOH converte it into acetic and a-8-di-methylsuccinic acide.

ACETYL OXIDE and Perceide v. Acerto

ACETYL-OXY-COMPOUNDS POUNDS.

ACETYL-PHENYLENE-DIAMINE v. PHENY-LENE-DI-AMINE.

8-ACETYL-a-PHENYL-PROPIONIC C11H12O3 i.e. CO2H.CHPh.CH2Ac. Benzyl-acetone-y-carboxylic acid. [126°]. Prepared by boiling the ethers of acetyl-phenyl-euccinic acid with baryta-water or dilute HCl. Plates. V. eol. alcohol or other. On reduction with codium. amalgam it gives the lactone of y-oxy-a-phenylvaleric acid, CH3.CH(OH).CH...CHPh.CO2H.

Salts. - * A', Zn: long white needles. - * AgA': white pp. - *CuA'₂: green; insol. water, sol. alcohol. - *CuA'₂ and *BaA'₂ are easily soluble (Weltner, B. 17, 72).

ACETYL-PHENYL-SUCCINIC ACID C,2H,2O i.e. CO.H.CHPh.CllAc.CO.H. [121°]. Formed by saponification of the di-ethyl-ether, which is prepared by the action of phenyl-bromo-acetic ether on sodio-acet-acetic ether. Large platea. (Weltuer, B. 17, 71.) When boiled with dilute hydrochloric acid or baryta water, it eplits off CO₂, giving acetyl-phenyl-propionic acid, C₂H₃.CII(CO₂H).CII₂.CO.CH₂.

A"K₂ easily soluble glistening needles.

(a)-Mono-athyl-ether C₁₄H₁₆O, i.e. CO_Et.CHPh.CHAe.CO_H. [133°]. Formed together with the di-ethyl ether by the action of phonyl-brom-acetic other upon sodio-aceto-acetic ether. On heating, it evolves CO, giving phenyllevulic ether (Weltner, B. 18, 790).

C, H,O, (B)-Mono-ethyl-ether CO.H.CHPh.CHAc.CO.Et. [128°]. White pearly plates; easily soluble in alcohol and ether. Formed by the action of sodium phenyl-bromeacetate upon sodio-aceto-acetic cther. By heating to 200° CO, is not split off. Boiled with baryta, it yields phenyl-levulic acid. It is reduced by sodium amalgam to a-phenyl-valeroγ-lactone-β-carboxylic acid:

CHPh.CO.O

CH(CO,IL).CH.CH,

With alcoholic NH, it yielde C12H12N2O2.

I'henyl-hydrazide C.,...H.,...N.,..O., [149°], plates.
Di-ethyl-ther A"Et., [70°], plates.
DI-ACETYL-PHOSPHORIC ACID G.H.,...PO,
i.e. H.,..Ac.,...PO, A viscid liquid, formed by the
action of Accl on Ag.,...PO, (Carine a. Kämmerer,
A. 121 120, D. Killing, and A. 121 120, D. 120, A. 131, 170). Boiling water decomposes it into acetic and ortho-phosphoric acide. It forms a calcium ealt, CaHAc,PO,2aq, cryetallising in ncedles.

ACETYL-PIPER - PROPYL - ALCEINE OXYPEOPYL-PIPERIDINE.

ACETYL-PROPIONIC ACIDS C.H.O. a-Acetyl-propionic Acid CH, CHAc.CO,H, v. Methyl-aceto-acetic acid under Aceto-Acetic Acid.

B. Acetyl-propionio Acid

CH, CO.CH, CH, CO.H. Levulic acid. (33.5°). (289°). S.G. B 1.135. $\mu_{\rm D}$ 1.443 at 15°. R_{∞} 45.3. Formation.—1. By spoilings acetyl-succinic ether with dilute HCl (Conrad, B. 11, 2177). 2. By boiling the following substances with very dilute H.SO,: Levulose, innlin, cane sugar (Grote a. Tollens, A. 175, 181), gnm arabic, or caragheen moss (Bente, B. 9, 1157). Filter paper and deal shavings give a small quantity. Small quantities may also be got from gincose, milk-sugar, and galactose, by heating with aqueous HCl.

Preparation .- Cane-sugar (1500 g.) is heated for 20 hours with water (1500 g.) and H₂SO₁ (150 g.), with occasional shaking. A large quantity of humie substance separates. The thick liquid is filtered under pressure, mixed with CaCOs (150g.), and the whole evaporated (till it weighs 1500 g.). The liquid is again filtered, mixed with H2SO4 (50 g.) and shaken with other. After evaporating the ether, the levulic acid (100 g.) is rectified (Grote, Kelmer a. Tollens, A. 206, 210). If glucose be used, the yield is not so good, and it is then better to use 1fCl (r. Conrad a. Guthzeit, B. 18, 112). Formic acid is also formed in these reactions:

 $\begin{array}{c} \textbf{C}_{a}\textbf{H}_{12}\textbf{O}_{b} = \textbf{C}_{b}\textbf{H}_{b}\textbf{O}_{a} + \textbf{C}\textbf{H}_{a}\textbf{O}_{c} + \textbf{H}_{b}\textbf{O}_{c}, \\ \textbf{\textit{Properties.}} + \textbf{Deliquescent} & trimetric & plates. \end{array}$ V. sol. water, alcohol or ether. On distillation it produces (a)- and (f)-angelico lactone, and also nectic acid, and another acid possibly C10 H10O3 [208°] (Wolff, A. 229, 260). Not attacked by bromine in the cold.

Reactions .- 1. Chromic mixture produces CO, and acetic acid. 2. Dilute HNO, gives succinic, oxalic, acetie, and hydrocyanic acids : (Tollens, B. 12, 334; A. 206, 257).-3. Reduce l, by P and IIIAq at 150 - 2000, or by sodiumamalgam in acid solution, to n-valeric acid.-4. In alkaline solution sodium amalgam produces γ -oxy-valeric, acid (q, v_*) . 5. Gives the iodoform reaction with NaOH and L.-6. Reacts with hydroxylamine, forming an oxfin.

Salts.—CaA', 2aq: minute needles. - AgA'; six-sided tables.—NaA': minute needles.—CaA', (at 150°); bluish-green flat needles or prisms. Barium, magnesium, and cadmium salts are

gummy. Ethers.—McA'. (191.5°). S.G. $^{\circ}_{6}$ 1.0681. μ_{0} 1.4216 dt 15°.R. $_{2}$ 52·2. – E.A'. (200.5°) (G.K.a.T.) (204°) (W.), S.G. $^{o}_{6}$ 1.0325. μ_{B} 1.421. R_{χ} 60.2. -PrA'. (215·5°). S.G. 6 1·0103. μ₀ 1·1246. R_∞ 69.5.

Amide,—C,H,O,NH, [108]. From ethyl levulate and alcoholic NH, or from (a)-angelicolactone (q. v.) and aqueous or alcoholic amnonia. Six-sided tables (from alcohol-chloroform, Wolff, A. 229, 260).

References .- See also Bromo- and Chroro-

ACETYL-PROPYL ALCOHOL C. H110O2 i.e. CH, CO.CII, CH, CH, OH. Methyl \(\gamma \)-oxy-propyl ketone. A colourless liquid, soluble in water formed by boiling bromo-curyl accto-acctic ether BrCH, CHAC, CO, Et with dilute HCl. 1t readily reduces ammoniacal AgNO, Aq but not Felding's solution. It is converted by heat into

an anhydride. Sodium-amalgam reduces it to w.y. di-oxy-n-pentane, CH₃.CH(OH).CH₂.CH₂.CH₂OH (Perkin jun. a. Freer, B. 19, 2566).

ACETYL-PYRO-PHOSPHOLIC ACID.
The barium talt, BaHAcP.O.2aq, is got as a orystalline pp., sl. sol. dilute acids, by adding aqueous hydrogen peroxide to a solution of barium acetyl-pyrophosphite (Menschutkin, A. 136, 254).

ACETYL - PYRO - PHOSPHOROUS ACID, AcHaPaO, 2aq, is got by heating AcCl with 1IaPO at 50° (Menschntkin, A. 133, 317). Crystalline mass.

Salts. - K. HAeP. O. 2! aq: slightly sol. water. BallAcl.O3: insol. water. - PbllAcl.O3: insol.

ACETYL-PYRO-TARTARIC ACID v. ACETYL-METHYL-SPECIAL ACID.

ACETYL-PYRROL v. Pynnol.

Pseudo-acetyl-pyrrol v. PYRRYL METNYL

ACETYL-SUCCINIC ETHER C1011 11005 i.e. CO_Et.CHAc.CH_.CO_Et. (c. 255°); (210° i. V.) at 330 mm. S.G. ²¹/_{17.5} 1 079; ¹⁵/_{17.5} 1 088; ³⁵/_{27.5} 1 080. M.M. 10 313 (Perkin, C. J. 45, 517). Formed by action of chloro-acetic ether upon sodium. aceto-acetic ether (Conrad, A. 188, 218). Oil. Sol, alcohol or ether. Gives no colour with FegCls. Cone. alcoholic KOH splits it into acetic and succinic acids; boiling baryta-water forms B-acetyl-propionic other and CO.

Phenyl-hydrazide Cull No. 0, [80°]. At 150° it splits off EtOH and gives methyl-oxy-quinizyl-acetic ether (Knorr a. Blank, B. 17, 2051),

Di-aceto-succivic Ether C121118O6

CO.Et.CHAc.CilAc.Co.Et. [79°].
Sodium acetacetic ether is treated in ethercal

solution with iodine (Rügheimer, B. 7, 892): 2CO_ELCHNa.CO.CH + I, = (CO_ELCHAe)a

The ethylic di-aceto-succinate crystallises from the other (Harrow, C. J. 33, 427). It forms trimetric tables, v. e. sol. alcohol, ether, or benzene. Reaction, -1. Boiled with dilute H.SO.

(1:10) it gives off GO2 and forms pyro-tritario or nyic other Colling, and carbo-pyro-tritario ether C.H.O.Et. - 2. Hydroxylamine forms a dioxim (needles; Münchmeyer, B. 19, 1849), and a neutral ether C12H1: NO , (Knorr, B. 18, 1568) .-5. Ammonia forms di-methyl-pyrrol di-carboxylie ether C.HMe.N(CO.Et), or

CMc:C.CO2Et CMe: C.CO₂Et.

Primary bases act in a similar manner, thus methylamine forms

,CMe:C.CO,Et NMe CMe:C.CO.Et

(Knorr, B. 18, 299). 4. Planyl-hydraxins acts in a similar way: $C_{12}H_{18}O_4 + N_2H_2C_6H_3 = C_{18}H_{12}N_2O_1 + 2H_2O_1$. The new compound, which N1h, CMc:C.CO,Et

may be NI1 . CMe:C.CO, Et

is called phenyl-di-meth pyridazine di-earboxy-lic ether. It contains 11,0 less than the monophenyl-hydrazide of di-acetyl-succinio ether, CO.Et.Clf(CMe:N.PhII).Cl1Ac.CO.Et (Knorr, B. 17, 3058; 18, 305). It crystallises in prisme, [127°] (from bonzoline). See also PHENYL-LLY DRAZINE.

ACETYL SULPHIDE OH,OS or Ac.S. Di-acetyl sulphide, Thio-acetic anhydride (120°). Preparation.—1. From Aco and P.S. (Ke-kulé, A. 90, 312) Yield 10 p.c.—2. From Acol and K.S.—8. By distilling Physic.

Properties.-An oil, slowly decomposed by

water into HOAo and HSAc.

Di-acetyl Di-sulphide C, H,O,S, or Ac,S, [21°] Formation. - 1. From KSAc and I (Kekulé a. Linnemann, A. 123, 279). - 2. From BaO, and Ao,S in ethereal solution (Beckmann, J. pr. 125, 465): $2Ac_2S + BaO_2 = Ac_2S_2 + Ba(OAc)_2 - 3$. By electrolysis of thio-acetic acid (Bunge, B. 3, 297).

Properties. - Crystallino. Insol. water, v. sol. alcohol or CS2. Decomposed by warm water or by alkalie forming thio acetic acid and eulphur.

Decomposed by distillation.

ACÊTYL SULPHOCYANIDE Callanso or (133°). S.G. 16 1·151. From CH. CO.S.Cy. AcCl and lead sulphoeyanido (Miquel, A.Ch. [5] 11, 295). Pungent liquid. Decomposed by water into HOAe and HSCN. Forms with NH, in etheroal colution a non-volatile liquid which dissolvee in water and gives a red colour with FegCls. ACETYL-THIO-UREA v. THIO-UREA.

ACETYL-TOLYLENE-DI-AMINE v. LENE-DI-AMINE.

ACETYL-UREA v. UREA.

ACETYL-VALERIC ACID C,H, O,.

a-Acetyl-n-valeric Acid v. n-monul-acetoacetic acid under Aceto-Acetic Acid.

a-Acetyl-s-iso-valeric Acid v. iso-propyl-acetoacetic acid under Aceto-Acetic Acid.

a-Acetyl-u-iso-valeric Acid v. methyl-ethylaceto-acetic acid under Aceto-Acetic ACID.

β-Acetyl-u-iso-valeric Acid

CH, Ac. CHEt. CO, H.

(250°<u>-</u>252°). Got by boiling a-acetyl-B-ethyleuccinic ether (q. 4) with dinte KOH (Thorne, C.J. 39, 340). Liquid, miscible with water, alcohol, and ether. Turns brown in air. It is gradually decomposed by heat into II.O and an

oil C, H₁₀O₂ (219°). S.G. 25 1·0221. Reaction.—HNO₃ oxides it to othyl-succinic acid.

Salts.-Gummy, coluble in water.

Ether.—EtA' (224°-226°). Lighter than water. ACHILLEA.--The Iva plant (A. Moschata) has been cliemically examined by v. Planta-Reichenau (A. 155, 145), who has extracted from it the following substances: 1. Ivain C24H22O5, obtained by distilling the dried herb (freed from the roots) with water to romove volatile oil, exhausting the dried residue with absolute alcohol, precipitating with lead acetate, removing excess of lead with H.S, and exhausting the evaporated recidno with acetic acid to removo achillein and moschatin. Ivain then remains as a dark yellow resinoue mass, insoluble in water, easily soluble in alcohol, yielding an inteneely bitter colution. — 2. Achillein C₂₀H₂₈N₂O₁₃ and Mocchatin C₂H₂₈NO, are obtained by distilling the herb, gathered before flowering, with water. oxhausting the concentrated filtrate with absolute shohol, evaporating off the alcohol and adding water, which throwe down mosohatin; and on treating the liquid filtered therefrom with Pb(OH), again filtering, remov-ing lead with H.S. and evaporating, Aclallein remains as a brown-red mass, vory soluble in water, less readily in alcohol, insoluble in ether; | ¿¿is (acid) and γεννάω (I produce), generalised

very bitter; not precipitated by lead salts. Resolved by prolonged boiling with dilute sulphuric acid into sugar and achilletin C,H,NO, a dark-brown powder, insoluble in water, vory slightly soluble in aloohol; not bitter. Moschatin C2, H2, NO, is pulvorulent, nearly insoluble in water, somewhat soluble in absolute alcohol; tastes bitter.

A. Ageratum, growing in Italy and Provence, yields an essential oil boiling at 165°-182°; sp. gr. 0.819 at 14° (De Luca, J. Ph. [4] 18, 105). H. W.

ACHROO-DEXTRIN v. Dextrin and Starch. ACIDIMETRY. The estimation of acids by volumetric methods. V. Analysis.

ACID-FORMING OXIDES. Same as ANHY-

DRIDES (q. v.).

ACIDS .- Salts of hydrogen. The word acid (ac, sharp; acere, to be sour; compare acetum, vinegar, δξύς, ύξος) was originally loosely applied to all sour liquids. The term cannot new be accurately defined; but it may be stated generally that an acid is a compound of hydrogen, which, when mixed with, or dissolved in, water, is capable of exchanging the whole, or a portion, of tho hydrogen it contains for a metal, with simultaneous" formation of water, by the action on the aqueous solution of the acid of a metallic exide or hydroxido.

HISTORY .- The corrosive action of acids, and their power of dissolving metals and other substances have been known from early times. Thus Geber, who lived during the eighth century, was acquainted with impure nitric and sulphurie acids, and described them under the name aqua dissolutiva. Paracelsus (16th contury), from whom the school of Iatro-chemiets sprang, held that the human body in health consisted of certain acid and alkaline principles which balanced each other, and that disease was due to a preponderance of one or other of these principles. He was the first to propound a theory to account for the properties common to all acids; he supposed that they all contained an acid principle, which conferred taste and solubility on all substances into which it entered. This theory was accepted by Becher (17th contury), who named the acid principle acidum primogenium; and he added that it consisted of a compound of earth and water, both of which he believed to be elemente. The distinctive properties of acids:-their solvent power, their power of changing the colour of certain vegetable tinctures, and the fact that they form nentral bodies with alkalis; -were catalogued by Boyle (17th century). Stahl, in 1723, adopted Becher'e theory, and endeavoured to prove that while acids were the basee of all ealine bodies, the principle of all acids was ellphuric acid. Stahl's view continued to find supporters for a long timo, but ite defects were at length perceived. Many of the supporters of the phlogistic theory held that inorganic acids, such as sulphuric and phosphoric acide, were simple substances; and that by their combinations with phlogiston they gave rise to bodies such as sulphur and phosphorus, which were then regarded as compounds, but which we now know to be elements. After the discovery of oxygen by Priestley and Scheele, Lavoisier, in naming that element from

the facts discovered by him, that many acid bodies are produced by the union of 'combustibles' with oxygen; and although it was pointed out by Berthollet in 1789 that sulphydric and prussic acids contain no oxygen, the view of Lavoisier generally prevailed until the researches of Davy, and of Gay-Lussac and Thenard, on muriatic and oxymuriatic acids (hydrochloric acid and chlorine) in 1810, and the discovery and examination of hydriodic acid, and the investigation of prussic acid by Gay-Lussac in 1814 and 1815, compelled chemists to recognise the existence of true acids containing no oxygen, and led to a distinction being drawn between acids which contained oxygen, and those which did not.

Lavoisier also regarded acids as binary oxygenated compounds; and he supposed that the water which must be present in order that an acid shall react on other bodies merely played the part of a solvent. This view was supported and extended by Berzelius, who taught that certain oxides are capable of uniting with each other to form 'ternary' compounds or salts, and that these salts are decomposed by electrolysis into their 'binary' constituents, which are an acid and a base. Berzelius therefore applied the term electronegative to that oxide which appeared at the positive electrode on electrolysis of a salt, and the term electropositive to that exide which separated at the negative electrode. nogative oxides he classed as acids, and the positive oxides as bases. This theory ignored the fact that water is associated with these oxides in their various reactions; and, moreover, it overlooked the evident analogy between acids containing oxygen and acids containing no oxygen, but formed by the union of tho halogens, or haloid groups, with hydrogen. . To restore this analogy, Davy proposed to abandon the old view that acids were compounds of certain elements with oxygen, and suggested that all acids, whether they contained oxygen or not, should be considered as compounds of hydrogen. Dulong supported Davy's view, and extended it; he regarded acids as compounds of hydrogen with elements such as Cl, I, S; or with radicles such as CN, NO, SO,. As it was at that time supposed that such radicles were capable of separate existence, and as Dulong's hypothesis involved the creation of a large number of hypothetical substances, this hypothesis did not meet with general support. It was reserved for Gerhardt, led by a study of organic substances, to prove that most acids, when vaporised, do not separate into an oxide and water, but pass into the state of vapour as a whole. From this it followed that hydrogen, replaceable by metals, must be a constituent of all true acids.

CHARACTERISTIC FEATURES OF ACIDS. - Bodies possessing properties corresponding with tho definition of an acid given at the beginning of this article always contain hydrogen in intimate combination with one or more of the following elements; fluorine, chlorine, bromine, iodine, oxygen, sulphur, selenion, tellurium, or certain groups of elements (e.g. eyanogen) of whicl? carbon is one (comp. Acids, Oscanic, p. 53). It is true that water is not accounted an acid, nor is it usual to include hydrogen dioxide | Moreover, on treatment of the acid chloride with

among the soids; yet if the definition of soid were strictly applied hydrogen dioxide would find a place in this class, for it has an acid reaction with test paper, and on addition, for example, of bagum hydroxide to a solution of β it in water, the reaction characteristic of acids takes place;-

 $Ba(OH)_2.8H_2O + H_2O_3 = BaO_2.8H_2O + 2H_2O.$ Again, the reactions of hydrogon sulphide, selenide, and telluride, with alkalis, would lead to their inclusion among acids. The nams acid must also be applied to most compounds of hydrogen and one of the elements above mentioned with a third element. The following examples will illustrate the definition given :-

Simple. HF. HBF = HF.BF HCl. $H_2PtCl_s = 2HCl.PtCl$ $H_sFe(CN)_s = 4HCN.Fe(CN)_s$ $H_sGO_s = H_sO.SO_s$ HCN. (H,O.) H.S. $\mathbf{H}_{2}\mathbf{CS}_{3} = \mathbf{H}_{2}\mathbf{S}.\mathbf{CS}_{2}.$ åс.

Such bodies as H₂ZnO₂ (Zn(OH)₂), and H₃AlO₂ (Al(OH)₃), may be classed either among acids or basic hydroxides, inasmuch as they possess the characteristics of both classes.

Among the compounds of carbon the acids form an important class. The formulæ of these compounds may be supposed to be derived from the formula either of formic acid, or of carbonic acid. If formic acid be taken as the type, then most acids containing carbon may be viewed as substituted formic acid; thus:

HCOOH. CH,COOH. C,H,(COOH). Succinic acid. Formic acid. Acetic acid. CaH,(OH)(COOII)a. Citric acid.

It is to be noticed that in two cases more than one molecule of formic scid is employed; and that succinic acid, by this view, is to be regarded as two molecules of formic acid, in which two atoms of hydrogen are replaced by the group C.H.; while citric acid is derived from three molecules of formic acid by replacement of three atoms of hydrogen by the group Calla(OH). The carboxylic acids may be similarly derived from carbonic acid (CO(OII),), if one hydroxyl group be regarded as replaced by an alkyl or similar group. But it is clear that unless this view of the composition of earbon acids helps to render prominent the actual relations existing between these compounds, it can be of no value. In this view of the constitution of carbon acids these compounds are all represented as containing the characteristic group CO OH; this group has been named 'carboxyl,' a word derived from 'carbonyl,' CO, and 'hydroxyl,' OH, and implying the presence of these two groups. That most of the acids of carbon contain the group CO.OH is rendered probable by the following considerations: when one of these acids is distilled with phosphorous chloride, PCl, the hydroxyl group is replaced by chlorine, thus; $3\mathrm{CH}_{s}\mathrm{CO.OH} + 2\mathrm{PCl}_{s} = 3\mathrm{CH}_{s}\mathrm{CO.Cl} + \mathrm{P}_{s}\mathrm{O}_{s} + 3\mathrm{HCl}.$ And on warming such a chloride with water the

acid is re-formed : $CH_1 \cdot CO \cdot Cl + H_2O = CH_1CO \cdot OH + HCl_{\bullet \bullet}$ It is thus proved that oxygen and hydrogen can be removed together from the acid molecule.

nascent hydrogen, the chlorine is replaced by

hydrogen, and an aldehyde is produced, thus;

CH₁COCl + 2H = CH₂CHO + HCl.

This aldehyde, when treated with phosphoric chloride, PCl₂, exchanges its oxygen for two atoms of chlorine, thus;

CH2CHO + PCl3 = CH2. CHCl2 + POCl3.

It is therefore inferred that the atom of oxygen replaced by chlorine in the last reaction is differently related to the other atoms in the molecule from that atom of oxygen which is replaceable by chlorine only when hydrogen accompanies it. The formula of the characteristic group, CO.OH, thus appears reasonable.

But there are many compounds of carbon exhibiting the property of exchanging hydrogen for a metal by the action of an oxide or hydroxide, which do not contain the carboxyl group. Among these compounds may be mentioned bodies such as ethane sulphonic acid, C.H., HSO,, and ethane phosphonic acid, C2H, H2l'O3, &c.; many of these bodies may be regarded as acid ethereal salts of inorganic acids. There are, however, others which, in spite of their acid preperties, it is not usual to name acids, although many of them might be legitimately included in this class. For instance the mercaptans, of which ethyl hydrosulphido may be chosen as an example, react with oxides or hydroxides in a similar manner to sulphydrie seid, $\mathbf{H}_2\mathbf{S}$, thus, $\mathbf{C}_2\mathbf{H}_2\mathbf{S}\mathbf{H} + \mathbf{K}\mathbf{O}\mathbf{H} = \mathbf{C}_2\mathbf{H}_2\mathbf{S}\mathbf{K} + \mathbf{H}_2\mathbf{O}$: and the corresponding selenion and tellurium compounds exhibit a like behaviour. Again, many of the nitro-compounds of the alkyl radicles have the power of exchanging hydrogen for a metal, under the usual limitations, as for

CH,NO. + KOH = CH,KNO. + H,O:-C(NO₂)₃H yields C(NO₂)₃R, etc. Hydroxyquinones, such as alizarin C₁₁H₈O₂(OII)₂, act as dibasic acids, forming compounds such as C14H8O2(OK)2; phenols, and their substitution derivatives, also yield metallie derivatives, e.g. sodium phenate C_eH_aONa, sodium pierate C_eH₂(NO₂)₂ ONa. On comparing such compounds with each other, and with other acids, the following deductions may be drawn: - (1.) That a powerfully electro-negative element such as fluorine, chlorine, broinine, or iodine, confers acid properties on its compound with hydrogen. (2.) That in compounds of elements exhibiting less markedly electronegative properties than the halogens, the presence of an electronegative element is necessary to the development of acid character. This may be seen from the following considerations. Hydrocarbons, such as methane, CII, exhibit no acid properties; if an atom of an electronegativo element such as oxygen or sulphur is introduced into the molecule in place of one or more atoms of hydrogen, the compound so formed, although not generally a true acid, yet exhibits a more or less acidic character. Thus, methylio alcoliol, CII,OII, forms metallic derivatives (CH3ONa, &c.) by the action on it of strongly positive metals; but as such compounds are decomposed by water, they cannot be formed in presence of that substance. Here, however, we may note that phenol, C. H. OH, and similar compounds, reacts with the hydroxides of strongly positive metals to form metallic derivatives which, although comparatively unstable, are nevertheless capable of existence in VOL. I.

presence of an excess of alkaline hydroxide. But if a derivative of a hydrocarbon contain two or more electronegative atoms or groups of atoms in the molecule, then, as a rule, this compound forms metallic derivatives of considerable stability. Thus, the replacement of two atoms of hydrogen in the molecule of an alcohol by an atom of oxygen (converting the group characteristic of primary alcohols, CH,OII, into the carboxyl group, CO.OH) is attended by a marked increase of acid properties. Similarly the existence of oxygen combined with carbon in hydroxyquinones (as earbonyl, CO) confers on hydroxyl groups present the power of exchanging their hydrogen for metals by reactions common to acids. And in presence of a large amount of an electronegative element the exchangeable bydrogen need not even be present as hydroxyl; for as shown above, such bedies as nitromethane, CH₃NO₂, form metallic derivatives, like CH₄KNO₂. Comp. Acete-acetic acid, p. 17. Regarding the relations between the nature of different elements and the acidic character of their compounds v. further Classification.

Basicity of acids. - Some acids, en treatment with the oxide or hydroxide of an alkali metal, may exchange all their hydrogen for metal, thus producing a salt; and it may not be possible to obtain from them a body intermediate between the salt and the acid; such an intermediate derivative is usually termed an acid salt. From other acids sueli intermediate derivatives are obtainable. The acids of the former class are termed 'monobasic'; those of the latter class are termed 'polybasic,' including the terms 'di-' 'tri-' 'tetra-' basic. The conception of the basicity of acids was introduced by Graham. Before his researches in 1833, it was supposed that an 'acid salt' contained, as its name implies, both acid and salt, and on the binary theory it was considered to be a compound of the two. But Graham showed that in neutral potassium phosphate there are, as he expressed it, three equivalents of petash for one equivalent of phosphoric acid, or in modern language, three atoms of potassium for one atomic group PO,; and that the acid phosphates differ frem the neutral phosphate in centaining water instead of potash, or as we should say, hydrogen in place of potassium. The composition of hydrated phosphoric acid being expressed by the symbol (old notation) PO_s.3HO, the composition of its different salts might be expressed by the symbols:

PO₅.2110.KO; PO₅.HO.2KO; PO₅.3KO. Phosphoric acid was therefore termed by Graham a 'tribasic acid.' In 1838, Liebig pointed out the necessity of considering the following acids as polybasic, because of the fact that they form acid as well as neutral salts;—cyanuric. malonie, comenic, citric, aconitie, aconic, tartaric, malic, and fumaric. In consequence of this change of view, Liebig argued that it was better to give up the binary thecey of acids held by Berzelius, and to go back to the older theory of Davy, viz. that acids are to be regarded as formed by the combination of hydrogen with simple or a compound radiole, the nature of this radicle having no part in defermining the number of stages in which the replacement of hydrogen by metal takes place. Thus by addition of oxygen or sulphur to sulphuretted

hydrogen (sulphydric acid) the following dibasic acids are obtainable:—_

Sulphydric acid . H.SO.
Sulphnrons acid . H.SO.
Sulphnrons acid . H.SO.
Sulphuric acid . H.SO.
Thiosulphuric acid . H.SO.
Dithionio acid . H.S.O.

It was, however, known that many acids, having a claim to be considered monobasic, such as hydrofinoric, acetic, benzoic, and steario, gave rise to double salts by addition of a molecule of acid to a molecule of salt. Laurent and Gerhardt pointed out that the relative densities, in the gaseous state, of many acids belonging to this class imply that a molecule of each acid contains only one atom of hydrogen; further, that an acid of this class forms only one alkyl (or ethereal) salt, and one amide; that while polybasic acids generally yield anhydrides by some direct process, often by the action of heat alone, the anhydrides of monobasic acids are usually obtained indirectly, and that anhydrosalts such as dichromate of potassium, are obtainable only from polybasic acids.

The number of atoms of hydrogen contained in a molecule of an acid is no criterion of its basicity; this fact was noticed by Gerhardt, but its bearings were more fully clucidated by Wurtz and by Kekullé. The basicity of an acid is determined, not by the number of atoms of hydrogen which it contains, but by the number of stages in which the hydrogen can be replaced, or in other words, by the number of salts which it is capable of forming with a specified monovalent metal. Thus a study of the salts of the following acids has led to their classification as shown below.

Monobasic.-HF, HCl, HNO, H(H.PO.),

H(HCO₂), HBF₄, HAuCl₄.

Dibasic.—H₂SO₄, H (HPO₄), H₂C₂O₄,

H₂PtCl₂, C₂H₄(COOH)₂.

Tribasic.—H₂PO₄, II, Fe(CN)₁₀, H₃ASO₄, COOH)₂,
C₂H₁(OH)/(COOH)₃, C₁H₂N(COOH)₄.

Tetrabasic.—H₁P₂O₃, H₄Fe(CN)₅, G₃H₂(COOH)₄.

Hoxabasic. - C. (COOH).

The number of salts of a monovalent metal which an acid is capable of forming corresponds, as a rule, with its basicity. Thus tribasic orthophosphoric acid forms three salts with potassium, viz. H_KPO₄, IK₂PO₄, and K₃PO₄; and similarly with other acids.

This classification, as already stated, is founded on a study of the salts of acids containing monovalent metals, practically of the salts formed by the action of potash or soda on the acids. The researches of Thomsen on the quantities of heat produced when acids and bases mutually react in equivalent quantities have confirmed the conclusions drawn from a study of the composition of salts. The principle of the thermal method may be thus stated :--If a diluto aqueous solution of a monobasic acid is mixed with an equivalent quantity of an alkali also in dilute aqueous solution, a definite quantity of heat is produced; if more than one equivalent of acid is used for one equivalent of base, the same quantity of heat is produced, This is shown by the examples which follow:

" Figures represent gram-units of heat.

		oase (NaOHAq)
3,700	13 700	6.850
3,090	16,300	8,200
800	2,800	3,900 1,400
	13,700 15,200	6,800 7,60 0
200	14,400	8,600
	,700 2,800 3,600 3,400	3,090 16,300 7,700 7,700 2,800 2,800 3,600 13,700 5,400 15,200 1,200 14,400

In most of these instances, the acid forms no acid salt; its hydrogen is replaceable in only one stage. But although acid salts of acetic acid (e.g., C.H.,O.,C.H.,NaC.), and of hydrofluoric acid (HF.KF), are known, the formation of these salts by the action of the neutral salt and the acid is accompanied by a very small thermal change. This fact forms a reason, in addition to those adduced by Gerhardt, for classing hydrofluoric and acetic acids with the monobasio acids.

The thermal value of the action of a base on a polybasic acid, unlike that of the action of a base on a monobasic acid, is dependent on the proportion between the number of equivalents of base and acid used. This is shown by the following examples:

Acid.	Number of equivalents of acid to one equivalent of base (NaOHAq).				
H.SO	2 1 11,200 14,600	15.500	1	7,800	1
H.SO H.PHO	- 15,900 14,900 14,800	14,500	9.600	7,300	=
H,co, ". H,Po, .	11.000 14.700 14.800	10.100 13.500	11,300	! =	5,900
H.P.O.	- 11,400	14,300	_	13,200	9,100

Again, a small thermal change is noticed whon solutions of a monobasic acid and of the potassium or solimm salt of this acid mutually react; but if a solution of a polybasic acid is allowed to react with a solution of a neutral salt of the same acid, a marked thermal change occurs. Thus the formation of KHSO₄ from K₂SO₄ and H₂SO₄ at 23° is accompanied by the disappearance of about 8,000 gram-units of heat.

ORTHO-ACIDS AND ANHYDRO-ACIDS.—The acids containing oxygen have been most completely investigated, owing to the fact that most of them are stable at ordinary temperatures, and in presence of air and water. It is inferred that in these acids oxygen and hydrogen are in intimate union, forming a hydroxyl group; the chief reason for this view, viz., that when these aoids are treated with phosphorous, or phosphoric, chloride they yield the chloride of the acid radicle, has already been stated. Thus sulphuric acid, SO2(OH)2, yields sulphuryl chloride, SO₂Cl₂; and phosphoric acid, PO(OH)₃, yields phosphoryl chlorido, POCl. Such groups as SO., sulphuryl, or Per phosphoryl, are termed acid radieles, and their compounds with hydroxyl are acids. The torm ortho-acid is employed especially in the nomenclature of carbon acids.

An ortho-acid, strictly speaking, is one in which the element to which the hydroxyls are

united is not combined with any other oxygen. Such compounds are in most cases unknown but their existence is inferred from that of their metallicor ethereal salts, e.g. Sl(@Na),; C(@CH,),;

CH₂C(OC₂H₆)₃, &c.
Many of the commonly occurring acids may be regarded as derived from such ortho-acids by removal of water; thus looked at, these soids are partial anhydrides. Their formation is

illustrated by the following examples:

S(OH), SO(OH). $SO_2(OH)_2$ SO, Unknown Sulphurio Unknown. Sulphuric Boid. anhydride.

I(OH), IO(OH), IO₂(OH), IO₃(OH) Inknown. Periodic Salts known. Salts 1.0 Unknown. Periodic known. acid. anhydride (?known)

P(OH), PO(OH)₃ P,0, PO2(OH) Unknown. Orthophosphoric Metaphos Phosphorio anhydride. acid. phoric acid.

Partial anhydrides are sometimes also formed by the condensation-products of two or more molecules of an acid, with removal of water, thus:

Unknown. Unknown. Pyrosulphuric acid. In most cases the composition of such acids is inferred from that of their salts; the very numerous natural silicates may be conveniently classified as salts of such condensed acids

(v Silicates).

Affinity (or avidity) of Acids .- By measuring the thermal changes which occur when one equivalent of an acid, in dilute aqueous solution, reacts an one equivalent of the neutral salt of another acid, also in dilute aqueous solution, it is possible to determine the proportion in which the base divides itself between the two acids. Measurements have been made by Thomsen, and he has named the proportion in which the base combines with either acid, the relative avidity of the acid. Thus when hydrochloric acid (36.4 parts) is added to potassium nitrate (101 parts), both in dilute aqueous solution, the thermal changes which occur point to an equal partition of the base between the two acids; i.e. half the potassium exists in the solution as chloride, and half as nitrate. On mixing nitric acid (63 parts) with potassium chloride (74.4 parts), the heat-change points to the same equal partition of the base. Hence it is concluded that the relative avidity, or affinity, of hydrochloric and nitric acids for potash is equal, and is expressed by the number 0.5. The relative avidity seems to be independent of the nature of the base within certain limits; it is also modified only to a small extent by the concentration of the reacting liquids, or by small changes of temperature. This conclusion of Thomsen has received thorough confirmation by the researches of Ostwald; and this is the more valuable inasmuch as Ostwald measured the partition of acids between bases by a method depending on the alteration of volume attending the mixture of an acid with the salt of another acid. The following table gives the relative affinities of some acids towards the base soda; the affinity of hydrochloric soid being taken as unity :-

Acid Avidity	HOI 1	HBr 0.89	HI 0-79	EP 0-66
· Acid	HON .	H.80,	H.SeO.	
Avidity	very small	0.49	0.45	
Acid	HNO,	H,PO,	H,B,O,	CCICOOH
Avidity Acid Avidity	(COOH), 0-24	С.Н.О. 0.03	Very small U.H.O. 0.03	0-38

For more details v. Affinity, p. 67; Acros, Basicity of, p. 51. Regarding acids v. also CLASSIFICATION. An acid with a large avidity or affinity is frequently now spoken of as a strong acid, tho term weak being applied to those acids the affinities of which are expressed by small numbers.

References .- Lavoisier, Traité élémentaire de Chimie, ed. 1789, i. 69 et passim; Kopp, Geschichte der Chemie, i. 308; iii. 17; Davy, Journal of Science and the Arts, i. 285; also G. A. 54, 377; T. 1815, 212; Berzelius, J. 6. 184; Graham, T. 1833, 253; P. M. 3. 451 and 469; Liebig, A. 26, 138, 170; A. Ch. 68, 5, 70; Laurent, A. Ch. [3] 24, 163; Methode de Chimis (1854), 62, Translation of Cavendish Soc., 39-45; Gerhardt, Gerh. (1856), 4.641; Wurtz, A.Ch. [3] 55. 466; 56, 342; 61. 161; Kekulé, A. Ch.
 60, 127; Odling, P. M., 18. 368; Thomsen,
 Thermochemische Untersuchungen, i.; P. 138. 65, 208, 498; 139, 193; 140, 88, 530; Berthelot, C. R. 75. 261, 435, 480, 538, 583; 87. 671.

ACIDS, BASICITY OF .- It has been shown in the art. Acros (q. v.) that some acids react with the hydroxide (or oxide) of potassium or sodium to form only one salt, whereas other acids by a similar reaction produce more than a single salt. The former acids are called monobasic, the latter polybasic. It was also shown in the art. Acros that the basicity of an acid may be determined by an examination of the heat of neutralisation of the acid. The thermal value of the reaction of a monobasic acid with a base, in dilute aqueous solutions, is independent of the ratio between the numbers of equivalents of acid and base used, provided not less than one equivalent of base is mixed with a single equivalent of acid; but the thermal value of the reaction of a polybasic acid with a base varies according as 1, 2, 3, &c. equivalents of base react with one equivalent of acid. If the thermal reactions which occur when acids and bases react in equivalent quantities, and in dilute aqueous solutions, are more closely examined it is found that the dibasic and tribasic acids fall into certain classes. Thomsen has especially examined this subject (Th. 1). The quantity of heat produced during the neutralisation of a dibasic acid is sometimes divisible into two exactly equal parts, according as one or two formula-weights of soda are allowed to react with one formula-weight of the acid. In other cases the thermal value of each stage of the total operation is different. Thus consider the following data: following data: [H*SiF*Aq, NaOHAq]== 18,300 [H*SO*Aq, NaOHAq]== 14,780.

[H'SiF'Aq, 2NaOHAq]=2×13,300 [H'SO'Aq, 2NaOHAq]

[H'SO'Aq, NaOHAq] = 15,850 [H'SO'Aq, 2NaOHAq] = (2 × 15,850) - 2,750. Each of these three acids represents a group.

2nd

8rd

"

Thomsen divides the dibasic soids examined by him into three groups according as the thermal value of the action of the second formula-weight of soda is (1) equal to, (2) greater than, or (3) smaller than, the value of the action of the first formula-weight.

The data are presented in the following table:-

GROUP I.

Heat produced in action of	Aci	đ
1st formula-weight	H,Sif, 13,300	H_PtCl_ 13,600
2nd ,, ,,	13,300	13,600
Gr	OUP 11.	

1st 2nd	17	33 93	14,750	14,750	13,850	н _. С.п.о. 12,450 12, 850
			~	4 -4		

			GROUP .	AIA.		
				H.SeO.		
1st	91	23	15,850	14,750	11,000	11,100
2nd	39	37	13,100	12,250	9,150	8,900
			H,CrO,	H_PHO.	C_H_(CC	O,II),
1st	**	**	13,150	14,850	12,40	00
2nd	"	93	11,550	13,600	11,78	50

The tribasio acids examined by Thomsen may also be classified according as the thermal value of the action of the second formula-weight of soda is greater or smaller than that of the first, and the value of the action of the third formula-weight is greater or smaller than that of the second. The data are as follows :-

GROUP I. Acid on H.C.H.O. H.C.II.O. (Aconitic Acid) (Citric Acid) (12,850 12,650 Heat produced in action of NaOH 1st formula-weight 12,950 12,800 13

13,350 13,550 GROUP II.

Heat n	roduced	In action	of Ac	:14
•	NaO	П	H _a AsO ₄	паро
1st for	rmula-	weight	15,000	14,850
2nd	11	*11	12,600	12,250
3rd	••	.,	8,350	6.950

Group I. of the tribasic acids corresponds to Group II. of the dibasic, and Group II. of the tribasic, to Group 111. of the dibasic, acids.

Thomsen suggests that this classification of dibasio and tribasic acids may be summarised in the following typical formula:-

DIBASIC ACIDS.

			-
Ottoab 1.	Typical formula	RH,	e.g. SiFH.;
Aold of Group II.	} "	R(011),	e.g. SO _a (OH),;
Acid of Group III.	1 4	к(он)п	e.g. 80 ₂ (0H)H.

TRIBASIO ACIDS.

Acid of Group I, Typical {R(OH), e.g. C,H,O,(OH),; Acid of Y ypical {HR(OH)H e.g. HPO,(OH)H. e.g. HPO,(OH)H.

As regards dibasic acids: in the case of every acid examined by Thomsen, except two, the thermal value of the action of the first quantity. of sods added is different from that of the second, coqual, quantity of soda. The first of the typical formulæ suggested by Thomsen for the three classes of dibasic acids is probably to

be assigned to H.PtCl, and H.SiF, only. Why should the formula R(OH), rather than R(OH)H be assigned to the acids of Group II.? The formula R(OH)H would indicate he easy separation of the acids into anhydride (R) and water (OHH). But the acids placed in Group III. are, as a class, more easily separable into anhydride and water than those placed in Group II. If the differences between the thermal values of the first and second quantities of soda acting on the acids of Group III. are tabulated we have this result: $H_2SO_4 = 2,750$; $H_2SO_3 = 2,500$; $H_2CO_3 = 1,850$; $H_2B_2O_4 = 2,200$; $H_2CO_4 = 1,600$; $H_2PHO_3 = 1,250$; $C_2H_4(CO_2H)_2 = 650$. These differences vary from 9.5 (H,SO₃) to 2.7 (C,H₄(CO,H)₂) per cent. of the total heat of neutralisation. We have good evidence in support of the statement that succinic acid is a dihydroxyl compound; therefore, although it occurs in Thomsen's third group, we must place it with those acids the typical formula of which is R(OH), i.e. with the acids of Group 11. The other acids of Group III. are fairly easily separable into anhydride and water. The formula CO, (OH)H for carbonic acid is to some extent confirmed by the fact that the higher homologues of this scia although dihydric are distinctly monobasic. If the differences between the thermal values of the first and second quantities of soda acting on the acids of Group II. are tabulated we have this result: H₂SO₄=1,900; H₂ScO₄=900; H₂C₂O₄= 600; $H_{\perp}H_{\downarrow}C_{\downarrow}O_{\downarrow}=400$. These differences vary from 6 ($H_{\perp}SO_{\downarrow}$) to 1.5 ($H_{\perp}H_{\downarrow}C_{\downarrow}O_{\downarrow}$) per cent. of the total heat of neutralisation. The differences in the case of acids of Group 111, arc considerably larger than these. When the difference between the thermal values under consideration is small, and, as a rulo, the value of the second quantity of soda is greater than that of the first, Thomsen regards the acid as, generally speaking, belonging to the type R(OH),; when the diffor-ence in question is large and the value of the second quantity of soda is, as a rule, smaller than that of the first, the acid is regarded as belonging to the type $\hat{R}(OH)H$.

These thermal investigations made by Thomsen point to the performance of definite functions by the different hydrogen atoms in the chemically reacting unit, or group of atoms, of many polybasic acids. Although the reacting unit of a tribasic acid contains three atoms of hydrogen all replaceable by metal under similar conditions, nevertheless the energy-change which accompanies any one of theso replacements is often different from the energy-change which accompanies the other replacements; hence we seem justified in concluding that each of the replaceable atoms of hydrogen in these acids is related to the rest of the atoms, which with the specified atom make up the chemically reacting unit of the acid, in a way different from that wherein the other replaceable atoms of hydrogen are related to the rest of the atomic complex in

question. In such aoids as H2SO, H2PO4, &c., it is necessary to exhibit the differences of function of the different replaceable atoms of hydrogen by formulæ which represent some of these acids as containing one OH group, others as containing two OH groups, and others three OH groups; but acids are known the reactions of which

oblige us to say that they contain more than one OH group, and at the same time to assert that each of these groups plays a different part in the reactions of the acid. Thus, glycollic acid CH_OH_COOH is a monobasic acid; the heat of

neutralisation of this acid is [CH*OH.COOHAq, NaOHAq] = 13,600 (De Forcrand, C. R. 96, 582); but the addition of a second equivalent of soda to the neutral salt is attended with the production of a small

quantity of heat

[CHOH.COONAAq, NaOHAq] = 4,200 (ib., Bl. [2] 40, 104). The disodium glycollate thus formed is, however, an easily decomposed compound. Another monobasic acid, glyoxylic, is known, having the composition CII(OH), COOH; this acid forms a definite sodium salt, an aqueous solution of which reacts with soda with the production of nearly one-sixth the quantity of heat produced by the reaction of the first equivalent of soda on the acid. The data are these (De Forcrand, C. R. 101, 1495) :--

 $[CH(OH)^2.COOHAq, NaOHAq] = 13,230;$

 $[CH(OH)^2.COONaAq, NaOIIAq] = 2,000.$ Here we have a very distinct illustration of the econnections between thermal changes and the modification in the nature of the reaction of a specified group of atoms produced by the relations of that group to the other atoms, or group of atoms, in the chemically reacting unit of au acid (v. further Affinity; especially pp. 74, 75).

M. M. P. M.

ACIDS, ORGANIC. The empirical formula of acetic acid C2H4O2 has been expanded into the structural formula CII, CO.O.H by reason of the following cousiderations. One fourth of the hydrogen of acetic acid is displaceable by metals hence we write C.H.O.II. By the action of PCl, acetic acid may be made to exchange the same quantity of hydrogen togcther with half its oxygen for chlorine, producing acetyl chlorido, C.II.OCl: hence we write C.H.O.OH. In the electrolysis of potassium acetate, ethano and carbonic acid are produced at the positive pole, potassium being formed at the negative pole. This decomposition may be represented thus; $C_2H_3O_2K = K + CO_2 + CH_3$; but methyl, CH3, is immediately polymerised, becoming ethane, C.H. This experiment shows that half of the carbon in acctic acid is intimately connected with oxygen, the other half being connected especially with hydrogen: hence we write, finally, CH3.CO.O.H.

Analogous reasoning applied to other organio acids, very frequently leads to a similar formula, s.g. in the case of succinic acid to the formula C₂H₄(CO.O.H)₂. The acid character of theso bodies is undoubtedly connected with the group The acid character of these CO.O.H or CO.H called carboxyl, and it is easy to generaliso and say that all organic acids that are free from sulphur, phosphorus, arsenie or silicon, contain carboxyl. Kekulé, therefore, considers that the basicity of an organic acid is determined solely by the number of carboxyls it contains. Such a conclusion cau, however, only be maintained, by defining an organic acid as a substance containing carboxyl. If this definition be accepted, it follows of course that all organic acids do contain carboxyl. But if we wish to let experiment guide us, we must (b.) Di-carboxylic acids: a. Di-hy. adopt some other definition, such as that an dric: Series I, C_aH_{m-x}O₄ or Oxalic Series

acid is a substance that contains hydrogen which can be displaced by metals with the formation of a metallic compound not decomposable by water. According to this definition, phenol, pyrogallic acid, nitro-ethane, and even the propargyl derivatives and perhaps acetylene, are raids. Compounds like sugar-lime are not necessarily salts, for the calcium need not have displaced any hydrogen in the sugar, but may have added itself in some way.

Sodie carbonate gives off CO2 when mixed with solutions of strong acids; if we adopt effervescence with sodic carbonato as a test of acidity, we shall consider the compounds just mentioned to be neutral bodies, but the nitrophenols and barbituric acid will still be acids. In testing with sodie carbonate we assume that carbonic acid is the weakest of all acids: this is a mere convention, the fact being that there is no definite line of demarcation between acids and neutral bodies, the two series shading off imperceptibly into one another.

It will be noticed that the acidity of phenol is greatly increased by the introduction of nitroxyl. In general, the displaceable hydrogen in an acid must be directly and indirectly attached to strong chlorous (or electro-negative) clements or radicles, for it is the balance of affinities between these elements or radioles and the metal that produces the stability of the salt. In carboxylic salts one O directly, and CO indirectly, neutralise or balance the metal, say sodium, forming the stable group CO.O.Na.

In sodium nitro-phenol, NO, C,H,O.Na, the sodium is balanced by O directly and by NO. indirectly. In sodium uitrate, NO, O.Na, the condition of the molecule is similar (v. also ACETO-ACETIC ACID, p. 22). Too many or too few chlorous groups weaken an acid, for the equilibrium of its salts is thereby destroyed. Thus aldehyde, CII, CO.II is a neutral body, while hydric hypochlorite, Cl.O.II, is a weaker acid than Cl.II.

For purposes of classification, it is most convenient to arrange acids according to their structural formulæ. Compounds whose structural formulæ exhibit closed rings, each containing more than two atoms, are classed as argmatic, a term that is more particularly applied to the derivatives of benzene; all other organic compounds belong to the fatty scrics.

Carboxylic acids of each series may be arranged according to their formulæ and general

characters as follows:

A. Fatty Series. (a.) Monocarboxylic acids: a. Mono-hydrie: (a.) Monocaroxytte acass: a. Mono-hydris; Series II, C. H..., O. or Actolic Series; Series III, C. H..., O. or Propiolic Series; Series IV, C. H..., O. or Propiolic Series; Series IV, C. H..., O. e.g. tri-cthenyl-butyric.—\$\beta\$. Di-hydric: Series I, O. H..., O. or Lactic Series; Scries II, C. H..., O. or Cavy-acrylic; Series III. $C_n II_{2n-1}O_n$, e.g. oxypentinoie.— γ . Pri-flydrio: $C_n H_{2n}O_n$ or Glycerio Series. — δ_n Ketonio: Series I, $C_n H_{2n-2}O_n$, e.g. aceto acetio acid; Series II, C_nH_{2n-1}O₃, e.g. allyl-aceto-acetic acid; Series III, C_nH_{2n-2}O₃, e.g. di-allyl-aceto-acetic acid.—

c. Di-ketonic: C_nH_{2n-4}O₄, e.g. ace l-aceto-acetic acid.

Series II, C_aH₂₋₁O₄, s.g. fumaric acid; Series III, C_aH₂₋₁O₄, s.g. acetylene di-carboxylic acid; Series IV, C_aH₂₋₁O₄, s.g. di-acetylene di-carboxylic acid.—β. Tri-hydrio: Series I, CaH2n-2Os, malic series; Series II, CaH2n-4Os. $C_aH_{2n-2}O_3$, mante series; Series II, $C_aH_{2n-2}O_3$, e.g. oxy-itaconie acid. — γ . Tetra -hydrio, $C_aH_{2n-2}O_4$, e.g. tartaric acid. — ξ . Penta-hydrio, $C_aH_{2n-2}O_4$, e.g. tri-oxy-adipio acid. — ξ . Hexanhydrio, $C_aH_{2n-2}O_5$, e.g. saccharic acid. — ξ . Ketonio, $C_aH_{2n-2}O_5$, e.g. acetyl-succinic acid. — ξ . Di-ketonic, $C_aH_{2n-3}O_5$, e.g. di-acetyl-succinio acid. cinic acid.

(c) Tri-carboxylic acids: a. Tri-hydric: Series I, $C_nH_{2n-4}O_s$, e.g. tricarballylic acid; Series II, $C_nH_{2n-4}O_s$, e.g. acontic acid.—\$\mu\$. Tetra-hydric: $C_nH_{2n-4}O_s$, e.g. citric acid.—\$\mu\$. Penta-hydric: $C_nH_{2n-4}O_s$, e.g. desoxalic acid.—\$\mu\$. Kotonic: $C_nH_{2n-4}O_s$, e.g. acetyl-triaghall-lic acid. carballylic acid.

(d) Tetra-carboxylic acids: a. Tetra-hydric

C.H. -O., e.g. ethanc tetra-carboxylic acid.

B. Aromatic Series. It is obvious that when rings of atoms are introduced into the structural formule, the empirical formulæ beoome very complicated. We shall therefore not attempt fully to classify the aromatic acids. The most important scries are as follows:

(a.) Mono-carboxylic acids, a. Mono-hydric: $C_nH_{2n-10}O_x$, e.g. benzoic acid; $C_nH_{2n-10}O_x$, e.g. cinnamic acid; $C_nH_{2n-12}O_x$, e.g. phenyl-propiolic acid; $C_nH_{2n-14}O_x$, e.g. naphthoic acid; $C_nH_{2n-10}O_x$, e.g. di-phenic acid; $C_nH_{2n-10}O_x$, e.g. anather phenyl-cinnamic acid; $C_nH_{2n-20}O_{2n}$ e.g. anthracene carboxylic acid; $C_nH_{2n-20}O_{2n}$ e.g. tri-phenyl-acetic acid.— β . Di-hydric: $C_nH_{2n-3}O_{3n}$ e.g. salicylic acid; $C_nH_{2n-10}O_3$, e.g. coumaric acid.—

7. Tri-hydric: $C_nH_{2n-8}O_4$, e.g. protocatechnic acid, $C_nH_{2n-10}O_{,i}$, e.g. oxy-coumaric acid.—5. Tetra-hydrio, $C_nH_{2n-8}O_{,i}$, e.g. gallic acid. e. Ketonic: CaH2n-10O4, c.g. oxy-acetophenone carboxylic acid.

(b.) Di-carboxylic acids: a. Di-hydric: C.H.2n-8O4, e.g. hydro-terephthalic acid; C. H2n-10O4, e.g. phthalic acid. - B. tri-hydric:

CaH_{2n-10}O₂ e.g. oxy-phthalic acid.
The more complicated aromatic acids may

be classified in a similar way. It will be seen that they are all poorer in hydrogen than the corresponding fatty acids.

Organic Acids in general .- Occurrence: In the vegetable kingdom, e.g. oxalic, malic, tartaric, benzoic, salicylic, cinnamic, veratric, gallic, and tannic acids. In animal juices and secretions, e.g. lactic, sarcolactic, nric, hippuric, glycocholic, and taurocholic acids. In decaying organised matter, e.g. acetic, butyric, valcric, amido-propionic, amido-hexoic, and glutamic acids.

Formation.-1. By decomposing products of the animal or vegetable kingdom by boiling with dilute acids, e.g. amido acetic, aspartic and glutamio acids.—2. From fats and fatty oils by toiling with alkalis, e.g. stearie, palmitie, and oleio acids—3. From resins by potashinsion, e.g. poly-benzoio and protocatechnic acids—4. By boiling a variety of substances with dilnte nitric acid (S.G. 1.2), e.g. oxalio and traverse acids from average acids. tartarie soids from sugar and other carbo hydrates.—5. By oxidising aromatic hydroearbons and other bodies with chromic mixture (2 pis. of K,Cr,O,, 8 pis. of H,SO, and 8 to 5

parts of water), e.g. benzele and terephthalic soids.—6. By oxidation with KMnO, e.g. vanillies acid from coniferin, pyridine oarboxylic acids, from methyl-pyridines.—7. From nitriles by boiling with KOH, e.g. acctic and snecinic soids. Unstable nitriles must be first converted into amides by cold cone. HCl, and the amides may then be turned into acids by boiling dilute HCl, e.g. pyruvic acid (Claisen). The nitriles may be prepared from alkyl ohlorides or potassio alkyl sulphates by distilling with KCy or digesting with HgCy2. No nitriles of the form XYC(CN), are known (Claus), hence derivatives of malonic acid cannot be prepared in this way. -8. By the oxidation of primary alcohols: X.CH2.OH + O2 = X.CO.OII + H.O. Secondary and tertiary alcohols can only produce acids with a less number of carbon atoms, e.g. CH₃.CH(OH).CH₂+50=CH₃CO₂H + CO₂H₂ + H₂O.

Preparation.—The acids may be separated

from insoluble neutral and alkaline substances by solution in aqueous potash; they may then be liberated by H2SO4 and purified by one of the

following methods:

(a.) If they are volatile, they are distilled alone or with steam.

(b.) By conversion into a lead, barium, or silver salt and, if possible, purifying the salt by crystallisation. The lead salt is then decomposed by H2S, the barium salt by the calculated quantity of H.SO, and the silver salt either by H₂S or by the calculated quantity of HCl.

(c.) By acidifying and extracting with ether. A large number of acids are soluble in other.

Reactions .- 1. Organic acids may be converted into ethers in two principal ways: (a.) By distilling with an alcohol and dilute H,SO, The reaction may be supposed to take place in two stages; the preparation of acetic ether may be thus represented:

EtOH + H2SO4 = EtHSO4 + H2O $EtHSO_1 + HOAc = EtOAc + H.SO_4$

(b.) If an acid is non-volatile, it is dissolved in the alcohol and the liquid is saturated with HCl. After some hours the solution is ponred into water and the ppd. ether distilled, if possible, in vacuo; the reactions may be thus represented:

 $EtOH + HCl = EtCl + H_2C$ EtCl + HOAc = EtOAc + IICl.

It is not necessary that IICl or H2SO4 should be present in order that etherification may take place, for if equivalent quantities of an acid and an alcohol be left in contact or heated together for a sufficiently long time, from 64 to 74 p.c. will react upon each other, forming an other. The rate at which the reaction takes place is greatest for acids of the formula X.CII. CO.H., slower for so called secondary acids, XYCH.CO.H., and slowest for tertiary acids of the type XYZ.C.CO.H., where X.Y and Z are alkyls (Menschutkin, v. CHEMICAL CHANGE).-2. Chlorides of phosphorus convert acids or their salts into acid chlorides of the form X.CO.Cl. These are usualty soluble in ether, and are decomposed by water, more or less rapidly, into HCl and the acid X.CO.OH. Oxy-acids exchange not only their carboxylio hydroxyl for Cl, but also their other hydroxyls; but the chlorides so produced are not reconverted by water into the original acid but only into chloroteids; thus lactic acid, CH, CH(OH), CO, H, is converted by PCI, into lactyl chloride, DH, CHCI, COCI, whence water reproduces chloropropionic said, CH2.CHCl.CO.H. The chlorides set upon dry nitrates of the heavy metal (Ag, Pb, Cu, Zn, and Hg) producing anhydrides, e.g.: 2Ph.CO.Cl + Pb(\$0_3)_2 =

 $(Ph.CO)_2O + PbCl_2 + N.O_1 + O$ (Lachowicz, B. 18, 2990).—3. Amides are formed by the action of NII, either upon the chlorides: $X.COCl + 2NH_3 = X.CO.NH_3 + NH_3Cl$, or ethers: X.CO.OEt + NH3 = X.CO.NH2 + HOEt. The amides are usually crystalline substances, and their melting-points form important means of recognising the various acids .- 4. Acetyl chloride converts acids into anhydrides (v. Acetyl chloride).-5. COCl, converts salts into anhydrides.-6. By heating with CaO or BaO, or sometimes by heating alone, CO, can be climinated from the carboxyls.—7. Dry distillation of calcium er barium salts usually produces ketones (q. v.) .- 8. Distillation of calcium salts with calcium formate usually produces aldeliydes (q. v.).

Salts .- Salts are formed by neutralising the acids with metallic oxides or carbonates. They can be conveniently obtained by the addition of "metallic sulphates or soluble carbonates to a solution of the barium salt of the acid, or of soluble chlorides to the solution of the silver salt. Sodium, added to ethereal or alcoholic solutions of oxy-neids, displaces not only carboxylio but also hydroxylic hydrogen. compounds so produced are, in many cases, partly decomposed by water, the sedium that has displaced alcoholic hydroxyl being turned out again, e.g. CH₂.CH(ONa).CO₂Na + H₂O =

CII3.CII(OII).CO.Na + NaOII. The silver salt is usually the least soluble, and is frequently used in determining the molecular weight of an acid; for when the basicity of an acid is known the molecular weight can be deduced from the percentage of silver left after strongly heating the salt. Silver salts seldom

contain water of crystallisation.
Acetic Series CallanO2. Nomenclature. The following names are employed in this dictionary, the numbers denoting the value of n: 1. formic acid, 2. acetic acid, 3. propionic acid, 4. butyric acid, 5. valeric acid, 6. hexoic acid = caproic acid, 7. heptoic acid = cenanthic acid, 8. octoic acid = caprilic acid, 9. ennoic acid = nonylic acid = pelargonic acid, 10. decoic acid = capric acid, 11. hendecoic acid = nudecylic acid, 12. dodecoic acid = lauric acid, 13. tridecoic acid, 14. tetradecoic = myristic acid, 15. pentadecoic acid, 16. palmitic acid = hexadecoic acid. 17. heptadecoic acid, 18. stearic acid = octodecoic, 19. encudecoic acid = arachic acid, 20. behenic acid - icosoic acid.

Formation.—Besides the general methods

described above, the following may be noticed: -1. The action of CO₂ upon sodium alkyls, e.g. NaC₂H₃ + CO₂ = C₂H₃.CO₂Na. This gives one method for preparing fatty acids from compounds containing a fewer number of atoms of carbon in the molecule; another method depends upon the saponification of alkyl cyanides (v. XYCH.CO,H+CO, where X and Y may be

alkyls or hydrogen. Other di-basic acids are decomposed in a similar way when their solutions are mixed with uranium nitrate solution and exposed to sunlight .- 5. By heating sodium alcoholates with CO gas: NaOEt + CO = EtCO, Na. --6. By reducing exy-acids by heating with HI. -7. By reducing unsaturated acids by HI or sodium-amalgam. •

Reactions .- 1. Dry distillation of salts of the alkaline earths or alkalis produces ketones: e.g. $Ca(O.CO.Me)_{ij} = CaCO_{ij} + COMe_{ij}$

2. Distillation of a mixture of such salts of two acids produces mixed ketones:

 $KO.CO.Mc + KO.CO.Et = K_aCO_s + Me.CO.Et$.

If one of the salts be a formate the product is an aldeliyde:

KO.CO.Mo + KO.CO.H = K.CO, + Me.CO.H. 3. Distillation of a salt of a fatty acid with an alkaline hydrate produces a hydrocarbon: $KO.CO.Me + KO.H = K.CO_3 + MeH.$

4. Distillation of the alkalino saltswith As.O. gives organic compounds containing Ausenic (q.v.) 5. Electrolysis gives saturated hydrocarbons:

 $2C_nH_{2n+1}.CO_nK = K_2 + 2CO_n + C_{2n}H_{4n+2}$.

6. Chilorine and bromino act by substitution, not by addition .-- 7. Distillation in a current of steam of the mixture of stearic, palmitic, and oleio acids got from fat slightly decomposes them, forming all acids of the series from formic

to octoic (Cahours a. Demarcay, C. R. 90, 156).

Synthesis.—The acids of the acetic series may be built up in the following way:-(a) NaMe is converted into NaCO, Me, or sodic acctate, by CO2 (Wanklyn) .- (b) Sodic acctate is converted into ethyl alcohol in one of three ways: a. It is converted by POCl, into Ac.O, and this is reduced by sodium-analgam (Linnemann) .β. Ammonic accente is prepared, and is converted by P₂O₅ into acetomirile: NH₄CO₂Me= 2H₂O + NCMo; the nitrile is then reduced by Zn and H.SO, (Mendius) to an amine : NCMc + 2H2= II N.CII Me, which is converted by nitrons acid into an alcohol: II N.CH Mo + HNO₂ = HO.Cll.Me + N2 + HO. This last reaction is, however, accompanied by an intra-molecular change in the case of all the amines except ethylamino and methylamine; as a result of this change n-propylamine gives rise to secondary as well as n-propyl alcohol. - y. The sodio acctate is mixed with sodic formate and distilled: the aldehyde thus got is reduced to alcohol by sodium-amalgam (Lieben a. Rossi), or the oxim of the aldeligde is reduced to an amine which is then treated with nitrous acid .- (c) Ethyl alcohol so prepared can now be turned into ethyl iodide. zinc ethide, and sodium ethide, successively.

A repetition of processes (a), (b) and (c) upon Nalit will produce sodic propionate, propyl alcohol, and sodic propide successively, and so we can build up the series of fatty acids.

Instead of using the sodium alkyls, it is more convenient to use alkyl cyanides; process is then: (a) convert methyl alcohol into methyl cyamide, and this, by saponincation, into acetic acid; (b) convert acetic acid into ethyl alcohol by one of the three processes, a, B, or y, just mentioned; (c) convert ethyl alcohol into supra).—2. The action of strong KOH upon all tethyl cyanide, and proceed as before to prepare kylated aceto-acetic ethers (q, v_c) .—3. The distillation of alkyl-malonio acids: XYG(CO₂H)₂= the acetic series may also be prepared synthetically with the aid of accto-acctic ether (p. 22) or

of malouic ether (q. v.). In this way any acid of didentical melting-point that acid may be con the form CHXY.CO₂H, where X and Y are alkyls, didered pure, otherwise the process must be re can be prepared.

The descent of the acetic series may be effscted by distilling each acid with soda-lime, whereby a hydrocarbon containing one atom of carbon less is got; this hydrocarbon is oonverted by chlorino into an alkyl chloride, whence by successive treatment with AgOAc and KOH

an alcohol may be forused.

The descent may also be effected by converting the acid into an amide, mixing this with bromine and pouring the mixture into a 10 per cent. solution of NaOH. An amine, a nitrile, and a derivative of urea are then formed, the amine and the nitrile contain one atom of carbon less than the amide. The amine may bs turned into an alcohol by nitrous acid, and then oxidised to an acid; while the nitrile gives the acid on mere saponification. The amides containing at least 8 earbon atoms yield large quantities of nitrile, while the lower amides produce chiefly amino (Hofmann, B. 17, 1108). The descent through nitrile from canoic to octoic acid may be thus represented:

 $C_4H_{17}CONH_2 + 3Br_2 + 8NaOII =$ $C_7H_{18}CN + 6NaBr + Na_2CO_3 + 611_2O_2$ $C, H_{13}CN + KOII + H_{2}O = C, H_{13}CO, K + NH_{3}$

Melting Points.—While the boiling points of the acetie series of acids gradually rise with each increment of CH, the melting points of those acids that contain an odd number of atoms of carbon appear to be lower than those of the acids that contain one atom of earbon

caprilic	[16·5°]	pelargonic	$[12.5^{\circ}]$
capric	[30°]	hendecoic	[28.50]
lauric	$[43^{\circ}]$	tridecoic	[10.5]
myristic	[63·8°]	pentadecoic	[51°]
palmitic	$[62^{\circ}]$	margarie	$[59.9^{\circ}]$
stearic	[69°]	e nendecoic	[59·9°]
arachic	[750]	medullie (?)	[72·5°]
T		1	*11 . 1*

Isomerism among the fatty acids will be discussed under Classification (v. also Isomerism).

Separation of two volatile acids.—Divide the acid into two equal parts, neutralise one with ; potash, add the other and distil. The most volatile acid will pass over in preference to the other; and if it constitutes more than half the entire mixture, the distillate will consist solely of this acid. If, however, the less volatile acid be in excess, the residue will consist of its potassium salt in a pure state. The operation is repeated upon whichever portion is still a mixture. Acctic acid is an exception to the rule, for although it bo the more volatile acid, it will remain behind as acid potassium acetate (Liebig, A. 71, 355). If the distillation be performed in aqueous solution in a current of steam, the acid of highest molecular weight goes over first (Hecht, A. 209, 319).

Separation of fixed acids. - An alcoholic solution of the mixture of acids is fractionally precipitated by a cone, aqueous solution of magnesium or barium acctate or by an alcoholic solution of lead acetate. In the series of pps. so got, the first contains the acid of highest molecular weight and the last the acid of lowest molecular weight. Each fraction is decomposed by boiling dilute IICl and the melting-point taken. If a sidered pure, otherwise the process must be repeated upon each fraction (Heintz, J. pr. 66, 1; A. 92, 295).

Actylic Series C.H. -O. Nomenclature. --n=3, acrylic; 4, orotonie; 5, angelic; 6, hexen-oic; 7, heptenoic; 8, octenoic = subcronic; 9, ennenoie; 10, decenoio = campholio; 11, hendecenoic = undecylenio; 12, dodecenoic; 14, tstradecenoie; 15, pentadecenoie = cimicie; 16, hexadecenoic = hypogeic; 18, oleie = octodecenoic; 19, doeglie = enendecenoic; 22, orucie and brassie acids.

Occurrence. - As compound others in fats and

oils, e.a. ofeic acid.

Formation. -1. From β -, and sometimes from α-, bromo or iodo- derivatives of the acstic series by boiling with alkalis or Ag.O:

CH.I.CH, CO.K + KOH = $CH_{u}^{2}:CH_{v}^{2}CH_{v}^{2}CH_{v}^{2}+KI+H_{u}^{2}O_{v}^{2}$

2. From β-oxy-acids, by distillation: CH_OIT.CH_CO_H = H_O + CH_:CH.CO_H.

 From certain β-oxy-ethers by PCl₂: 3CMe_OH.CO_Et+2PCl_=

 $3CMe_2Cl.CO_2Et + P_2O_3 + 3IICl$

 $CMe_*Cl_*CO_*Et = CH_*:CMe_*CO_*Et + HCl$ (Frankland a. Duppa, C. J. [2] 3, 133). Similarly Me.C(OH).CH..CO.Et gives Mc.C:CH.CO.Et.— 4. From derivatives of aceto-acetic or malonio ethers containing allyl, othylene, &c .- 5. By Perkin's reaction, by heating aldehydes, C,H2,O, with sodic acctate and Ac.O (v. Aldentdes).

Properties .- As in the acetic series, the lower members of the acrylic series are volatile liquids miscible with water. Solubility and specific gravity diminish as molecular weight and boiling point increase. The higher members are nonvolatile and insoluble in water.

Reactions. - 1. The acids of this series contain the group C:C and consequently combine directly with bromine and chlorine, usually with HBr or HI in conc. solution, and frequently with If, the latter combination is effected either by action of sodium amalgam on a solution in water or alcohol or by heating with conc. III .- 2. Fusion with potash produces two acids, one of which is almost always acctic acid. The molecular formula is split up in the middle of the group C:C, e.g.

CII, CH: CII. CO. H + 2KOH = CH,CO2K + H,CH.CO2K + H,

3. Boiled diluto with H2SO, they often change into the lactone of saturated oxy-acids: thus hydro-sorbic acid, CH, CH, CH, CH, CH, CO, H, changes into oxy-hexo-lactone,

CH, CH, CH, CH, CH, CO,

4. Many of the higher members are polymerised by nitrons acid.

Series $C_n H_{2n-1} O_2$. Nomenclature. -n=3. Propiolic; 4, tetroAc; 5, pentinoic; 6, hexinoic = sorbice 7, heptinoic = benzoleie; 8, octinoic = di-allylacctic; 9, lauronolic = enninoie; 10, camphic = decinoic; 11, hend inoic = undecolic; 14, myristolie = tetradecinoie; 15, pontadocinoie; 16, palmitolic = hexadecinoic; 17, elcomargarie = heptadecinoic; 18, stearolic = octodecinoie; 22, bebenolic.

Formation. - From di-prominated (or di-chlorseries of consecutive fractions contains acids of inated) acids of the acetic series, or mone-

brominated acids of the acrylic series by treatment with alcoholic KOH.

Reactions.—Combine with Br, or with Br, also with HBr or 2HBr, and with H, and sometimes with H ..

Lactic Series C_nH_{2n}O₃. Nomenclature.—n = 2, Glycollio; 3, lactic and hydraerylic; 4, oxybutyric; 5, oxy-valerie; 6, oxy-hexoic &c .-oxy being prefixed to the names used in tho

Formation.—1. By the general methode: thus (7) by saponification of oxy-nitriles, (8) by oxidation of glycole .- 2. From bromo-, chloroor iodo- derivatives of the acetic series by displacing the halogen by hydroxyl: (a) by boiling with much water, (b) by moist Ag₂O, or (c) by KOHAq.—3. From amide-acids by nitrous acid. 4. From aldehydes or ketones by addition of HCN followed by eaponification of the resulting oyanhydrin by HCl:

 CII_3 .CHO + HCN = CH_3 .CH(OII).CN CH_3 .CH(OII).CN + $HCl + 2H_2O =$ CH, CH(OH).CO, H + NH, Cl.

5. By oxidation of acids containing methenyl: $(CII_3)_2CH.CO_2H + O = (CH_3)_2C(OII).CO_2H.$ 6. By action of zinc alkyls on oxalic others:

CO2Et.CO2Et + 2ZnEt2 = CO,Et.CEt,.OZnEt + EtZnOEt, CO₂Et.CEt₂.OZnEt + 2H₂O = CO₂Et.CEt₂.OII + Zn(OH)₂ + C₂H₆.

7. By reduction of ketonic acide.

Reactions .- The action of PCl, and of Na, and the characters of the resulting compounds have been discussed above. The oxy-acids act as half alcohol and half acid. Thus they form two kinds of monoethylicethers, oncof the form R"(OH).CO,Et, the other of the form R"(OEt).CO.H. The ethere R"(OH).CO.Et possess all the characters of an ethyl salt of an acid. Thus they may be prepared in the usual way from the acid, alcohol, and HCl; they may be saponified easily by alkalie; they are converted by ammonia into amidos R"(OH).CONM2; they are neutral to litmus. The ethere of the form R"(OFt).CO2H can be prepared by saponifying the diethylic ethers R"(OEt).CO₂Et and these are got from R"Cl.CO₂Et by action of NaOEt. The ethers R"(OEt).CO₂H cannot be saponified by alkalis; are only convorted into ammonium ealts, R"(OEt).CO2NH, by ammonia; and are acid to litmus.

a-oxy-acids ars split up by boiling conc. HCl

into formic acid and aldeliydee:

 CH_s .CH(OH). $CO_2H = CH_s$. $CHO + IICO_2H$. 8-oxy-acids boiled with conc. HCl give II2O and an acid of the acrylic series:

 $CH_2OH.CH_2.CO_2H = CH_2:CH.CO_2H + H_2O.$ y-oxy-acids split up, when their solution is warmed, into water and lactones (q. v.).

CH₃.CH(OH).CH₂.CH₂.CO₂H = CH₃.CH.CH₂.CH₂CO.O + H₂O

On dry distillation, the q-oxy-acids of the form R'.CH(OH).CO.II produce lactides, or compound ethers of the form R'.CII < O.CO CII.R'. 3 oxy-acids are converted by dry distillation into unsaturated acids:

 $R'.CH(OH).CH_2.CO_2H = R'.CH:CH.CO_2H + H_2O_3$

or into an aldehyds and an acid;

 $CH_1.CH(OH).CRR'.CO_1H =$

OH, CHO + HCRB'. CO2H.
Ketonic acids. Carboxylio acids represented by formulæ in which carbonyl is united to two atoms of carbon. The preparation and properties of the ketonio acids got by displacing one or two atoms of hydrogen in aceto-acetic acid by hydrocarbon radicles are described under Acero-ACETIC ACID. Homologues of aceto-acetic acid in which carbonyl and carboxyl are not both united to the same atom of carbon are described as alkoyl-derivatives of fatty acide; thus, CH, CH, CO, CH, CH, CO, CH, CH, CO, II is described as Pro-PIONYL-PROPIONIC ACID.

Ketonic acids containing two carboxyls are described as derivativee of di-basic acide, thus CII, CO.CH(CO.H).CII, CO.H is described as acctyl-succinic acid.

Ketonic acids of the form R'CO.CO.H can be prepared from evanishes of the form R'.CO.CN. and also, in the aromatic series, by the action of HgPh, &c., on ClCO.CO,Et.

Di-basic acids, C,H2n-2O

Formation.—1. By oxidation of the corresponding glycols.—2. By holling the oyanides of alkylenes with potash. Alkylidene cyanides do not exist (Clans). -- 3. By saponification of cyanoacids, C_nH_{2n-1}CyO₂.-4. By reduction of unsaturated di-basic acids.-5. By action of reduced eilver upon iodo-acids (e.g. formation of adipic from iodo-propionie acid). - 6. By oxidation of fats, fatty acids, ketonic acids, unsaturated acids and many other bodies .- 7. By reducing polyhydric di-carboxylic aoids by HI, e.g. tartaric acid to succinic .-- 8. From acetoacctic ethers by acting with NaOEt and the ethyl salt of a chloro- or bromo-acid, and caponifying the product with cone. KOH (v. ACETYL-SUCCINIC ETHER). -9. From eodio-malonio ether and alkyl iodides:

 $CHNa.(CO_2Et)_2 + IR = CHR(CO_2Et)_2 + NaI.$ The product still contains hydrogen displaceable by sodium: CHR(CO2Et)2 + Na = CNaR(CO,Et), + II whence alkyl iodides form

di-alkylated malonio ether:

 $C\tilde{N}aR(CO_2Et)_2 + R'I = CR'R(CO_2Et)_2 + NaI$ (v. Malonio acid).

Properties .- Solid and not volatile. Frsquently produce anhydrides when heated. Malonic acid and its derivatives are eplit up by heat into CO, and acids of the acetic series. In the oxalic ceries the acids containing an even number of carbon atoms in the molecule have higher melting-points, and lower solubility in water than the acids with an uneven number of carbon atoms (Baeyer, B. 10, 1286; Henry, C.R. 99, 1157; 100, 60).

For the characters of the homologues of benzoie and salicylio acids see Aromatio Series. Ses also AMIDO-, BROMO-, CHLORO-, 1000- and NITRO-ACIDS, and SULPHONIO ACIDS.

ACONIC ACID C,H,O, i.e. CO,H.CH, C CIP 0

M. w. 128. [163°-164°]. S. 17·8 at 15°. Formed by boiling itadibromopyroturtaric acid with a causctio alkali, C.H.Br.O. = 2HBr + C.H.O. (Kekulé, A. Suppl. 1, 338), or with water (10 pte.) (Beer, A. 216, 92). Iu like manner from bromoitaconic acid. O.H. BrO, (Swarts, J. 1873, 584). To prepare it, a solution of itadibromopyrotartaric acid neu alised with soda is heated to the boiling point, then gradually mixed with more soda till it contains 3 mol. NaOH to 1 mol. of the acid. On evaporating the solution, sodium aconate crystallises out and aconic acid may be obtained from it by decomposition with sulphuric acid and agitation with ether (Meilly A. 171, 158).

Separatee from water in rhombie crystals;

from ether in elongated lamins (M.).

Reactions.—1. Decomposed slowly by boiling water, forming a brown syrup.—2. Does not combine with bromine.—3. Unites with HCl and HBr forming chloro- and bromo-itaconic acids (Swarts).—4. Reduced by Sn or Zn to itaconic acid.—5. Boiling baryta produces formic, succinio and oxy-itaconic acids:

$$C_{5}H_{4}O_{4} + 2\Pi_{1}O = CH_{2}O_{2} + C_{1}H_{2}O_{4}$$
 and $C_{5}H_{4}O_{4} + H_{2}O = C_{1}H_{2}O_{3}$.

Salts.—Ba A'_2 : v. sol. water, ppd. by alcohol; crystallises from hot alcohol in shining prisms; gives off 21 aq. at 150° .—Cu Λ'_1 faq: blue prisms.—Ag Λ'_2 : sparingly soluble lamine. Na Λ'_2 3aq: efflorescent triclinic crystals: a:b:c=538:1:699; a=10396, $\beta=104^\circ$ 27', $\gamma=84^\circ$ 49'. Got by beiling ita di-bronne-pyrotartaric acid with the calculated quantity of aqueous Na $_2$ CO $_3$. It is decomposed by long boiling with water (B).—Zu Λ'_2 8aq: large shining crystals, melting below 100° in their water of crystallisation.

Methyl Ether. MeA' [85°]. From AgA' and MeI. Long thin prisms, v. sol. ether, m. sol. alcohol, sl. sol. water. II. W.

ACONITANILIC ACID v. ANILINE.

ACONITE ALKALOIDS. — 1. Aconitine. $C_{siH_{18}NO_{12}}[185^{\circ}\text{ cor.}]$. S. ·13; S (benzenc or chloroform) 18; S. (ether) 1·56; S. (alcohol) 2·7; S. (petroleum) ·036 (Jürgens, Ar. 1·h. [3] 24, 127).

Occurrence.—In the root of monk's hood, Aconitum Napellus (Geiger a. Hesse, A. 7, 276;

Planta, A. 74, 257).

Preparation.—The ground root is exhausted with alcohol containing tartaric acid; the concentrated extract, after exposure to the air in shallow dishes to remove the last traces of alcohol, is mixed with water; the aqueous solution is filtered to separate resin, the last portions of which are removed by agitation with light petroloum, and then precipitated with potassium carbonate; the precipitate, consisting chiefly of aconitine, is dissolved in ether, which leaves behind a small quantity of humons substance; the colution is mixed with aqueous tartaric acid and precipitated with sodium carbonate; the precipitate is dissolved in ether; and the otheric colution left to evaporate. The residue consists of nearly pure aconitine, which may be further purified by converting it into hydrohromide, decomposing this salt, after recrystallisation, with sodium carbonate, and finally recrystallising the precipitate from other (Duquesnel, C. R. 73, 207; Wright, C. J. 31, 150).

Properties.—Crystallises in rhombic or hexagonal plates. Soluble in alcohol, ether, benzene, very soluble in chloroform, insoluble in light petroleum. Extremely poisonous; the minutest particles, inhaled or blown into the eye produce excessive irritation (W., C. J. 31, 154; Wright a Luff, C. J. 33, 325). Lævogyrate (D.).

Reactions .- Slightly alkaline; forms well-

orystallised salts. Revolved by heating with alcoholic KOH into benzoic soid and a coninc:

O₃₃H₄₃NO₁₂ + H₂O = O₂H₃O₂ + G₂₈H₃₈NO₁₁;—
partly also in the same manner by dilute mineral
acids, but another portion is at the eame time
resolved into water and apon conitine. Acetio
and benzole anhydrides convert it into a cetyl
and benzole anhydrides convert it into a cetyl
and benzole anhydrides convert it into a cetyl
and benzole forms crystals containing
G₃₃H₁₈NO₁₂, HBr₁2 | H₁O; the hydrochloride
C₃₃H₁₈NO₁₂, HBr₁2 | H₂O; forms with auric chloride
the salt C₃₃H₁₈NO₁₂, HGl, Au Cl₃, which separates
in pale yellow amorphous flakes very elightly
soluble in water (W.).

Apouconitine C33H11NO11-[186°].-Formed, together with benzoic acid and aconine, by prolonged boiling of aconitine with sulphuric acid (5 p.c.) or with a saturated solution of tartario acid. The benzoic acid which separates is dissolved out by other and the apoaconitine is precipitated by sodium carbonate, while the aconine remains dissolved. Crystals soluble in ether. As poisonous as aconitine. The hydrobromids C33H 11NO 11, HBr. 2 H2O is crystalline (W. a. L.). Acetyl-a poaconitine C33H40AcNO11 [1816] Soluble in ether, separates therefrom in small . crystals. Dissolves easily in acids, forming amorphous salts (W. a. L.). Benzoylapoaconitine C₁₀H₁₀NO₁₂ = C₂₃H₁₀BzO₁₁, obtained by heating aconitine or aconine with Bz₂O, is indistinctly crystalline, softcus at about 130°, forms amorphous salts. Nitrate nearly insoluble (W. a. L.).

A conine C₂₅H₃₈NO₁₁ [130°]. Formed, together with benzoic acid, by the action of aqueous alkalis, or more completely by that of alcoholic NaOH, on aconitine (v. sup.). V. sol. water, alcohol, alkalis and chloroform, insol. ether. Amorphous. Reduces gold and silver salts at ord. temp., Fehling's solution when heated.

Salts.—3C_{2.}H_{.0}NO_{11.}2HCl and B',H.SO₄ are amorphous and probably only mixtures, B'HClAuCl₃ is a yellow floculent pp. B'HHHgI₄ is a white floculent pp. (W. a. L.).

Apoaconine, C25H4NO10 is formed by heating

the hydrochloride of aconine at 140°.

C₂₀H₃₇NO₁₀HCl is amorphous, soluble in water, precipitated by alkalis.

2. Pieroacenitine $C_{3l}H_{15}N_{10}$ —Found by T. B. Groves in a connuercial aconite root, supposed to be that of A. Napellus. Amorphous varnish, having a bitter taste, but not producing any prickly sensation on the tongue. Not poisonous. Does not melt at 100°. Salts crystallise well. The hydrochloride contains $C_{3l}H_{15}NO_{10}HCl_{23}^{-2}I_{12}O$. The gold salt B'HClAnCl₃, is a canary-yellow amorphous precipitate very slightly soluble in water (W.).

Picroaconine C₂₁H₁₁NO_p. Formed, together with benzoic acid, by the action of alcoholic KOH on picroaconitine. Closely resembles aconine. Forms C₂₁H₁₁NO_pHIHgI₂ (W. a. L.).

3. P seud & conitine $C_{ss}^{-}\Pi_{ss}NO_{12}^{-}[104^{\circ}-105^{\circ}]$. The chief basic constituent of the root of Acontum ferox, from which it is obtained by exhausting with alcohol and sulphurio acid (0.05 p.o. of the weight of the alcohol), or with methylated spirit (640 c.c.) containing a little HClAq (1 o.o.), leaving the alcohol to evaporate, precipitating the remaining solution with ammonia, dissolving the precipitate in ether, and evaporating. The crystals which separate are purified by

recrystallisation from a mixture of other and light petroleum, or by means of the nitrate.

Properties .- More soluble in alcohol and ether than aconitine; orystallises in transparent needles and eandy crystall; remaine eyrufy after rapid evaporation. The air dried base contains 1 mol. H.O. given off at 80° in a stream of air, more quickly at 100°; docomposes, with separation of water at 130°-140°. Decomposed by alcoholic soda at 100° into pseudaconine and

veratric acid:

 $C_{46}H_{49}NO_{12} + H_2O = C_{27}H_{41}NO_6 + C_9H_{10}O_4$ at 140° into veratric acid and apopseudaconine, Cy, H, NO. By mineral acids it is resolved into water and apopseudaconitine; with acctic acid it forme acetylapopseudaconitine. Salts mostly amorphous; nitrate C_{3a}H₄₈NO₁₂ NO₃H 3H₂O, crystalline. B'HClAuCl, crystallises from alcohol in small needles, slightly soluble in cold alcohol. B'HIHgi is an amorphous flocoulent pp. (W. a. L.)

Apopseudaconitine C36H47NO11 [1030]. Formed, together with pseudaconine and veratric acid, by heating pseudaconitine with dilute mineral acids, or with veratrio acid alono when pseudaconitine is heated at 100° with a saturated aqueous solution of tartaric acid. Crystallises from ether in the same forms as pseudacomitine. The nitrate is crystalline. The aurochloride C36H17NO11HClAuCl3 crystallises from alcohol in

small needles (Wright a. Luff).

C36H 16AcNO11aq. Acetyl-apopseudaconitine Formed by heating pseudaconitine at 100° with acetic anhydrido or glacial acetic acid, and separated by agitation with ether. Crystalline. Nitrate and aurochloride crystalline. Benzoytapopseudacovitine C36H 16BzNO11aq. Indistinctly orystalline. Nitrate and aurochloride crystallise, the latter from alcohol in anhydrous rosettes (W. a. I.).

Pseudaconine. C₂,H₄₁NO, [100°]. Formed, together with veratric acid, by heating pscudaconitine with dilate mineral acids, or better with alcoholic soda. Light yellow varnish, moderately soluble in water forming a strongly alkalino bitter solution, which, however, does not produce any prickly sensation on the tongue. Soluble in ether. Forms amorphous salts, Reduces silver solution and alkaline copper solution

when heated.—C₂₇H₄₁NO₉HIHgI₂ is a white

amorphous precipitate (W. a. L.). Apopseudaeonine C2, II, 19, NO8. Formed, together with veratric acid, by heating pseud aconitine with alcoholic soda at 140°. Closely resembles pseudaconine. - C₂₁H₂₁Ac₂NO₃ is an amorphous varnish melting below 100°, sparingly soluble in water. Salts amorphous. ingly soluble in water. Salts amorphous. C₂₁H₂₁B2₂NO₂ is nearly insoluble in water (W.

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4. Japaco sitine. C_{ss}H_{ss}N₂O₂₁. In the root of Aconitum Japonicum. Prepared by exhausting the root with alcohol containing 1 p.c. tartaric acid, concentrating the extract when adding water, and repeatedly agitating the filtered liquid with ether to remove resinous conetituents; precipitating the alkaloids with sodium earbonate; agitating it with other; dissolving it in aqueous tartaric acid; again precipitating it with Na,CO3, and dissolving in ether. The resulting solution when left to evaporate deposited orystals which after being from ether, and after repeated fractional crystallisation gave by analysic numbers agreeing with the formula $C_{e_0}H_{*_0}H_{*_0}O_{21}$, confirmed by the analysis of the gold-salt. The hydrobromids $C_{s_b}H_{s_b}N_2O_{21}$ 2HBr 5H₂O, and the *nitra* crystallise well (Wright a. Luff, C. J. 35, 387). the nitrate

Japaconing $C_{2a}H_{2}NO_{10}$, ie obtained, together with ber zoic aoid, by heating japaconitine with alcoholic potash: $C_{aa}H_{aa}N_{2}O_{21} + 3H_{a}O = 2C_{1}H_{4}O_{2} + 2C_{2a}H_{4}NO_{10}$. Yellowish varnish, easily soluble in other, alcohol, and chloroform; insoluble in water. C26H 11NO 10HHHgI2. Forms a mercuriodide

Japaconitine and japaconine heated with benzoic anhydride yield the same product, viz. C_{2.}H₃₈NO₇(OC₂H₃O), which is floculent, dissolves in ether, and does not crystallise. Salts non-crystalline, nitrate very sparingly soluble in

water (W. a. L.).
5. Lycaconitine C. Har N. O. 2aq. A non-crystallisable alkaloid obtained from wolf's bane, aconitum lycoctonum (Dragendorff a. Spohn, J. Ph. [5] 10, 36t; C. J. 48, 403). If heated with water under pressure it is converted into erystalline lycoctonic acid, C111118N2O7, and two alkaloids, lycaconine and acolyctine.

6. Myoctonine, C2:H2:N2Os5a4. Is an amorphous alkaloid also present in A. lycoctonum. H. W.

ACONITIC ACID CaH,O, i.e. CaH,(CO,H), or CO.H.CH.C(CO.H): CH.CO.H [186°-187°] S. 186 at 13°, S. (80 p.e. alcohol) 50 at 12°. Equisetic acid, citridic acid. Occurs as calciam salt in the roots and leaves of monk's-hood (Aconitum Napellus) and other aconites, in the herb of Delphinium Consolida collected after flowering (Wicke, A. 90, 98); in the horse-tail (Equisetum flaviatile) (Baup, A. 77, 293); in millefoil (Zanon, A. 58, 21; Hlasiwetz, J. pr. 72, 429); in the juice of the sugar-cane (Behr, B. 10, 351), and in that of sugar-beet (O. v. Lippmann, B. 12, 1649); as calcium and potassium salt in the leaves of Adonis vernalis (Linderos, A. 182, 365).

Formation .- 1. By the action of heat on citric acid, or by prolonged boiling of that acid with hydroculoric acid: C. II, O, -H2O = C.H.O. (Dessaignes, C. R. 42, 494); more quickly by heating citric acid with HCl in a scaled tube at 130° – 140° (Hergt, J. pr. [2] 8, 372), or by boiling it with HBr (Mercadante, G. 7, 248) .- 2. In small quantity, together with citraconic acid, by heating citric acid with III in a sealed tube

(Kämmerer. A. 139, 269).

Preparation. - Citrio acid, in portions of 100 grams each, is heated in small flasks provided with bent distillation tubes a met. long, till the whole tube is lined with small oily drope, and the residue is heated of a water bath with 15 g. water till it solidifies to a crystallino mase. On pulverising this mass and treating it with pure ether, acouitic acid dissolves and citric acid is left behind (Pawolleck, A. 179, 150). Hunaus (B. 9, 1751) heats citric acid at 140° for a day in a stream of HCl-gas, dissolves the product in a small quantity of water, evaporatee, and treats the residue by Pawolleck's method.

Properties and Reactions. - Cystallises in small four-sided plates, melting at 187° and resolved at the same time into CO2 and itaconic freed from adhering syrup, were re-crystallised and CallaOa; also when heated with water as 180 (Pebal, A. 98, 94). Dissolves easily in absolute ether, whereby it is distinguished from nitric acid. Convorted by sodium-amalgam into tricarballylio acid C₂H₂O₃ (Hlasiwetz, J. 1864, 896). Unites with fuming HBr, at 100°, forming bromocitrio acid C.H.BrO,, and with hypochlorous acid, forming ohforocitric acid C.H.Clo. The calcium salt fermented with cheese yields succinic acid (Dessaignes, C. R. 81, 482).

Salts. The acid is tribasic. The NH4, K, Na, Mg and Zn salts dissolve readily in water, the rest are insoluble or only sparingly soluble. The soluble aconitates form, with lead and silver solutions, white flocculent precipitates (distinc-

tion from fumaric and maleic acids).

(NH₁)H₂A": nodules or lamine. S. 15.4 at 15°.—(NH₂)₂HA": nodules or lamine. S. 15.4 at 15°.—(NH₂)₂HA":—K₂A"(2aq. S. 9 at 15° (Baup, A. 77, 299). Slender, silky, very deliquescent needles; lose aq at 100° and aq at 190° (Guinochet, C. R. 94, 455).—K.HA" 2aq. S. 37.7 at 16°: small prisms (G.).—KH₂A". S. 11 at 17°: minute alegarated wisnes.—No A" 2aq. at 17°; minute elongated prisms. - Na, A" 2aq: retains its water in a current of air at 60° but gives it up at 15° in vacuo.—Li, A" 2aq: v. sol. water; solution is alkalino.-CallA"aq: gummy; v. sol. water.— Ca₃A", 3aq: gummy; v. sol. cold water, but at 80°-100° this solution deposits rhombic prisms, sl. sol. cold water. The latter gradually dissolve, changing to the gummy variety.—C.a.A", 6aq. S. 1·01 at 15° (B.).—Sr.A", 8aq. S. ·625 at 16° (G.). Ppd. on boiling the solution.—BaH, A", 2: prisms. S. 4·2 at 17° (G.).—Ba, A", 3aq: gelatinous pp. got by adding BaCl, to aconitic acid or a solution of an aconi-BaUl₂ to acontic acid or a solution or an acontate.—Mg₂A"₂ 3aq. S. 10·1 at 17°. Elongated octahedra (G.).—Co₃A"'₂,3aq. S. 3·5 at 16°. Pink powder.—Ni₂A"₂. Pp. changed by long boiling to Ni₂A"'₂ 6aq.—Cd₂A"'₂ 6aq. S. ·113 at 17°. Prisms.—Zn₂A"'₂3aq: insolublo in water.—Pb₂A"'₂3aq(?): flocculent precipitate (Buchner).—Pb₃A"'₂2PbO2H₂O: obtained by prolonged believe of the NH self with heart lead acoustic. boiling of the NH4-salt with basic lead acctate (Otto, A. 127, 180).-Mn3A", 12aq: small rose-coloured octahedra, slightly soluble in water (Baup) .-- Ag, A" prepared by adding AgNO, to the normal ammonium salt, is a thick flocoulent precipitate, becoming crystalline on drying; slightly soluble in water.

A solution of aconitic acid mixed with ferrio chloride is precipitated by ammonia, but the presence of citrio acid even in small quantity prevents the precipitation (Bn. 1, 648).

Ethers.—The ethers of aconitic acid are formed by heating the acetyl derivatives of the corresponding citric ethers, C3H4(OAc)(CO2R)3, at 2500-2800, acetic acid being split off; yield 75 p.o. of the theoletical (Anschütz a. Klingemann, B. 18, 1953)

Me₄A" (271°) (Huneus, B. 9, 1750); (161°) at 14 mm. (A. a. K.) From aconitic acid, McOII,

and HC! (H.). Et. A''' (275°) (Mercadante, G. 1, 248); (252°) at 250 mm. (Conen, B. 12, 1655); (171°) at 14 nun. (A. a. K.). S.G. 20 1.1064 (C.); 14 1.074 (Crasso, A. 34, 59). From tetra-ethyl citrate and PCl, at 100° (Conen) F Pr.A''' (195°) at 13 mm. (A. a. K.)

Di-anilide. [217°]. Yellow needles; formed by boiling aqueous aniline aconitate (Michael, B. 19, 1874).

Iso-aconitic ether CO.Et.CH:CH.CH(CO.Et),(?)(248°)S.G.131.0505. A product of the action of hot HCl upon di-oarboxy-glutaconic ether (q. v.). An oil; sol. alcohol or ether (Conrad a. Guthzeit, A. 222, 255).

Pseudo-aconiticacid

CO2H.CH2.CH(CO2H).C".CO2H [217°] Formed at 180° from propylone tetra-carboxylic acid (q. v.), obtained from bromo-maleic ether and sodium malonio ether (Schacherl, A. 229, 95).

Salt.—Ba₃A'''₂aq. H. W. ACORIN C₃₆II₅₀O₅. A glucoside extracted from the common reed (Acorus calamus). Sol. alcohol or ether; ppd. by benzene from its ethereal solution (A. Faust, Bl. [2] 9, 392; Thoms, Ar. Ph. [3] 21, 465)

ACRALDEHYDE v. ACROLEÏN.

ACRIDINE C., II, N i.e. CII CH HC CH CH N

M. w. 179. [106°] (Bernthsen); [111°] (Fischer a. Körner). V. D. 6 10 (Graebe, B. 5, 15).

Occurrence. - In coal tar (Graebe a. Caro. A. 158, 265). The portion that boils between 300° and 360° is extracted with H2SO4Aq and the extract ppd. by K_Cr_O,

Formation.—1. By heating formyl-diphenylamino (23 g.) with ZnCl₂ (45 g.) at 190°-220°:

H.CO.N(O₆H₃)₂= G₁₃H₂N + H₂O.

2. From crystallised oxalic acid, diphenylamiue, and ZnCl₂ at 120° - 260°.—3. From chloroform, diphenylamine, and ZnCl2 of AlCl3. In this way 2 g. of acridino can be got from 25 g. diphenylamine. It is better to heat chloroform (1 pt.) with diphenylamine (1 pt.), ZnCl, (1 pt.), and ZnO ($\frac{1}{2}$ pt.), for 8 hrs. at 200° (Fischer a. Körner, B. 17, 101). (C₅11.), NH + CCl₂H + ZnO = C₁₅11., N, HCl + ZnOl₂ + H₂O. — 4. By passing phenyl-o-toluidine through a red hot tube (Graebe, B. 17, 1370).—5. In small quantity (5 p.c.) by heating aniline and ZnCl2 with oor p- oxy-bonzoic aldehyde or even with benzoic

aldehyde (Möhlau, B. 19, 2151).

Preparation.—Heat formic acid (50 g. of S.G. 1'22) with diphenylamine (175 g.) and ZnCl₂ (100 g.) gradually from 150° to 270°, avoiding evolution of CO. Dissolvo the product in alcohol, and pour into aqueous NaOII. Aeridine and diphenylamine are in the alcoholic layer; evaporate this, and dissolve the residue in ether; shake the ether with diluto hydrochlorio The acciding is then in the acid solution. The yield is small (Bernthsen, A. 224, 3).

Properties.-Long needles (from much water) or prisms, a: b: c = 656:1: 335. Pungent odour and burning taste. The base and its hydrochloride attack the tongue even when in minute quantities. Volatile with steam. Very slightly soluble in water. Dilute solutions exhibit a characteristic greenish-blue fluorescence.

Salts.—(Bernthsen, A. 224, 3; B. 16, 1802 Graebe, B. 16, 2828; Medicus, B. 17, 196.)-B'HCl: yellow plates, soluble in water imparting a bluish-green fluorescence.-B',H,PtCl, sparingly soluble minute yellow needles.

B'HNO, Saq: [1519] yellow pp. got by adding sodium nitrite to a solution of an acridine salt. Long yellow silky needles; sl. sol. ether or cold water, m. sol. het water, v. e. sol. alcohol; somewhat volatile with steam.—B'H.SO; formed by adding aqueous SO, to a solution of the hydrochloride. Yellowish-red needles, v. sl. sol. water.—B'HNaSO; got by mixing solutions of sodium sulphite and acridine hydrochloride. Colourless, easily soluble, prisms.

Picrate. C₁₁H₂NC₂H₄(NO₂)₂OH. Minnte yellow prismatic needles. Melts at a high temperature. V. sl. sol. cold alcohol, cold water or cold benzene. Boiling water partially decomposes it (Anschütz, B. 17, 438).

Acridine forms no carbonate.

Reactions.—1. IIgCl₂ gives a yellow crystal-line pp. (O₁₂H₄N,1ICl)₂H₂Cl₄—2. K₂Cr₂O, gives a yellow pp. C₁₃H₂NH₂CrO₄—3. I dissolved in KIAq gives a brownish pp. (C₁₃H₂NHI)₂I.—4. Reduced in alcoholic solution by sodium amalgam to hydro-acridine which is soluble in alcohol is the same time a white powder insoluble in alcohol is formed. Hydro-acridine, C₆H₄ CII₂ C₆H₄, is not a base. It forms prisms, [169°], sl. sol. cold alcohol, v. sol. hot alcohol or ether, insol. water. Sol. conc. II₂SO₄ and reppd. by water, unaltered. It is oxidised by Ag₂O or CrO₃ back to acridine.—5. KMnO₄ oxidises acridine to a quinoline di-carboxylic acid (acridinic acid) (Graebe a. Caro, B. 13, 99).

Octe-hydro-aeridine (aeridine-octo-hydride) C₁₉H₁₇N [84°]. (320°). Colourless plates or tables. Formed by heating aeridine or hydroaeridine with III and P at 220°.—B',HCl: colourless tables, soluble in hot water, sparingly in cold

(Graebe, B. 16, 2831), ACRIDINES.—Compounds having the general

formula

 $C_{e}H_{4} < C_{N} > C_{e}H_{4}$

They are characterised by basic properties, fluorescence in dilute solitions, capability of directly uniting with MeI, and of forming neutral dihydrides which may readily be reconverted into the original base. v. Butvl.-Aoridine, Metivil-Acridine, and Phenyl-Acridine. See also Amido-Phenyl-Acridine, Oxyphenyl-Acridine, Amido-Directorian Retrons.

ACRIDINIC ACID v. (Py. 2:3)-Quinoline-di-CARBOXYLIC ACID.

ACRIDYL-BENZO1C ACID v. PHENYL-ACRI-DINE CARBOXYLIC ACID.

ACROLACTIC ACID

C.H.O., i.e.CHO.CH..CO.H or CH(OII):CII.CO.H. Formed by boiling ethyl \$\beta\$-chloro-acrylate CHCI:CH.CO.Et, with baryta water (Pinner, B. 7, 250; A. 179, 92). The acid is a thick syrup. Its silver salt, AgA', blackens quickly on exposure to light, and is m. sol. water.

ACROLEIN C₂H₂O, i.e. CH₂:CH.CHO. Acrylic aldehyde, Acradehyde. Mol. w. 56. (52·4°). V.D. 1897. S. 2·5. S.G.²⁰ ·841; μ_p1·4089; R_∞ 25·31 (Brühl).

Formation.—1. By oxidation of allyl alcohol CH₂:CH.CH₂OH, with platinum-black or chromio acid nixture.—2. By dehydration of glycerin, C₂H₂O₃, and therefore in the destructive distillation of fats.—3. By distillation of acctone dibromide: C.H.OB₂, = 2HB₂+C.H.O.—4. From dibromide: C.H.OB₂, = 2HB₂+C.H.O.—4.

di-iodacetone and silver cyanide (M. Simpson, J. pr. 102, 880).—5. By exploding ethylene with a large excess of oxygen, the carbon being partly oxidised to CO, which with the ethylene forms acraldehyde, $C_2H_1 + CO = C_2H_1O$. This effect, however, is produced only by nascent, not by ready-formed CO (E. v. Meyer, J. pr. [2] 10, 113).

Preparation.—Anhydrous glycerin (1 pt.) is distilled with KHSO₄ (2 pts.), and the vapour, after passing over calcium chloride and lead oxide to remove water and acrylic acid, is condensed by a freezing mixture (Aronstein, A. Suppl. 3, 180).—Obtained also in large quantity as a by-product in the preparation of conanthaldehyde, from castor-oil (Schorlemmer).

Properties.—Mobile strongly refracting liquid. Vapour extremely irritating to the nose and eyes. Taste pungent and burning. It is readily converted into dis a cryl, a white amorphous body (isomeric or polymeric?), insoluble in water,

alcohol, acids, and alkalis. Reactions .- 1. Oxidised quickly in the air, or by silver-solution to acrylic acid, in the latter case with formation of a silver speculum; by nitric acid to glycollic and oxalic acids (Claus. A. Suppl. 2, 118).—2. Converted by nascent hydrogen (Zn and HCl) into allyl alcohol, C₃H₄O, isopropyl alcohol, C₃H₄O, and acropinacone 2C₃H₄O + H₂=C₃H₁₀O₂ (Linnemann, A. Suppl. 3, 257).—3. With PCl, acrolein yields allylene chlorido C,H,Cl2 (84.5°), the isomerio dihydrochloroglycideor B-epidichlorhydrin (102°), and trichlorhydrin C₃H₃Cl₃ boiling at 152°-156° (Geuther, Z. 1865, 24).—According to Romburgh (Bl. [2] 36, 549) the three liquids are allylidene chloride C₃H₁Cl₂ (85°), its isomeride, (110° cor.), and β-chloro-allyl alcohol CHCl:CH.CH2OH (153° cor.).—4. Bromine forms di-bromo-propionic aldehyde (q. v.).—5. Acrole in heated with ethyl-alcohol and its homologues and a little acetic acid, yields glycerides; e.g. triethylin C₃H₃(C₂H₅)₃O₃, from 1 vol. C₃H₄O and 1 vol. alcohol and 0.5 vol. acetic acid; trimethylin C3115(CH3)3O3, from 1 vol. C3H4O, 3 vol. methyl alcohol, and 0.5 vol. acctic acid; and triamylin $C_3\Pi_5(C_5H_{11})_3O_5$, in like manner. On passing HClgas into a unixture of acrolcin and 2 vol. absolute alcohol, diethylchlorhydrin C, H, (C, H,), ClO, is obtained as a heavy oil having a sweetish ethercal odour and S. G. 1.03 at 10.5° (Alsberg, J. 1864, 494).--6. Acrolein acts strongly on aniline, forming diallylidene-di-phenyl-di-amine, (C₆H₅), (C₃H₄), N₂ (Schiff, J. 1864, 414).

Combinations,—1. With Sodium Hydrogen Sulphite bydirectombination C, H, O, 2NaHSO, = CH_x.CH(NaSO₃). Crystalline nodules. With acids gives off SO₂ but no acrolern. By NH₃ and BaCl, only half the sulphurous acid is precipitated as BaSO₃, the asulphopropionic aldelyde remaining in solution. Sodium amalgam converts it into expropane sulphonic acid. Silver oxide oxidises it to a sulpho-propionic acid (Max Müller, B. 6, 1441.—Bn. 360).

2. With Acetic Anhydride. C.H.O.C.H.O. or C.H.(OAc). Formed by direct combination at 100°. Liquid (180°). S.G.21 1.076 (Hübner a. Geuther, A. 114, 47).

lation of fats.—8. By distillation of acetone dibromido: C₂H₂OBr₂ = 2HBr + C₂H₂O.—4. From Liquid boiling at 140°-145° (Aronstein).

4. With Ethyl Chloride.

CH₁:OH.CHCl(OC₂H₃). Formed together with
scrolein-acetal, by the action of sodium ethylate on allylidene chloride, CH.: CH.CHCl, at 120°.

Liquid. (115°-120°).

5. With Ethyl Aloohol. C₃H₁₀O₂ CH, OH.CH(OH)(OC.IL). From acrolein hydrochlorido and sodium ethylate:

CH.:CH.CHO,HCl + NaOEt = NaCl + CH.: CH.CH(OH)OEt.

Liquid boiling, with partial decomposition at 130°, S.G. 20916. Soluble in water, alcohol and ether (Geuther a. Cartmell, A. 112, 3).

6. With ammonia, acrolein forms a condensation-product $C_0H_0NO = 2C_3H_4O + NH_3 - H_2O_4$ prepared by passing the vapour of anhydrous scrolein into a coholic ammonia (llübner a. Geuther, A. 114, 35), or more readily by passing the vapour of crude acrolein into aqueous ammonia, expelling the excess of ammonia by a gentle heat and precipitating the remaining liquid with a mixture of ether and alcohol (Claus. A. 130, 186) .- Red amorphous body easily soluble in water and in acids, sparingly in hot alcohol, insoluble in cold alcohol and in other. Converted by dry distillation, first into a non-volatile oxygenated base (Clans. A. 158, 222), then into pico-lino and water (Baever, A. 155, 283). Acroleinammonia unites directly with bases, forming brown anorphous salts. The platinochloride (C.H.NOIICI) PtCl, is a yellow amorphous pp.

Polymerides. I. Metacrolein (CaH,O)a.[50°]. V.D. 5.9. Formed with evolution of hydrogen, when acrolein hydrochloride is heated with potassinm hydroxide (not NaOH). Needle-shaped crystals lighter than water, having an aromatic odour. Partly reconverted by distillation into acrolein. Volatilises undecomposed with aqueous vapour. Insoluble in cold, sparingly soluble in hot water, easily in alcohol and ether. Exerts only a feeble reducing action on ammoniscal silver solution. Not affected by dilute alkalis, but changed more or less into aerolein by heating with mineral acids. Does not combine with ammonia. Unites with dry HCl-gas forming β-chloropropionic aldehyde

CH, Cl.CH, CHO (Genther a. Cartmell, A. 112, 3). 2. Acrolein resin. Formed by heating scrolein for a week with 2-3 vol. water at 1003, as a brown resin which begins to melt at 100°, ls moderately soluble in hot water, easily in alcohol and in ether. Heated with ammoniacal silver solution, it reduces the silver in specular form. Reconverted into acroleïn at 100° (G. a. C.).

3. Hexacroleio acid C₁,H₂₁O₆. Formed by treating acrolein with alcoholic or aqueous potash or with moist silver oxide. Yellow amorphous body, insoluble in water, easily soluble in alkalis, alcohol, and ether. Has a elight acid reaction. Salts: NaC18H23O6: brown and amorphous. Ca(C181123O4)2: yellow flocculent precipitate insoluble in water and in alcohol. Barium salt; amorphous; decomposed by CO. (Claus. A. Suppl. 2, 120). II ACROLEIN - DIPHENYLAMINE v. II.W.

Dı. PHENYL-AMINE-ACROLEÏN.

ACROLEIN-UREA C.H.N.O, i.e. CON,H,(C,H,).

Formed by the action of acrolcin on nrea in clooholio solution (Leeds, A.C.J. 4, 58; B. 15, 1560). White powder; sol. alcohol, ether, or forming 8-chlorolactic soid

OS. Other bodies are 151, 206; B. 15, 1898) Other bodies are also formed (Schiff, A.

ACROLEIN-m-XYLIDINE v. m-XYLIDINE

ACROPINACONE C.H.O.

OH2:CH.CH(OH).CH(OH).CH:CH2 (1600-180°) S.G.17 99. Formed by action of zino and dilute II SO, upon aorolein (Linnemann, A. Suppl. 3, 268; L. Henry, J. pr. [2] 9, 477). It is extracted with ether. It turns brown in air. V. sol. alcohol or ether, insol. water.

ACROTHIALDINE. C. H13NS,5aq. produced by the action of ammonium sulphydrato on acrolein at 0° (Schiff, Bl. [2] 8, 444). Insol. water, v. sl. sol. alcohol, ether, or CS2.

ACRYL-COLLOIDS v. B-Baomo-Acayllo Acid. ACRYL-ALDEHYDO. PHENOXY- ACETIC ACID C11H10O4 i.e. CHO.CH:CH.C6H4.O.CH2.CO2H o [153°]. m [100°]. p [182°]. These three acids are prepared by adding a cold aqueous solution of aldehyde to a dilute solution of sodium oaldehydo-phenoxy-acetato at 50° (Elkan, B. 19, 3018)

ACRYLIC ACID C3H4C2, i.e. CH2: CH.CO2H. Mol. w. 72. [8°] (140°) (Liunemann, A. 171, 294). Formation.-1. By oxidation of acrolein (p. 61).-2. By heating β -iodopropionic acid with sodium ethylato: CH_I.CH_.CO_H + NaOEt =

NaI + EtOII + CII : CII.CO II. (v. Schneider a. Erlemneyer, B. 3, 339).—3. By heating β -iodopropionic acid with lead oxido.— 4. Together with propyl alcohol and other products, by heating allyl alcohol with KOH (Tollens, Z. [2] 6, 457).-5. From iodoform and solium ethylate (Butlerow, A. 114, 204).-6. By debromination of a-B-dibromopropionio soid with zinc-dust.

CH_Br.CHBr.CO₂H - Br₂ = CH_2 ; CH+ CO_2H . 7. By heating dichlorallyleno with water:

 $CCI_2: C: CII_2 + 2II_2O = 2IICI + CII_2: CH.CO_2H$ (Pinner, B. 7, 66). -8. By the distillation of hydraerylates $C_3H_4O_3 = C_3H_4O_2 + H_2O$ (Beilstein, A. 122, 372).

Preparation .- Acrolein mixed with 3 vol. water is poured upon recently precipitated silver oxide suspended in water in a vessel protected from light; the liquid is hoated to boiling; sodium carbonato added to slight alkaline reaction; and the mass, after evaporation to dryness, is treated with dilute sulphuric acid. The liquid is filtered (hexacroleic acid and reduced silver remaining on the filter) and the filtrate is distilled, acrylic acid then passing over (Claus. A. Suppl. 2, 117).

Properties.-Colourless liquid having a pungent odour like that of acetio acid; solidifying at low temperatures; misciblo with water.

Reactions. -1. Converted by sodium-amalgam and by boiling with zinc and dilute sulphurie acid into propionio acid (Linnemann, A. 125, 317) .- 2. Fusion with KOH gives formio and acotio acids: $C_3H_4O_2 + 2H_2O = CH_2O_2 + C_2H_4O_2 + H_2$ (Erlenmeyer, A. 191, 376).—3. Unites directly with bromine forming aβ-dibromopropionic soid, CH,Br.CHBr.CO,H, and with hydriodic acid. forming β-iodopropionio acid, CH-1.CH-CO-H (Wislicenus, A. 166, 1). -4. Its alcoholic solution saturated with HOI, yields ethyl \$\beta\$-ohloropropionate, \$\text{CH}_2\text{Cl.CH}_2\text{CO}_2\text{H}_4\$ (Linnemann, \$\beta\$: 163, 98).—5 Unites with hypochlorous soid, OH.: OH.CO.H + CIOH = CH.Cl.CHOH.CO.H (Mollikow, B. 12, 2227; 13, 2154).

Salts.—All except the silver salt are easily soluble in water. Give off part of their gold at 100°, leaving basic salts; the K., Ba., and Zn. salts decomposing in this manner oven at ordinary temperatures. - KC3H2O2 is very deliquescent

(Clausius).—NaA', microscopio needles. 100 pts. cold aloohol dissolve 0.7 pt. of this salt (Zotta, A. 192, 105). Dissolves easily in 90 p.c. alcohol. Melts with decomposition above 250° (Linnemann). Converted by heating with aqueous soda at 100° into the isomeric hydracrylio acid CH_OH_CH_CO_H (Liunemann, B. 8, 1095). —CaA'2: needles (Caspary a. Tollens). -SrA'2: small rhombic plates very soluble in water.—ZnA'₂: small scales (Clausius).—PbA'₂: shining needles soluble in alcohol.—AgA': flocsulent precipitate crystallising from boiling water in prisms (Caspary a. Tollens, A. 167,

Ethers.-MeA' (80.3°) (Weger) S.G.2 .934. From methyl a-\$-di-bromo-propionate, McOH, Zn and H₂SO₄ (C. a. T.; Kahlbaum, B. 13, 2319).

—Eth' (98·5°) (W.); (101·5°) (C. a. T.). From ethyl α·β-di-bromo-propionate, EtOH,Zn and H₂SO₁.—PrA' (122·9°)(W.). From propyl α·β-di-bromo-propionate, PrOH, Zn and H₂SO₄.—Allyl ether. CaHsA' (119°-124°) (C. a. T.).

Other derivatives of acrylic acid are described ES: BROMO-ACRYLIC ACIDS, CHLORO-ACRYLIC ACIDS, CHLORO-BROMO-ACRYLIO ACID, AMIDO-ACRYLIC ACID, Bromo-iodo-acrylic acid, Iodo-acrylic acid. See also Hydrackylic acid.

Paracrylic Acids (C,H,O,)a.-An acid probably having this composition is formed by the action of potassium cyanide on othyl a-chloropropionate at 150°. Short prisms melting at 180°-182°. Gives a brown red pp. with ferric chloride (Karetnikoff, J. R. 9, 116).—Another paracrylic acid is formed by boiling aqueous B-iodopropionic acid with execss of silver exide, till the solution becomes coloured, and metallic silver begins to separate. The same acid is formed when hydracrylic acid C3II,O3 is left in contact for several days with 1 mol. bromine .--Small crystals melting at 69°; insol. in water, slightly sol. in cold, easily in hot, alcohol. Easily takes up HI at 157° and is converted into β-iodopropionio acid.-Tho sodium salt is indistinctly crystalline, deliquescent, does not melt at 180°. The lead salt is soluble in water (Klimenko, J, R, 12, 102).

Di-acrylio acid C, H,O,.

At 250° sodium hydraerylate is decomposed into water, sodium acrylate, and sodium diacrylate. On treating this mixture with water it becomes very hot, and the sodium di-acrylato takes up aq being converted into para-adipomalato, Na₂C₆H₈O₃. The latter is thrown down as a visoid syrup when an equal volume of alcohol is added. At 200° 250° it loses aq, changing to sodium di-acrylate, an amorphous deliquescent mass, which becomes warm when breathed upon, combining again with aq. Salts.— Na,A".—BaA".—CaA": from calcic hydracrylate at 220° (Wisliconus, A. 174, 285).

Para-adipomalic acid is a syrup. It is reduced by HI to para-adipo acid C₄H₁₀O₄. Salts.—Na₂C₄H₂O₅ aq.—BaA".—CuA" aq.—PhA". All are amorphous.

ACTINOMETER.—Instrument for measuring chemical intensity of light. V. PHYSICAL METHODS, sect. OPTICAL.

ADENINE C.H.N.3aq. Occurs amongst the decomposition-products of the contents of all growing animal and vegetable cells. Formed, amongst other products, by boiling nuclein with dilute H2SO4.

Long rhombio needles (from NH, Aq. (V. sol. hot water, and in NaOHAq, v. sl. sol. Na.CO,Aq. Neutral to litmus. Insol. ether or CHCls.nitrous acid it is converted into hypoxanthine (Kossel, B. 18, 79, 1928; H. 10, 248). C, H, Ag, N,:

(Ausset, B. 16, 13, 1928); H. 10, 220); C₂H₃Ag₂N₃; insol. NH₃Aq. (C₃H,N₃)₂H₂SO₄2Aq. Sl. sol. water, ADIPIC ACID C H₁₀O, i.e. CO₄H.CH₂CH₂CH₂CH₂CH₂CQ₄H. Mol. w.146. [149°]. S. 1·44 at 15°; S. (ether) ·633 at 19° (Dicterle a. Hell, B. 17, 2221); S. 7·73 at 18°

(Wirz, A. 104, 257).

Formation.—1. By the oxidising action of nitric acid on sebacic acid, and on natural fats, e.g. hog's lard, cocoanut oil, &c .- the first product of the action being sebacic acid, which by further exidation is converted into adipic acid (Laurent, A. Ch. [2] 66, 166; Bromeis, A. 35, 105; Malaguti, A. Ch. [3] 16, 84).—2. By the action of HI and phosphorus at 140° on mncie acid (Crum Brown, A. 125, 19), or saecharic acid (De la Motte, B. 12, 1572). 3. From muconio acid $C_6\Pi_aO_1$ and sodium-amalgam (Marquardt, B. 2, 385). -4. From β -iodopropionic acid and silver at 100° - 160° ; $2(CH_2 1.CH_2 \cdot CO_2 H) + Ag_2 =$ 2AgI + (CH₂)₄(CO H)₂ (Wislicenus, A. 149, 221), 5. By reduction of di-acetylene di-carboxylic acid, CO.,11.C:C.C:C.CO.,H, or of hydro-muconio acid, C₁H_s(CO₂H), with sodium-amalgam (Baeyer, B. 18, 680).—6. By heating butano-wtetra-carboxylie acid (Perkin, B. 19, 2010).

Preparation.—Schacio acid is boiled with uitric acid, whereby it is converted into a mixture of adipic and succinic acids, which are soluble in water. The uitric acid is then eva-porated off, and the residue crystallised from water. It is then fused and the solidifie I mass is pulverised and treated with other, which dissolves the adipic acid, leaving a small quantity of succinic acid (Arppe, Z. 1865, 300).

Properties .-- Monoclinio lamine, flat needles, or feathery groups of needles. Sparingly soluble in cold water, freely in alcohol and ether. It has a tendency to form supersaturated solutions. Converted into butane by distilling with large excess of CaO (Hanriot, C. R. 101, 1156).

Salts. - The ammonium salt (NH4)2A" forms monoclinic crystals resembling augite; a:b:c= $688:1:979:\beta=82^{\circ}14'(A.217,143)$. S. 40 at 14°, At 150° it gives off all its NH2-Na2A"2aq.-Na.A"3aq: very soluble penely plates.—K.A".—BaA": white pp. S.12.04 at 12°; 7.47 at 100°. SrA"2aq.—CaA"aq. -CaA"2aq: minute needles (from alcohol), giving up their water at 100° (Laurent, C. R. 31, 351).—MgA"4aq: prisms. S. 25 at 15°.— ZuA"2aq.—CdA"2aq.— CuA"aq.—CuA"2aq.—PbA": smalle glistening plates, S. 021 at 15°.—HgA": white orystalline pp. -Ag.A": small glistening platos, S. 016 at 14°.—The ferric salt is a brown-red insoluble pp. For more detailed description of salts v. Dieterle a. Hell, B. 17, 2221.

Ethyl ether .- Et.A" (245°) (Arppo, Z.

1865, 302).

Amide O.H. (CO, NH.), [220°] S. 44 (Henry, Bl. 48, 618).—Dimethylamide C.H. (CO.NHMe). [151°-158°] (H.).

Derivatives of adipic acid v. BROMO-ADIPIO ACID. OXY-ADIPIC ACID.

Para-adipic Acid .- Formed by the action of HI on paradipimalio acid, C.H.,O.-a decomposition product of eodium hydracrylate v. di-ACRYLIO ACID.—Syrupy.—ZnC,H,O,,3H,O. Viscid flocculent pp. (Wislicenus, A. 174, 295). H. W.

Iso-adipic acid C,H8(CO2H)2.[192°].S.1 at 22°. A product of the action of bromine on the sulphate of cyanethine (q. v.). The product is extracted with other; on evaporation this leaves an oil which reacte violently with strong NH3, forming crystals of the amide of butane dicarboxylic acid, C4H, (CONH2). Converted by boiling dilute H.SO, or HCl into the acid. (E. v. Meyer, J. pr. [2] 26, 358).

Properties.—Rows of prisms, or, from hot concentrated solution in water, globular aggregates. Begins to sublime at 100°. Readily

soluble in alcohol and ether.

Salts,-A"H(NH.). Solutions of this salt give the following precipitates: Fe2Cla, reddish-white; AgNO, white; CuSO, green; Pb(OAo), on agitation, prisme crossing one another; HgCl₂, CaCl₂, BaCl₂ give no pps. — A"Ag₂, — A"Cu. — A"Pb, laq.

Amide. - (Sec above.) - Does not melt at 260°. Prisms with pyramidal ends (from water).

Adipio acid $C_1H_1(CO_2H)_2$ (Hydro-pyro-cinchonic acid) [194°]. Small white needles or glietening prisms; easily soluble in alcohol and ether, less in water. Is the chief product of reduction of pyrocinchonic acid CO_H.CMe:CMe.CO.H or of the reduction of di-chloro-adipic acid CO.H.CCIMe.CCIMe.CO.H. By conversion into the anhydride and redissolving in water it is converted into the isomeric adipic acid melting at [240°].

Salts .- CaA" 1 aq: very sparingly soluble white silky needles. - SrA" 1 aq: sparingly soluble needles .-- PbA" 3aq: white crystalline pp .--A"Cu: green pp.

CH(CH₃).CO>O. [187°] (Otto Anhydride.-

 Betkurts, B. 18, 838; Roser, B. 15, 2012; Leuckart, B. 18, 2314). Constitution .- Probably identical with the

above iso-adipio acid.

Adipic acid (C,H,(CO,II),), [210°]. Glistening plates, or prismatic needles. Formed by icomerio change from the preceding adipic acid [1940] by conversion into the anhydride and rediesolving in water.

Salt. - Ag₂A": sparingly soluble white cryetalline pp. The acid does not give an anhydrido on heating (Otto a. Beckurts, B. 18, 843).

Adipio acid C,H,(CO,H), [165°-167°]. methyl-succinic acid (?) - From aceto-acetic ether by means of sodium, a bromo propionic ether, and MeI (Hardtmuth, A. 192, 142).

Salts.—PbA": floceulent pp.—Ag₂A".
Adipic acid C₆H₁₀O₄ [142°-143°].—From tropilene (q. v.) and HNO₄(S.G. 1·285) (Ladenhurg, A. 217, 140).

217, 140). Salk.—Ag₂A".—Ammonium salt forme tri-clinic cryetale: a:b:c=.8474:1:.6496 a=

20° 20'. B=95°10'. y=100° 56'.

Constitution.-Probably identical with w-dimethyl-succinic acid (q. v.), CO₂H.OMe₃.OH₂.CO₂H.

Other isomerides of adipic acid are described as METHYL-ETHYL-MALONIO, PROPYL-MALONIC, 180-Propyl-malonic, pi-Methyl-succinic, Ethyl-succinic, and Methyl-glutario acids.

Adipic (?) aldshyds C.H.O. Formed by treating acetic aldehyde with zinc-turnings at 100° 3C₂H₄O - H₂O = C₆H₁₀O₂. Smells like wild mint, and appears to be decomposed by prolonged distillation, with formation of H₂O and higher condensation products. Unites with alkaline bisulphites, forming oryetalline compounds (Riban, C. R. 75, 98).

ADIPOCERE. - A fatty substance produced in the decomposition of animal substancce in moist ground; first found by Foureroy in tho Cimetière des Innocens at Paris. Consists of palmitic, stearic, and oleic acids (Gregory, A. 61, 362; Wetherill, J. 1855, 517). According to Ebert (B. 8, 775) it consists essentially of palmitic seid, together with margaric and oxymar-

gario acids, $C_{11}H_{21}O_{2}$ and $C_{11}H_{21}O_{3}$. II. W. ADIPOMALICACID $C_{6}H_{10}O_{1}$, is formed by boiling bromadipic acid with potash, as a viscid mass which becomes crystalline. - PbC, H,O,5H,O. White precipitate which dissolves in hot colution of lead acetate and separates therefrom in nacreous scales. Gives off 21LO at a moderate heat (Gal a. Gay-Lussae, C. R. 70, 1175). H. W.

Para-adipo-malic acid v. di-Acaylic acid.

ADIPOTARTARIC ACID C.H, O. Formed by heating pulverulent dibromadipic acid with water at 150°. Moderately soluble in alcohol and other. Much more soluble in hot than in cold water, and separates in monoclinic lamina. Optically inactive. Solution agitated with KOH yields a crystallino pp. resembling cream of tartar (Gal a. Gay-Lussae).

ADONIDIN. A glucoside in Adonis vernalis, resembling digitalin in physiological action (Cervello, Ph. [3] 13, 129; Mordagno, Ph. [3] 16, 145).

ADONINIDINE. A poisonoue substance in Adonis cupaniana (Cervello, G. 14, 493).

ÆSCIGENIN Cirlino. Formed, together with glucose, by passing HCl-gas into a boiling alcoholic solution of telsescin (infra)

 $C_{18}H_{30}O_7 + H_2O = C_{12}H_{20}O_2 + C_8H_{12}O$ Indistinctly crystalline powder, insoluble in water, soluble in alcohol. Strong sulphuric acid, in presence of sugar, dissolves it with blood-red colour. Acetyl chloride converts it into a diacetate (Rochleder, J. 1867, 751). H. W. ESCINIC ACID C24H40O2. Occure in emall

quantity in the cotyledons of ripo horee-chestnut seeds. Formed, together with propionic acid, by boiling argyræecin with potash-lye

 $C_{27}H_{42}O_{12} + 2KOH = KC_{24}H_{39}O_{12} + KC_{4}H_{4}O_{29}$ and together with butyrio acid by similar treatment of aphrodescin :

 $C_{52}II_{82}\hat{O}_{23} + 3KOII = 2KC_{21}\Pi_{33}O_{12} + KC_{4}\Pi_{7}O_{23}$ Gelatinous mass, becoming partially crystal-

Resolved by hydrochloric acid into glucose and telescin:

 $C_2H_{10}O_{12}+H_2O=C_0H_{12}O_0+C_{13}H_{20}O_1^2$. The acid potassium sait $KO_{21}H_{20}O_{11}$. $C_{22}H_{40}O_{12}$ forms silky, needles, sl. scl. water (Rochleder).

sembling orders, formed by the action of ammonia-

vapour on moist paramsouletin: C₂H₂O₄ + NH₂ + O = H₂O + C₂H₁NO (Rochleder, J. 1867, 753).

ASCIORCIN C.H.O., is formed by the action of sodium amalgam on escaletin. Dissolves in alkalis with green colour quickly changing to red. Converted by ammonia into H. W. esciorcein (Rochleder, ibid. 751).

ESCIONALIC ACID C,H,O,H2O. Produced, together with formic and oxalic acids, sometimes also protocateelinic acid, by boiling asculetin with very strong potash lye. More easily obtained pure by boiling esculetin for several hours with baryta water in an atmosphere of hydrogen. Very minutely crystalline mass. Gives with forrio ohloride a red brown colour, changing to purple-violet on addition of sodium carbonate; with ferrous sulphate and a small quantity of sodium earbonate, a deep blue colour (Rochleder, J. 1867, 752). H.W.

ESCULETIC ACID C.H., O.,
i.e. C.H. (OH), CH: CH: CO.H. Formed by boiling
esculetin with baryta (Rochleder, J. pr. 69, 211).

Salts. BaA'2. -5PbC, H,O,C,H,O, which contain the group CH:CH.CO.II, such as fumario and maleic aoids and the coumarie acids, are usually capable of existing in two forms, one of which can be easily transformed into the other. Æsouletic acid and its methyl derivatives are at present known in one form only, but tri-othyl asenletic acid and its other have been obtained in two forms, which are described below as derivatives of (a) and (B) resculetio acid.

Tri-methyl derivative C.H. (OMe), C.H. CO,H [168°]. Needles. Soluble in alcohol, ether, benzene, and hot water, sparingly in cold water. Formed by the action of alcoholio KOH on the following body. Its neutralised solution gives pps. with

AgNO₂, CuSO₄, ZnSO₄, and Ph(OAc)₂.

Methyl ether C₅H₂(OMe)₃.C₂H₂.CO₂Mo [109°]. Prisms. Distils undecomposed at a very high temperature. Soluble in alcohol, ether, and benzene, insoluble in water. Formed by svaporating dimethylesculetin (1 mol.) with a solution of NaOH (2 mols.) nearly to dryness and digesting the residue dissolved in methyl sleohol with methyl iodide (Tiemann a. Will, B. 15, 2082).

(a)-ÆSCULETIO ACID C.H.2(OH).CH:CH.CO.H Tri-ethyl-derivative C.H. (OEt)3.C.H. CO.H [103°]. Prepared by saponification of its ether. Changes when heated to its beiling point or boiled with strong HCl into the (\$)-isomeride.

On reduction with sodium-amalgam the triethyl-derivatives of both (a)- and (β)- assculetie acid give the same tri-ethoxy-plenyl-propionic acid, C₄H₂(OEt)₂CH₂CH₂CO₂Et, and with alkaline KMnO₄ the same tri-ethoxy-benzoic aldehyde,

C.H. (OEt), CHO. Ethyl-Ether C.H. (OEt), C.H. CO.Et [51°], thick yellow prisms, very soluble in alcohol, ether, and benzene, insolublo in water; prepared by heating di-ethyl-asculetin with sodium ethylate and ethyl-iodide at 100°, avoiding an excess of ethyl-iodide and longer heating than four hours; on heating to its boiling point (above 280°) it changes into the (8)-isomeride (Will, B. 16, 2110).

(9) Asconatto Acto C,H,(OH), OH:CH.CO,H

Triethylderivative O.B. (OBi), O.H. CO.H. [144°], colourless silvery crystals, easily soluble in alcohol, other, and bensene, nearly insoluble inwater; formed by saponification of its ether, or by heating the (a)-isomer to its boiling point. Ethyl-Ether O.H. (OEt), O.H. CO.Et

[75], glistening tables, easily soluble in alcohol, ether and benzehe, insoluble in water, distile undecomposed above 360°; prepared by heating. di-ethyl-asculetin with ethyl iodide and sodium ethylate at 100° for 8 hours; it is also formed by leating the (a)-isomerio ether to its boiling point (Will, B. 16, 2108).

ESCULETIN

C,H,O, i.e. C,H,(OH),

Occurs in very small quantity in horse-chestnut bark. Formed by the action of diluto acids or of emulsin on resculin (Rochleder, J. 1863, 589).

Preparation.—A solution of asculin in warm strong hydrochloric acid is boiled till it solidifies to a crystalline pulp; this after washing with water is dissolved in warm alcohol; the solution precipitated with lead acetate; the precipitate of lead-asculetin is washed with alcohol and afterwards with boiling water, then suspended in boiling water and decomposed by hydrogen sulphide; the liquid is filtered at boiling heat; and the esculetin which separates on cooling is

reerystallised (Zwenger, A. 90, 63).

Properties.—Very thin shining needles or scales consisting of C.H.O., H.O.; bitter, slightly soluble in cold, more soluble in warm, water and alcohol, nearly insoluble in ether. Aqueons solution exhibits a very faint blue fluorescence, considerably exalted, however, by addition of a small quantity of ammonium earbonate. Deep green coloration with ferric ohloride; yellow precipitate with lead acetate. Æseuletin dissolves in hydrochlorie acid; and is oxidised by nitric acid to oxalic acid. By boiling with very strong potash-lye, it is converted into formic, oxalie, protocatechuie, and a scioxalio acids; by sodium amalgam into æseiorcinol.

A hydrate C,H,O,,†H,O, isomeric with daphnetin, eccurs in horse-chestant bark in larger quantity than anhydrons assouletin. It is less soluble in water than the latter, and erystallises therefrom in small granules. Sublimes at 203°, and melts above 250°, converted into asseuletin by heating at 200° in CO2-stream. also when crystallised from hot hydrochloric acid or from absolute alcohol mixed with strong hydroeliloric acid (Roehleder, J. 1863, 588).

Æsculetin unites with MgO (Schiff, B. 18, 1951), and with solution of lead acctate forms. a lemon-yellow precipitate having the composi-

tion PbC, H,O, (Zwenger, A. 90, 63).

Diacetylasculetin C18H10Os i.e. C9H4A02O4 [134°] formed by treating assouletin with acetic anhydride and sodium acetate. Crystallises from alcohol in prisms; from water in needles, dissolves in alcohol and ether; is not coloured by ferrio chloride; is easily saponified by heating with strong sulphurio acid at 30° to 40° (Nachbaner, A. 107, 248).

~ Bromoæsculetins. - C.H.Br.D., formed by treating dibromæsenlin with strong snlphurio acid, crystallises from alcohol in yellowish needles melting at 233°, slightly soluble in water.—C.H.2Br.O., formed by adding bromine to a hot solution of asculetin in glacial acetic acid, crystallies from alcohol in long yellow needles, melting with decomposition at 240° (Liebermann a. Knietsch, B. 13, 1591).

Di-acetyl-di-bromo-asculetin C13H_Br_O4 i.e. C4H_Ae_Br_O4 [177°]. Formed by acetylatida of fil-brom-asculetin crystallises from alcohol in slender needles.

Di-acetyl-tri-bromo-æsculetin O₁₈H,Br₃O₅ i.e. C₅HAe₂Br₃O₄, formed by acetylation of tri-bromesculetin, or by bromination of diacetylæsculetin, crystallises in long very thin needles, melting, with decomposition at 180°-182°; insoluble in water (L. and K.; Liebermann a. Maetbaum, B. 14, 475).

Methyl-æsculetin C10H3O, i. e.

O₂H_{O2}(OH)(OMe) [184°] is formed by heating sesculetin (6 pts.) with methyl iodide (15 pts.) and KOH (4 pts.) dissolved in a small quantity of methyl alcohol till the liquid becomes neutral. On treating the product, after the greater part of the methyl alcohol has been given off, with water and hydrochloric acid, methylæsculetin crystallises out, while dimethylæsculetin remains in colution.

Shining needles. Soluble in cold dilute alkalis and in ammonia, and precipitated therefrom by acids. Decomposed by boiling aqueous alkalis like esculetin. Insoluble in cold, but soluble in hot water; easily soluble in alcohol, ether and benzene, insoluble in light petroleum

(Tiemann a. Will, B. 15, 2075).

Dimethylæsculetin, C₁₁H₁₀O₄ é. e. C₂H₁O₂(OMe)₂ [144°], is deposited from the mother-liquor of the preceding compound on addition of ammonia. Shining needles. Easily soluble in alcohol, ether, and benzene, nearly insoluble in light petroleum; insoluble in cold, soluble in hot, water. Dissolved by H.SO₄ and precipitated by water. H. W.

Ethyl-asculetin C₂H₄O₂ (OH) (OEt) [143°], colourless crystals, soluble in alcohol, ether, benzene, alkalie, and hot water, insoluble

in celd water.

Di-ethyl-æsculetin C₃H₂O₂(OEt)₂ [109⁹], colourless silvery plates, soluble in alcohol, 6ther, and benzene, sparingly in hot water, incoluble in cold water and cold aqueous alkalis

(Will, B. 16, 2106).

Constitution .- Aseulctin contains two hydroxyls, for it forms a di-acetyl derivative. The formation of protocatechuic and of tri-ethoxypropionio acids and of tri-etboxy-benzoic aldehyde from asculetin and tri-ethyl-asculetic acids respectively show them to be aromatic bodies. If we compare the formulæ for cou-marin C₂H₂O₂, unibelliferon, C₂H₆O₂, and æsculetin, C.H.O., we see that the two latter may be regarded as oxy- and di-oxy-coumarin. umbelliferon is oxy-coumarin has been proved by eynth sis (Tiemann a. Reimer, B. 12, 993). All three bodies are fluorescent in alkalino solution, but methyl-nmbelliferon, CoH,Ou(OMe) and methyl-æsculctin fluoresce more strongly than nmbelliferon and asculetin respectively, while di-methyl-Asculetin, C.H.O.(OMe), fluoresces most strongly of all.

When coumarin (1 mol.) is evaporated with NaOH (2 mols.) and the residue digested with

MeOH and MeI, it takes up the elements of Me₂O, becoming methylio methyl-o-coumarate; OH:CH

C,H,(OK),OH:OH.CO,K+H,O+2MeI=

C_eH₄(OMe).CHeCH.CO₂Me + H₂O + 2KI. But two isomerides may be got in this way, one, (a), when excess of MeI is avoided and the digestion is for 3 hours at 100°, the other (β) by digesting for a longer time at 150° (W. H. Perkin, C. J. 31, 417, 39, 409). Precisely the same reaction occurs when di-ethyl-æsculetin is digested with NaOEt and EtI, the two isomerio ethers, C.H.₄(OEt)₃-CH:CH.CO₂Et, being formed, the (a) compound when excess of EtI is avoided and the heating kept up for only four hours, the (β) compound by more prolonged heating. In both cases tho (a) compounds are changed by distillation into the (β) compounds. These reactions indicato analogous structure.

It is howover, remarkable that di-methylesculetin does not form a dibromido as cou-

marin docs.

Paraesculetin.—Colloo, (?).—Obtained by treating asculetin with aqueous NaHSO, at boiling heat, then adding rather dilute sulphuric acid and afterwards alcohol, whereby Na2SO4 is first thrown down, and then the compound C, H, O, NaIISO, which when decomposed by sulphurie acid yields hydrated paraesculetin C,H,O,,21H,O in indistinct crystals easily soluble in water, sparingly in other, more freely in alechol, less easily in wood-spirit, nearly insoluble in acetono and chloroform, soluble in glacial acctic acid. Paraesculetin exerts a strong reducing action in alkaline solution, throws down metallic copper from Feliling's solution at 50°-70°, and reduces indigo at ordinary temperature. Not attacked by acetic anhydride. Exposed in the moist state to ammonia-vapour, it quickly turns red, then dingy-violet, and changes after a few minutes to a sky blue liquid, which when left over sulphurio acid gives off ammonia, and again turns rea, from formation of escioreein (p. 65). Converted by heating with aniline into esculetanilide (Rochleder, J.1863, 589; 1867, 752).

C,H,O,NaliSO,,1H,O forms small needles. According to Liebermann a. Knietsch (J. 1880, 1028), the true fermula of this compound is C,H,O,,NaHSO,; it probably therefore contains a hydro-æsculetin. H. W.

ESCULIN C₁₃H₁₄O₈ [204·5°-205°] (H. Schiff, B. 14, 302).—Occurs in the bark of the horsechestnut (Æsculus Hippocastanum) especially in March before the buds open (Minor,

B. J. 12, 274; Jonas, A. 15, 266).

Preparation.—1. Horsechestnnt bark is exhansted by boiling with water; the extract is precipitated with lead actate, and the filtrate, freed from lead by hydrogen sulphide, is evaporated to a gyrup. The esculin then erystallies out after a few days, and may be purified by washing with water, and crystallising, first from weak spirit (40 p.e.), then from boiling water (Rochleder a. Seltwarz, A. 87, 186).—2. The bark is exhausted with weak aqueeus ammonia; the solution evaporated to dryness; the residue, mixed with alumina and exhausted with alondo of 95 psc.; and the esculin which crystallies from the escoloil is agitated with water and

ether, and finally washed with benzene (Fair-

thorne, C. N. 26, 4).

Properties.—Small prisms, composed of C₁₈H₁₆O₈,2H₂O Bitter, sparingly soluble in oold, easily in boiling water, the solution coagulating on cooling. 1 pt. disselves in 24 pts. boiling alcohol. Very slightly soluble in absolute ether, solnble in glacial sociic acid and ethyl acetate (Trommsdorff, A. 14, 200). The aqueous solution is slightly acid, and exhibits a blue fluorescence which disappears on addition of acids, but is restored by alkalis. Dissolves in alkalis more readily than in water.

Reactions .- 1. Æsculin is resolved at 230° into glucosan and esculctin: C15H16O9= C_aH₁₀O_b+C_bH_aO₁(Schiff); by digestion with dilute mineral acids or by treatment with emulsin, into glucose and resculetin (Rochleder a. Schwarz, A, 88, 356); hy boiling with barytawater into glucose and esculctic acid (Rechleder J. pr. 69, 211).—2. Converted by sodium-amalgam into hydræsculin (Rochleder).—3. Agitated with a small quantity of nitrio acid, it yields a yellow solution which assumes a deep blood-red colour on addition of ammonia, this •reaction affords a delicate test for esculin (Sonnenschein).—3. Conc. H.SO. (4 drops) followed by NaOClAq gives a violet colour (Raby,

J. Ph. [5] 9, 402). Æsculin forms with magnesia, the compound 2C₁₅H₁₆O₉Mg(OH)₂, which is yellow and dissolves readily in water (Schiff, B. 13, 1952).

Pentacetylæsculin

C₂₅H₂₀O₁₄ i.e. C₁₅H₁₁Ac₅O₆.
Formed by heating esculin with acetic oxide orystallises from alcohol in small needles melting at 130° (Schiff, A. 161, 73; B. 13, 1952).

Di-bromo-æsculin C15H14Br2O9. Is obtained by gradually adding bromine in calculated quantity to a solution of esculin in glacial acetic acid. Crystalliscs from glacial acetic acid in small needles; melts and decomposes at 193°-105°. Sparingly soluble in alcohol, still less in all other solvents (Liehermann Knietsch, B. 13, 1594).

Pentacetyl-dibromo-æsculin C25H21Br2O14 i.e. C15H3Br2AO5O9, prepared in like manner from dibromesculin, forms slender needles, melting at 203°-206°, converted by strong sulphuric soid into dibromasculetin,

Pentabenzoylæsculin C.H.O. C. H. Bz.O., forms nodular groups of crystals sparingly soluble in other, freely in hot alcohol (Schiff).

Trianilæscnlin

 $C_{23}H_{21}N_2O_6i.e.C_{15}H_{16}(NC_6H_5)_3O_6$, from resculin and aniline by prolonged heating at 200°. Auorphous brown powder, soluble with red colour in alcohol (Schiff, B.4, 472).

ÆTHAL v. CETYL ALCONOL.

ETHOKIRRIN. The yellow colouring matter of the flowers of Antirrhinum Linaria. H. W.

AFFINITY .- Chemical affinity is that property of bodies in virtue of which, whon hrought into contact, they react on each other, forming new bodies. It can be called a force, in so far as by its action energy is produced, namely, heat, light, electrical or mechanical energy. And, vice versa, energy must be employed to reverse the action of chemical affinity, and to decompose the combined substances.

Nothing is known as yet about the nature of chemical affinity, nor has a satisfactory hypothesis been suggested concerning it. The oldest conoeptions concerning the reasons why substances react on each other reach back to Greek philosophy; nothing has survived of them except the name affinity, which preserves the notion that those substances which are of the same origin or of the same kind, and which therefore are as it were related to each other, possess the power of mutual reaction. It is now known that the contrary of this is more correct. Moreover, the reason of the greater or smaller facility with which substances react chemically was conceived te be somewhat similar to human qualities-sympathy and antipathy. These conceptions held sway as long as the philosophy of Aristotle reigned. The breach with these ideas, which was brought about by Galileo's mechanics, introduced mechanical ideas into chemistry also. The ultimate particles of substances were imagined as furnished with points, edges, and hooks, by the aid of which were brought about their decompositions and combinations. Sir Isaac Newton's discovery of the general mutual action of masses introduced a new phase into the conception of nature. The idea of an attractive action between one small particle and another at a distance was introduced by Newton himself into chemistry, in order to explain the mutual reactions of bodies. He did not, however, consider the cause of chemical actions as identical with that of general gravitation, but as different from it, especially as regards the law concerning action at a distance.

Later investigators, Buffon, Bergmann, Berthollet, assumed, on the contrary, that both forces are of the same nature, and that ouly the circumstances under which chemical forces actespecially the close proximity of the reacting particles -- cause an apparent difference.

An influence similar to that due to Newton's astronomical discovery was exerted at the beginning of this century by a physical discovery, that of the electric current. The great chemical activity of the current was soon noticed. By its help Davy decomposed the alkalis and earths; and Berzelius made use of the phenomena of electrical decomposition for the foundation of a theory concerning chemical compounds, which rested on the supposition that chemical attraction was nothing but the attraction of the opposite electricities concentrated on the smallest parts of substances.

The electro-chemical theory of Berzelius was the first chemical theory which was based on facts. Owing to this it obtained great importanco. Taking into account the needs of the time, Berzelius developed his theory only with a view towards classification; but it did not con-tribute anything towards the investigation of the nature of chemical affinity.

The last great change in the views concerning affinity took place in the middle of this century, and was brought about by Mayer's and Joule's discovery of the equivalence of 'forces,' or more strictly, of 'the actions of forces.' It classed with mechanical, electrical, and thermal, energy, in so far as it is convertible into any of these, and can be produced from each of there When this was known, the need to trace back the mode of action of the forces of affinity to other known actions of forces ceased to exist, as a great many inferences could be drawn from this experimental fact, and a special hypothesis did

not seem called for.

Two different views have been held concerning the way in which chemical forces act, and each of these has still its followers at the present day. First it was imagined that the force acting between two different kinds of matter is similar te that acting between two masses; it brings the ultimate particles nearer together, and, if under the given circumstances this is possible, it produces combination. It would be difficult to entertain different ideas concerning the simple process of combination. The task becomes far more difficult when it is a question of simultaneous decomposition and combination. often a enbstance acts on another which is a compound without combining with it as a whole, but only combining with one of its constituent parts, and expelling the other from the original compound. The hypothesis indicated above refers these facts to the opposite action of two ferces, similar to two mechanical forces opposite in direction and unequal in magnitude, which produce motion in the direction of the greater. It was imagined that the stronger cliemical affinity overcame the weaker, and produced chemical action in its own direction.

It was in accordance with this conception that 'tables of affinity' were compiled which were to give an account of the force of chemical affinity according to its order. Such tables were first published in 1718 by H. Geoffroy. They contained, under the head of any one substance, a series of others, all of which could enter into chemical combination with the specified body. They were so arranged that the preceding body would always replace all the following bodies from their combinations with the one at the head of the list. Such tables of affinity, which were very pepnlar during the last century, culminated in the works of Bergmann, who collected the ideas indicated above in a theory of affinity. Bergmann recognised that substances can react differently according to circumstances, and therefore he gave two tables for each substance, one for the action in aqueous solution-'in the wet way'; the other for the action at the temperature of Insion-'in the dry way.' Beyond this he kept to the idea that ohemical affinity always acts exclusively in one direction, so that under all circumetance a smaller affinity is overcome by

To Claude Louis Berthoilet ie due the great merit of having shown this view to be too narrow. According to him, besides 'the intensity of the forces,' the 'mass' of the reacting substances is of importance, in so far as under the me conditions the action is the greater the more the reacting substances there is present. 'Toute batance qui tend a entrer en combinaison, agit in raison de son affinite et de sa quantité' (Status chimique, p. 2). Thie is the concise expression of the new idea which Berthellet introducèd into the theoly of affinity. But it was reserved for a much later time to develop thie idea scientifically, since one of the chief applications which its author made of it—namely, the inference

that compounds according to fixed proportions do not exist, but only such as vary in composition within fixed limits -- was proved to be false. The proof of this error which was given by Prousts Bertholbt'e compatriot, brought also the correct foundation into discredit. To this the correct foundation into discredit. To this must be added, that in the discoveriee of Richter, Dalton, and Berzelius, the ecience found such important and productive tasks that there was no inducement to investigate what of truth was left in the partially refuted hypothesis of Berthollet. The idea of 'influence of mass' ie the first, but not the only, conception which the science owes to Berthollet. This ohemist further recognised the decided influence of the physical states of the reacting bodies on the final results of the actione of affinity. He taught that the simple action, according to the measure of affinity and mass, helds only for homogeneous mixtures, but holds no longer when by the appearance of certain substances in a different state of aggregation the chemical equilibrium is disturbed. The influence of 'cohesion' and 'elasticité,' on the results of chemical actions, were explained by him perfectly clearly; how first equilibrium is established in the usual. manner, but is afterwards disturbed by ene of the substances separating out in a different state of aggregation; in consequence of this, a fresh quantity of this substance is formed, and is again separated, and thus the process repeats itself until the substance in question-gaseous or solid -has been entirely, or almost entirely, removed from the changing system. A chemical reaction carried to completion, which had been taken by Bergmann to be the normal case, appears, according to Berthollet, as the exceptional case, and occurs only because of differences in the states of aggregation of the reacting cub-

The decision between these two fundamentally different views has not yet been completely made. Even now, Bergmann's theory is propounded by some investigators, though in a somewhat modified form. In place of the greater or smaller affinity, the greater or emaller production of heat is considered by these investigators as decisive of the course of a chemical action; the fundamental idea, however, the 'exclusiveness' of the reaction, is presupposed by them also. In order to explain those partial reactions in the opposite direction, which certainly occur, those who maintain the theory of Bergmann are obliged to admit that such partial reactions are possible under certain conditions. notably under the influence of heat. These authorities are therefore under the necessity of proving the presence of euch special conditions in all those cases wherein wevdeal with incomplete reactions. Berthellet's theory, on the other hand, by making use of one and the same principle, embraces both kinds of chemical action, and teaches us to consider the one as the limiting case of the other. Apart from this legical advantage it has another; it allows us to apply definite laws, which can be stated in a mathematical form, to the action of affinity taken in eenjunction with the action of mass. But this cannot be done by Bergmann's theory. whether in its old or in its new form.

Berthollet's views met with respect and ap-

prediction from his contemporaries, but not with continuation and development; because chemistry began at this time to follow another path which had been opened up by Dalton and Berzelius. Thus it was possible for the dundamental fact of the influence of mass, the fact, namely, that chemical action decreases and increases with the rolative quantities of the acting substances, to be denied; and it became necessary to prove this fact at longth by many special oases before it could be regarded as a secure property of science. One of the first investigators to whom the merit of this proof is duo was H. Rose (P. 82, 545), who showed that in the formation of many carbonates of the heavy metals, by precipitating aqueous solutions of the salts of these motals by sodium carbonate, the precipitates contained less carbonic acid and more metallic hydroxido the more water was present in the original solutions. It followed therefore that the water, according to its quantity, expelled the carbonic acid from its combination with the metal. Roso found further (P. 94, 481; 95, 96) that when barium sulphato was fused with an equivalent quantity of potassium ear-· bouate it was only partially converted into earbonate. If the quantity of potassium earbonato was increased, appreciably more barium sulphate was decomposed, but only with a proportion of 6 to 7 equivalents did the decomposition become approximately complete.

Another case of the influence of mass was investigated by Bunsen (A. 85, 131). If to a mixture of earbonic oxide and hydrogen a quantity of oxygen is added, less than sufficient for the complete combustion of the two gases, a division of the oxygen between the two combustible gases take place. The proportion in which the oxygen combines with either gas depends on the relative masses of these gases. Bunsen's further result, that these proportions can be expressed by some small multiples of the atomio weights of the gases, has been recognised to be an error. Debus (A. 85, 103) proved in Bunsen's laboratory the fact of the influence of mass on the precipitation of mixed solutions of lime and baryta by insufficient quantities of carbonic acid. In 1865 Gladstone (Tr. 1855, 179; and C. J. 9, 54) proved the general truth of the influence of mass in various ways. His method consisted in using certain definite physical properties, especially colour, and the rotation of the plane of polarisation of a ray of light, from measurements of which to draw conclusions concerning the arrangement of compounds in a homogeneous solution. Thus, by comparing the colour of pure ferric sulphocyanide with the colour produced in mixed solutions of iron salts and potassium sulphocyanide, the same amount of iron being present in both solutions, he established the fact, that by the reaction of three equivalents of potassium sulphocyanide and one equivalent of an iron salt, only 13 per cent: of ferric sulphocyanide was formed, and that even in the presence of 375 equivalents of potassium sulphocyanide the whole of the iron had not been converted into sulphocyanide.

By this and similar methods Gladstone established the following laws:

1. When two or more binary compounds are mixed so that all resulting compounds have the

power of reacting on each other, each electropositive element enters into combination with each electronegative element, and it does so according to fixed and constant proportions.

2. These proportions are independent of tha manner in which the different elements are initially arranged. They are also not only the resultants of the various forces of attraction between the different substances, but depend also on the mass of each of these substances.

3. An alteration in the mass of one of the binary compounds produces a change in the quantity of each of the other binary compounds, and it does so in a ratio which progresses regularly. Sudden changes occur only when a substance can combine with another in more than one proportion.

4. The equilibrium of affinities is generally established after a very short time, but in soma eases the elements attain their final condition

only after hours.

5. The resulting effects are completely altered when precipitation, volatilisation, crystallisation, and similar phenomena, occur; in such cases the equilibrium which had been established at first is again disturbed by the removal of some of the chemically active substances

Harcourt and Esson (C. J. [2] 5, 460) cxamined the reaction between hydrogen peroxide and hydriodic acid, and also that between potassium permangauate and oxalic and sulphuric acids. They concluded that 'when any substance is undergoing a chemical change, of which no condition varies except the diminution of the changing substance, the amount of change occurring at any moment is directly proportional to the quantity of the substance.

The principle, first established by Steinheil (A. 48, 153), of determining the chemical composition of a homogeneous liquid by means of physical measurements, was put to practical use by Gladstone in various ways. His measurements might have served as a direct basis of a theory concerning the influence of mass, had such a theory existed. But even the extensive investigations of Berthollet and St. Giles on the etherification of acids and alcohols (A. Ch. 61, 65; 66, 68), by which the chemical inflnence of mass had been confirmed and the magnitude of this influence had been measured, did not give rise to the formulation of a mathematical theory of affinity.

Meanwhile a number of theoretical concep tions, some of old standing, were pointing in the same direction. As far back as 1851, Williamson (A. 77, 37; and C. J. 4, 110) in his fundamental researches on etherification, had propounded a theory concerning the course of . chemical reactions, which explained, better than had been done before, the nature of the chemical influence of mass. According to him, substances which react on each other, when in contact or mixed together, are by no means in a state of . neutral equilibrium, but rather in one involving a continuous exchange of constituents. The final result of the reaction depends on the direction in which this exchange of constituents can take place most easily and most frequently. The state of chemical equilibrium arrived at under any conditions is thus not a statical one, in which the forces balance each other and so no more

produce any effect, but is rather a dynamical state, in which two opposite reactions occur continually to the same extent, so that the average state of the system remains the same.

Williamson's theory was accepted for the special case for which it had been propounded. Neither the author nor any of his contemporaries, however, made an application of it to the general explanation of chemical reactions. Meanwhile, hypotheses were developed on an entirely different basis, which agreed with this theory in a most remarkable manner. These are the views concerning the gaseous state, which, first propounded by D. Bernoulli, and afterwards, independently, by Herapath, Joule, Krönig, and Clausius, were developed by Clausius and Maxwell in a mathematically well-founded theory of the states of aggregation. According to this theory, bodies are made up of molecules, which are in a state of continual motion. Iu the case of gases this motion is rectilinear, until the molecules meet with some resistance, whereby they are caused to rebound according to the laws of collision of clastic bodies. The velocity of motion increases with the temperature, and is inversely proportional to the square root of the molecular weight of the gaseous body. Likewise, there is motion within the molecules, which motion is of the nature of oscillations, the intensity of which bears a constant ratio to the motion of the molecule as a whole. Moreovor, at any specified temperature, the molecules of a homogeneous gas have not all the same velocity, but have different velocities varying from the mean value in such a manner that deviations from this value are the fewer the greater they are. In the case of liquids, the molecules have no longer the power of translational motion, but are compelled to fill a definite space owing to the forces which act between them. In other respects the statements made for gases hold good for liquids also, especially those referring to the differences of condition of the various molecules of a homogeneous substance at a constant temperature. The theory has been least developed for the case of solids; here it is to be assumed that the intermolecular forces assign to the various molecules very definite relative positions of equilibrium (v. Aggregation, States of, p. 87).

The application of these ideas to chemical processes has been made by L. Pfaundler (P. 131, 55). It can easily be seen how they directly lead to conceptions which do not appreciably differ from those of Williamson. The molecular conceptions are, however, a decided improvement on those of Williamson, since, in the differences of the conditions of various molecules, they supply a reason for the continuous interchange of atoms which Williamson supposed to occur. When the velocity of motion execeds a certain amount, there will always be present some molecules in which this velocity is so great that the connection between the atoms is loosened or destroyed. These molecules are then ready to interchange their constituent atoms, while other molecules, which have a smaller velocity, will not do so. This is the explanation for partial reactions. The same conceptions, when slightly modified, lend themselves to the explanation of the influence of mass

and to that of reversible reactions.

At the same time as these hypotheses concerning the mechanism of chemical processes were developed, Guldberg and Waage (Etudessur les Affinites chimiques, Christiania, 1867) laid the foundamon for the exact development of the theory of affinity by establishing a mathematical law for the influence of mass. Their work marks an epoch in the history of affinity. It was they who first gave a possibility of determining numerically the intensity of chemical affinities, though at first only as relative magnitudes.

The law established by these two investigators states 'that chemical action is proportional to the active masses of each of the substances participating in the reaction.' By 'active mass' is understood that quantity of the substance participating in the reaction, measured in equivalents or in molecular weights, which is contained in unit volume of the system. This is the same idea as underlay the views of Berthollet; it is the same, only freed from the errors which were attached to it in his time and thrown into an exact mathematical form. Guldberg and Waago lay special stress on the fact that, when the action of several substances on each other is proportional to the active mass of each separately, the intensity of the reciprocal actions among the substances is measured by the product of these masses.

In their earlier paper Guldberg and Waage developed the equations for the chemical equilibrium of opposite reactions by putting the chemical forces as proportional to the product of the active masses. On the other hand, as proportional to these same forces they put the velocities of the reactions, i.e. the relation between the quantity of substance changed and the time necessary to effect the change. In their later papers it was found better to refer the considerations, not to the forces, but to the velocities of the reactions, since these are capable of exact definition, and to leave out of account altogether the somethat vague idea of chemical forces. According to this conception, chemical equilibrium results when the velocities of the opposite reactions have become equal, i.e. when the quantity of substance undergoing a certain change is equal to the quantity formed by the reverse process. It is evident that this is the same theory as had been framed by Williamson, and developed by Pfaundler. The empirical law of the influence of mass thus receives a reasonable foundation in the molecular theory of matter. The effect is proportional to the active mass, because the number of molecules which can react is proportional to the mass. On these lines Van't Hoff (B. 10, 669) has deduced the law of the influence of mass. He retains exactly the form given to it by Guldberg and Waage; and in a later paper (J. pr. [2] 19, 69) these authors accepted this formal improve-

In the following part I propose to give a short systematic review of chemical kinetics, or the doctrino of the course of chemical actions, and to deduce from it the conditions and equations of chemical equilibrium; further on, the theoretical conseptions thus gained will be used in the practical determination of the magnitudes of an antities. The theoretical part is based mainly

on the paper of Guldberg and Waage mentioned above. A book recently published by Van't Hoff (Etudes de Dynamique chimique) is also of importance and has been of great use to the author, though he by no means agrees with the whole of its contents.

CHEMICAL KINETICS .- Let us suppose some substance to be undergoing chemical change. Then in any time, dt, some quantity, dx, will have been changed. We define the velocity of the chemical reaction, c, as the ratio of the quantity changed to the time taken for the change, and we therefore put $c = \frac{dx}{dt}$. The quan-

tities of the reacting substances are in every

case measured by formula-weights.

The quantity of substance changed will be dependent on many conditions. Among these we find such as are constant, or can be kept constant, during the whole process. Such are temperature, pressure, volume, &c. One condition, namely, the quantity of substance undergoing change, necessarily varies during the pro-cess, and we have to find an expression for the velocity of the change as a function of this *quantity. This may be dene by putting $c = \frac{dx}{dt} = k f(x_2x_1)$,

$$c = \frac{dx}{dt} = k f (x_2 x_1),$$

where the constant factor k represents the constant conditions, and $f(x_1x_2)$ the conditions varying with the quantities x_1, x_2, \dots Concerning the ferm of the function $f(x_1x_2)$ information has been sought and found in various ways. the results arrived at are concordant, and show the function to be one of direct proportion. This rosult has been arrived at empirically, and also theoretically as a deduction from various assumptious. Guldberg and Waage did the first; Horstmann, and others after him, showed that for oertain cases, especially for gaseous compounds, tho law of direct proportionality between chemical action and mass follows as a necessary conscquence from the second law of thermodynamics. Gibbs has made this result perfectly general. The kinetical theory of the constitution of matter leads to the same result, by considering the probabilities for the occurrence of those arrangements of particles which make chomical change possible.

Before, however, the proof for the law of direct preportion between chemical action and mase can be attempted we must distinguish betwoen the various types of chemical reactions. As a rule, chemical action does not take place in the presence of one substance only, but more than one is needed to bring about the final result. Since the final result is preportional to the mass of each separately, we have quite generally $\frac{dx}{dt} = k, x_1, x_2, x_3, x_4, \dots x_n$

$$\frac{dx}{dt} = k, x_1, x_2, x_3, x_4, \ldots x_n$$

where $x_1, x_2, x_3, \ldots x_n$ are the quantities of tho various substances, and x the quantity of substance formed by their reaction. All these quantities are measured according to the ratios of the molecular weights of the various substances.

The eimplest case is that in which one substance only undergoes change during the reaction, or that in which the change of only one substance has to be taken into account. The first case occurs when, for example, a substance

is decomposed, or when it undergoes a molecular transformation; the second occurs when the other substances participating in the reaction are present in snoh quantities that the diminntion of them occasioned by the chemical change is not appreciable. Since, then, the factors $x_2, x_2, \dots x_n$ disappear or become constant, we have

 $\mathbf{c} = \frac{dx}{dt} = k x_1.$

The velocity of the reaction at any instant is preportional to that quantity of substance undergoing change which is still within the sphere of action.

This equation was first established by Wilhelmi (P. \$1, 413) in 1850 for the inversion of cane-sugar. Since that time it has been confirmed in many ways. In order to compare it with the empirical results it must be integrated. If we put the quantity of substance present at the beginning of the change = a, then, after any time t, a quantity x will be decomposed, and, sinco we are dealing with molecular units, x will at the same time represent the quantity of the product of decomposition formed. The quantity x_i of substance present at the time t is $(a-x)_i$ and we have therefore

$$\frac{dx}{dt} = k (a - x)$$

$$\frac{dx}{a - x} = k.dt$$
(1)

 $-\log(a-x)-k.t+\mathrm{constant},$

where log represents the natural legarithm. In order to determine the constant of integration, a definite pair of values for x and t must be taken. We put both simultaneously equal to zero, i.e. we begin the time from the instant when the decomposition begins, and thence we get

 $-\log a = \text{constant}$, and $\log a - \log (a - x) = \log \frac{a}{a - x} = kt.$

In order to pass from the natural to the common logarithms we have only to multiply the constant k by its modulus.

Some series of experiments illustrative of this equation will now be given. The first of these refers to a simple decomposition, the change of dibremosuccinie acid into dibromomaleic acid and hydrebromic acid,

 $C_2H_2Br_2(COOH)_2 = C_2HBr(COOH)_2 + \overline{H}Br_4$ a chango which occurs in aqueous solution at 100°. This reaction has been studied by Van's Hoff (Etudes de Dynamique chimique, p. 13). The progress of the decomposition can be traced by titrating with standard alkali. The amount ef alkali required for neutralisation before the action began was 10.25 (arbitrary units), and when the change was finished the amount of alkali was 15.32. The creess of alkali nsed, over 10.21, at any stage of the change is put as equal to x. For a we have 5.11, since two equivalents of dibremosuccinio acid give one equivalent of hydrobremic acid. Van't Hoff's results are presented in the table given on next page. The constancy of the value of k, as chown in the last column, furnishes a proof of the truth of the equation used.

A eccond eeries of experiments dealing with the inversion of cane-sugar by means of sulphuric acid is taken from the experiments of W. Ostwald (J. pr. 29, 394). The action is one of

# (minutes)	units alkali		log a-s	
	10:21	0.00	7_	_
180 × 2	10.53	0.82	0-0281	0.0141
4	10-79	0.58	0.0524	0.0131
. 6	11.05	0.84	0.0776	0.0128
. 8	11.25	1.04	0.0088	0.0124
10	11.55	2.34	● 0.1320	0.0132
18	11.94	1.73	0.1795	0.0138
16	12.29	2:08	0.2269	0.0143
. 19	12.53	2.31	0.2613	0.0138
32	12.84	2.63	0.3116	0.0143
26	13.03	2.82	0.3187	0.0131
30	13.30	3.09	0.4027	0.0135
84	13.57	8.36	0.1647	0.0137
. 89	13.71	3.20	0.5009	0.0129
45	14.05	3.84	0.6038	0.0135
52	14:32	4.11	0.7077	0.0137
60	14.52	4.31	0.8041	0.0135
71	14.69	4.48	0.9066	0.0128
90	15.03	4.82	1 2 1 1 1	0.0133

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the type called catalytic, i.e. the substance which brings about the change (in this case sulphurie acid) does not itself undergo any change. sugar, by taking up water, is changed into dextrose and lævulese. The rotatory power changes in the same preportion as the decomposition proceeds; the course of the change can therefore be traced by means of the polariscope. The original solution rotated 25°04'; when completely inverted it rotated -8°15', so that the total angle passed through was 33°19'; this number is at the same time the measure of the total amount of sugar, and must therefore be put =a. For x we have the difference $25^{\circ}01'-w$, where w is the angle of retation produced by the solution undergoing change at time t.

(minutes)	w	x	$\log \frac{a}{a-x}$	k
58	20020	4.81	0.0684	0.0001180
114	16028	8.76	0.1331	1163
197	11°34	13.70	0.2313	1175
263	8°30	16.74	0.3052	1161
894	3°35	21.69	0.1603	1169
585	-1°39	26:43	0.6909	1182

Similar experiments have been made for many other cases and have given like results. For non-reversible chemical reactions, which depend on the quantity of a single substance only, tho shove formula holds good quite generally; it can however be proved only in the case of reactions which are sufficiently slow to allow of measurement.

A second main division of chemical processes is formed by these which involve the presence of two different substances. In such cases the general equation (1) takes the form

$$\frac{dx}{dt} = k.x_1.x_2 \tag{2}$$

Two cases have to be distinguished here; either the quantities of the reacting substances are equivalent with respect to the chemical change considered, or one of thom is present in excess. Putting the quantities = a and b, then either -bor a > b. In the first case we have to put

$$\frac{dx}{dt} = k(a-x)^2 \tag{3}$$

and integrating

$$\frac{x}{a-x}$$
 = a.k.t

where the constant of integration is determined

on the same suppositions as before.

Decompositions of this type have often been, investigated experimentally. The example given investigated experimentally. All parameters here is a series of experiments by Ostwald (J. pr. 27, 1), on the docomposition of acetamide by acetamide by trichloracetio acid. This acids, especially by trichloracetic acid. change takes place according to the equation:

 CH_3 .CONH₂ + CCl_3 .CO₂H + $H_2O \Rightarrow$ CCl₃.COONH₁ + CH₃.CO₂H.

The quantities both of acetamide and trichloracctic acid diminish, with production of inert ammonium trichleracctate, and acetic acid, which acid under the conditions of the experiment exerts little or ne influence on the change. In the following table t stands for the time in minutes, x for the quantity of ammonium salt formed (or, what is the same thing, for the quantity of acetamide decemposed) measured in units such that the total quantity a = 26.80.

. ,	x	$\frac{x}{a-x}$	ak
15	3.13	0.132	0.0088
30	5.52	0.260	0.0087
45	7.61	0.397	0.0086
60	9.23	0.525	0.0088
90	12.01	0.811	0.0090
120	13.82	1.065	0.0089
150	15.51	1.375	0.0092
180	16.59	1.623	0.0090
240	18.33	2.169	0.0090

These results show that the quantity

$$ak = \frac{1}{t} \cdot \frac{x}{a-x}$$

is constant, as is required by theory.

A second scrics of experiments by R. Warder (B. 14, 1361) on the suponification of ethylic acctate gave similar results. Equivalent quantities of ethylic acctate and seda were mixed. and portions taken from time to time were neutralised by dilute acid. The quantities of acid used-these will be called s-give the quantities of the substances not yet decomposed. In each experiment the soda alone would have used 16.00 c.c. of acid; hence we have a = 16.00 and x = 16 - s, therefore also a - x = s

t (minutes)		x	x a-x	a.k
5	10.24	5.76	0.563	0.113
15	6.13	9.87	1.601	0.107
25	4.32	11.68	2.765	0.108
35	8.41	12.59	8.69	0.108
55	2.31	13.69	5.94	0.108
120	1.10	14.9	₹¥:55	0.113

Again ak is sufficiently constant. When a and b are different, x, becomes a-x, and x, becomes b-x; then

$$\frac{dx}{dt} = k(a-x)(b-x) = k\{x^2 - (a+b)x + ab\}$$
 (5)

the integral of this equation is

$$\log \frac{(b-x)a}{(a-x)b} = (a-b)kt.$$

The validity of this equation has been proved by T. Flood (P. M. [5] 6, 871).

These two types of non-reversible chemical notions which have been just considered comprise all non-reversible actions which have been accurately studied. To be consistent, we must assume that in ohemical reactions which involve more than two, say three, substances, an squation corresponding with those given ought to hold good. Thus when three substances are present

in equivalent quantities, $\frac{dx}{dt} = k(a-x)^3$; and

$$\frac{1}{2} \cdot \frac{2ax - x^2}{(a - x)^2} = a^2 \cdot kt. \tag{7}$$

But no reaction has been observed with sufficisnt certainty the course of which proceeds according to this, or according to a higher, equa-

Moreover, a complication may arise from the simultaneous occurrence of several reactions. For such a case the principle of the 'coexistence of reactions' is important; this principle states that every reaction proceeds as if it alone took place. This principle is of paramount importancs; it forms the connecting link between the simple reactions, and those of so-called chemical equilibrium. For the mathematical expression of the coexistence of reactions, when one and the same substance is affected by the various changes, we have the following,

$$\frac{dx}{dt} = k \cdot x_1 x_2 \cdot \ldots + k' x_1'' x_2'' \cdot \ldots + k''' x_1''' x_2''' \cdot \ldots$$

If, however, the coexistent reactions take place among different groups, which are themselves without effect on each other, the equation of velocity has to be developed for each separately without regard to the others.

No experimental investigation of the law of coexistence has as yet been published. The application of this law in the theory of affinity leads, however, to results which agree with expsrience, and the law may therefore be considered to be experimentally proved.

Rsysrsible Reactions.

The processes investigated above frequently represent only one part of the actual reactions. In many cases the substances formed mutually react to reproduce the original substances. In such cases the process does not end with decomposition; but a permanent final state is arrived at in which the original substances, as well as the products of their double decomposition, are coexistent. In such a case the final systsm is said to be in chemical equilibrium. Here we have to consider on the one hand the velocity of the reaction, on the other hand the proportion of the masses for which chemical equilibrium results. As aids in the first part of this inquiry we have the equations given abovo, together with the principle of enexistence: in investigating the second part of the problem we have the following condition;—chemical equilibrium results when the velocities of the opposite reactions have become equal.

The establishment of chemical equilibrium is connected with the second type of chemical roactions [equations (3) (4), and (5) (6)], respectively.

$$\frac{dx'}{dt} = k'x_1'x_2', \text{ and } \frac{dx''}{dt} = k''x_1''x_2''$$

and equilibrium results when

$$\frac{dx'}{dt} = \frac{dx''}{dt} \text{ or } k'x_1'x_2' = k''x_1''x_2''.$$

This is the equation first established by Guld. berg and Waago. Putting the initial quantities of the substances as p, q, p', and q', when the substances p' and q' are formed by the reaction of p and q and vice versa, equilibrium will result when a certain quantity ξ of p and q has been decomposed. Then the quantities $p-\xi$, $q-\xi$, $p'+\xi$, $q'+\xi$ are in equilibrium; and ξ has the same value throughout, since the quantities p, q, &c. are measured according to equivalents. The quantity ξ may be positive or negative.

The equation of equilibrium then takes the

following form :--

$$k'(p-\xi)(q-\xi) = k''(p'+\xi)(q'+\xi);$$
 (8) from which a value for ξ is found,

$$\xi = \frac{k'(p+q) + k''(p'+q')}{2(k'-k'')} \mp \sqrt{\left(\frac{k'(p+q) + k''(p'+q')}{2(k'-k'')}\right)^2 + \frac{k'p'q' - k'pq}{k'-k''}}$$
The minus size helds when $\frac{k'}{k'} = \frac{k'}{k''} = \frac{k'}{$

The minus sign helds when $\frac{k'}{k''} > 1$

By making certain assumptions, this expression may be considerably simplified. If at the beginning of the reaction the substances p and q only are present, in equivalent quantities, p = q and p' = q' = o, and it follows that

$$p=q$$
 and $p'=q'-o$, and it follows that $k'=\frac{\xi^2}{k''-(p-\xi)^2}$ and $\xi=p$

$$\sqrt{\frac{k''}{k''}}$$

$$\sqrt{\frac{k''}{k''}+1}$$
The equation of velocity takes the following ds

form. The resultant velocity $\frac{dx}{dt}$ is the difference of the partial velocities:

ence of the partial velocities:
$$\frac{dx}{dt} = k'(p-x)(q-x) - k''(p'+x)(q'+x) \quad (10)$$
Introducing a new constant h we have
$$\frac{k}{h} = \frac{k'(p+q) + k''(p'+q')}{k'-k''-k''-\xi};$$
the equation can then be brought to the form
$$\frac{dx}{dt} = (k'-k'')(\xi-x)(h-x);$$
and from this by integration, we obtain

$$\hat{h} = \frac{k'(p+q) + k''(p'+q')}{k' - k''} - \xi;$$

$$\frac{dx}{dt} = (k'-k'')(\xi-x)(h-x);$$

and from this by integration, we obtain
$$\log \frac{h-x}{\xi-x} \frac{\xi}{p} = t(h'-k')(h-\xi)$$
 (11)

The form of this equation is analogous to that already deduced for simple reactions (equation (6), p. 72). If now it is assumed, as before, that p=q, and p'=q'=0, ξ assums the simple form given in (9a)

and
$$h=p$$

$$\frac{\sqrt{\frac{k'}{k''}}}{\sqrt{\frac{k'}{k''}-1}}$$

The preceding equations are dednotions, from the laws of the influence of mass, and the coexistence of chemical remations. Both For the vslooity of each of the reactions we have plaws are of about the same importance in the theory of affinity as the laws of gravitation and , the coexistence of motions are in astronomy.

Each individual practical case really comprises several different relations; but there are combinations in which so great a part of the result depends on one single cause, only, that the observed phenomena may be represented almost as it this were the only cause. And as little as we doubt the law of gravitatiou because the motions of the moon cannot yet be expressed completely in equations, so little have we cause to doubt the laws stated above because certain phenomona cannot yet be represented as simple deductions from them.

We have hitherto assumed that the constant of velocity does not alter its value throughout the whole reaction. It is, however, not impossible that reactions exist in which the reason for the change of k is to be found in the chenical process itself; in such cases the problem becomes considerably complicated.

Chemical Dynamics.

In general dynamics the magnitude of any force is defined and measured by the velocity which it imparts to a mass of known magnitude. Another way of measuring forces consists in establishing equilibrium between the given force and a force acting in the opposite direction, which latter is of a magnitude already known or easily determined. This can be considered as a special case of the first method, as a case in which the velocity due to the given force is reduced to nothing, by means of one equal in magnitude but opposite in direction. The second method, though not a direct one, possesses all the important experimental advantages belonging to a zero method and is therefore the more usual. The measurement of the intensity of chemical forces can be accomplished by two methods, analogous to those employed in general dynamics. The more usually employed method (because of experimental advantages) is the statical, or the method of equilibrium, in which a chemical process is reduced in a certain proportion by another action occurring in the opposite direction. This corresponds to the statical method used in measuring mechanical forces. Analogous to the kinetical method, of the method of velocity, is the process of obtaining a measuro of the intensity of the acting forces by measuring the velocity of the chemical change. The two chemical methods are connected in a similar manner as the two mechanical methods, since, as has been shown above, the equilibrium of chemical changes can be regarded as the consequence of the mutual counteraction of changes which are equal in magnitude, but opposite in direction.

Statical Methods.

The first attempts to measure affinitios woro made by Wenzel, in 1777 (Die Lehre von der Chemischen Verwandtschaft, p. 28, Dresden, 1777). He used the method of velocities, but his process was very imperfect. His experiments related to the solution of metals by various acids. Later experiments dealt almost exclusively with the affinity between acids and bases, and were mostly carried out by the method of equilibrium.

Solutions of acids and bases were mixed in proportions such that different acids competed for an insufficient amount of a base, or vice

gersd; and an attempt was then made to delermine the distribution of the base between the competing acids (or vice versa). Ordinary analyses could give no information as to this distribution; since such analyses oculd determine only the absolute quantities of the acids and bases, and not their distribution. Steinheil (A. 48, 153) (although with an entirely different aim) was the first to show how we must proceed in order to get a knowledge of the arrangement of the constituents of a solution without interfering with its composition. Since each of the constituents of a solution changes the physical properties-such as density, refractive index, colour, &c .- of the solution, a knowledge of the laws governing these changes indicates how to solve the problem, by measuring a sufficient number of constants and forming the necessary

It has been already explained how Gladstone used these means for establishing a number of facts concerning the chemical statics of solutions of salts. He could not, however, utiliso his measurements further, as there did not then exist a general theory of chemical affinity. Such a theory was first given by Guldberg and Waago (Etades sur les Affinités chimiques, Christiania, 1867) and was also applied by them to a number of measurements. This theory has met with such wide confirmation that we hope to be able some day to reconcile with it those facts which do not appear at present to be in keeping with it.

J. Thomsen (P. 138, 65) was the first to apply the theory of Guldberg and Waage to the case of homogeneous solutions. He found that sulphuric acid when acting on soda gives a heatproduction of 31,378 gram-units, while nitric acid gives 27,234 units only. Now, when sulphuric acid and nitrie acid simultaneously act on soda, all three substances being present in equivalent quantities, three cases may arise. Either the sulphuric acid exclusively combines with the soda, or the nitric acid exclusively does so, or the soda divides itself between the two acids in some fixed proportion. In the first case 31,378 gram-units of heat, and in the second case 27,234 units, would be produced, while in the third case the heat-production would be represented by a number between these two. Therefore the number found by experiment gives a measure of the distribution of the soda between the acids.

Similarly sodium sulphato is allowed to react with nitric acid. If no chemical action results there will be no production of heat. If the nitric acid combines with all the soda, liberating all the sulphuric acid, a disappearance of heat must result, numerically equal to the difference between the two heats of neutralisation; that is to say, 31,348-27,234-4,144 units of heat will disappear. If, however, a division of the base between the acids results, a quantity of fleat less than 4,144 units will disappear. Experiment shows that 3504 nnits of heat are used; therefore the soda divides itself between the two acids.

If no secondary thermal action takes place between the substances used in the experiment, we can deduce directly from these numbers that

2504 4144 = 0.845 of the total quantity of sedium sulplate is decomposed. Free sulphuric acid docs, however, react with sodium sulphate, and the action is accompanied by production of heat. The extent of the reaction depends on the relative quantifies of the reacting substances present. Thomsen has measured this effect for a great many proportione, and has expressed his results by the equation:

$$Q = -\frac{n}{n+0.8}.8,300$$
 gram-units of heat;

where n represents the number of equivalents of eulphuric acid present for each equivalent of sodium sulphate. By the help of this formula Thomsen found empirically that soda divides itself in such a proportion that $\frac{1}{2}$ of it goes to the sulphurio acid, and $\frac{2}{3}$ of it to the nitric acid. Calculating the heat production on this supposition, the result is -3,547; experiment gives -3,504; the difference lies within the limit of experimental error.

Guldberg and Waage's theory givee an oxtremely eimple expression for this case. Sinco sodium sulphate and nitrie acid were present in equivalent quantities at the beginning of the reaction, but sodium nitrate and sulphuric acid were absent, we have to put in equation (8)

$$p = q = 1$$
 and $p' = q' = 0$;
this gives $k(1 - \xi)^2 = k'\xi^2$,
 $\frac{k}{k'} = \frac{\xi^2}{(1 - \xi)^2}$

As & is the quantity of soda combined with the nitrio acid, and $(1-\xi)$ that combined with the sulphurie acid, it follows that the ratio of tho velocities of the reaction is equal to the square of the ratio of distribution. Thomsen calls the endeavour of the acids to combine with bases the avidity of the acids, and defines it by tho ratio of distribution. According to this, the avidity of sulphuric acid is half as great as that of nitric acid, or putting the latter = 1, the avidity of sulphuric acid is = 0.5. Tho avidities are in the ratio of the square roots of the velocities of reaction.

Thomsen (Thermoehemis die Untersuchungen, i. 808) has made further experiments concerning the avidity of other acids towards soda, using a method similar to that described above. His numbers are given in the following table, where the avidity of nitric acid has been put = 1.00.

Hydrochloric acid .	= 1.00
Hydrobromie " .	0.89
Hydriodie " .	0.79
Sulphurie ,, .	0.49
Scionio ,, .	0.45
Trichloracetic	0.36
Orthophosphoric acid .	0.25
Oxalie , .	0.24
Monochloracetic ,, .	0.03
Hydrofluone ,, .	0.05
Tartaric ,, .	0.05
Citrio	0.05
Acetic	• 0.03

Borio, silicic, and hydrocyanic acids do not give any appreciable values.

Taking the squaree of these numbers we get the relative velocities of the reactions between tho acids and the eoda, which values cannot be obtained directly owing to their great magnitude.

The question now presents itself as to whether

soda is employed. Thomsen made similar ex-periments for hydrochloric and sulphuric acids, using different bases (P. 188, 497); viz. potash, ammonia, and magnesia, the oxides of manganese, iron, cobalt, nickel, zine, and copper; for the avidity of sulphuric acid he obtained numbers which increase from 0.5 up to 0.8, and vary, for the alkalis between 05 and 0.57, and for the bases of the magnesia series between 0.70 and 0.81. These results induced Thomsen to conclude that the relative avidity of acids depends on the nature of the base.

Berthelot (A. Ch. [4] 30, 516), however, raised the just objection that Thomsen's method does not allow of the measurement of the relative avidities without the introduction of errors. The free sulphurie acid reacts on the neutral salt, forming acid sulphate, and thus loses part of its active power, and it does this the more the greater the quantity of acid-sul-

phate which can be formed.

Thomsen's experiments were repeated by W. Ostwald (P. Ergzbd. 8, 167; J. pr. [2] 19, 468), who used a different method based on measuring the changes of volume which accompany chemical reactions in aqueous solutions. The volume of the solution of a salt is different from the sum of the volumes of the solutions of the acid aud the base, which by their mutual action produce the salt; and further this change of volume is different for different bases and acids. If we use solutions which coutain one gramequivalent of the acid or base per kilogram of solution then the volume of two equivalents of soda is 1913.26 c.c., and that of two equivalente of nitric acid is 1933.25 c.c.; the sum of these two is 3846.51 c.c.; but the volume of the corresponding solution of sodium nitrate is 3886 05 c.c., that is to say, 39.54 c.c. more than the sum of the volumes of acid and base. Repetitiou of the experiment with sulphuric acid gives an increase of volume of 29.96 c.c. ouly. Hence the volumechanges can be used for determining the composition of the solution, in the same manner as the heats of ucutralisation had been used by Thomsen. Results were obtained by this method exhibiting the behaviour of nitric acid and sulphuric acid towards soda; these results agreed entirely with those of Thomsen. Thomsen's conclusion that uitric and hydrochloric acids are stronger acids than sulphurio acid-a result opposed to the older views-was thus confirmed by Ostwald. The same chemist investigated at length the question as to whether the relative affinity of an acid varied with the nature of the basc. By experiments to which Berthelot's objection cannot apply, he arrived at the result that the relative avidity of an acid is independent of the base. Thus with hydrochlorie and nitries acids he found the following numbers :-

Potash			0.97
Soda .			0.96
Ammonia	. 4	٠.	0.96
Magnesia			0.99
Zine oxide		٠.	0.95
Copper ox	de		- 9⋅97

The differences are not greater that the probable errors of the experiment.

The question concerning the influence of the avidities thus found have constant values, temperature on the relativo affinities of acids or whether they change when a base other the has also been investigated by Ostwald. Has researed the expansion due to heating the same solutions which had served for the volumetric experiments with soda. The ratio of the avidity or affinity of hydroohloric and to that of nitric acid towards soda proved to be as follows:—

At 0° . 1.02 20° . 0.96 40° 4 0°98 60° . 1.00

In both cases the number for sulphuric acid is not quite constant; but, as already mentioned, this acid seems to be the etronger the less acid sulphate is formed, and vice versa.

At a subsequent time Thomsen (Thermoch. Uniters. i. 89) also attacked the problem of the influence of temperature on relative avidities, and arrived at the same results as Ostwald.

The refraction of light was used by Ostwald as another method for determining the couposition of a homogeneous solution by means of its physical properties. Dale and Gladstone (T. 1863, 317), and also Landolt (P. 133, 1), showed that the function v(n-1), in which n is the refractive index and v the specific volume of a liquid body, depends only on the elementary composition of the body and not on the temperature, nor (within certain limits) on the chemical arrangement of the constituents of the body. Hence the specific volume is inversely proportional to the refractive index diminished by 1, and the volume-changes attendant on chemical reactions must be accompanied by opposite changes in the refractive indices. Experiment has most fully borne out these conclusions. The optical method is, however, less accurate than the volumetric method, when the ordinary apparatus only is used.

The numerical results of Ostwald's experiments are collected in the following table. The numbers have the same meaning as those of Thomeen given before (p. 75), i.e. they give the relative avidities of the various acids, putting that of nitric acid = 1.

Acid		Thomsen
Nitrio	1.00	1.00
Hydrochloric .	. 0.98	1.00
Trichloracetic .	0.80	0.36
Dichloracetio .	0.33	_
Monochloracetic	0.070	0.03
Glycolic	0.050	_
Formie .	0.039	
Citric	0.033	_
Acetic	0.0123	0.03
Propionic	0.0104	1 —
Butyrie	. 0.0098	I —
Isobntyrio	0.0092	ł –
Succinic	0.0145	
Malio .	0.0282	I —
Tartarie	0.052	0.05

Thomsen's values have also been given as far as they refer to the acids considered here. They agree as well as can be expected; triabloracetic acid alone shows an appreciable difference. Thomsen's number for this acid is undoubtedly much too emall; this has been proved beyond doubt by other measurements.

The ratios of avidities given in the preceding

poissh, sods, or simmenis; they are independent of the nature of the base.

Besides these three methods which are of general application to the case of solutions, some investigators have employed others, which can, however, be used only in special cases. Thus G. Wiedemann (W. 5, 45) has shown that from the magnetic properties of ferric salts in eclution we can argue as to the amounts of these salts decomposed by the water into free soid and colloidal solublo iron oxide. This method is, however, restricted to the case of these special salts. A. Müller (P. Ergzbd. 6, 123) has drawn conclusions from the change in colour as to the distribution of iron oxide between hydrochloric acid and sulphuric acid. Jelett (I. 25, 871) determined the relative affinities of codeine, quinine, and brueine, by means of the rotation of the plane of polarisation, and found for the influence of mass the same law as had been established by Guldberg and Waage. The experiments of Dibbits (P. Ergzbd. 7, 462), Brücke (Sitz. W. 77, April 1877), and others, are of a more qualitative nature.

Besides these statical methods, based on the determination of the composition of a homogeneous solution, others are available in which the chemical reactions take place in heterogeneous media, viz. between solids and liquids, or liquids and gases, or lastly between solids and gasee. The theory of these methods has been also given by Guldberg and Wange, and Ostwald has developed the methods for the purpose of determining affinities.

According to Guldberg and Waage the chemical action of solids in contact with solutions is independent of their mass; in other words, the chemical mass of the solids is constant. Otherwise the lawe of the inflnence of mass hold good. If, for example, an acid acts on the salt of another acid, which latter salt is insolublo in water (or more strictly, searcely coluble), and with the base of which the first acid forms a soluble salt, then the same equation holds good as applies in the case of substancee which are all soluble, with this exception that the term corresponding to the insoluble salt becomes constant or independent of x. Putting in the equation k.p.q = k'.p'.q'p = hydrochloric acid, and q = calcium oxalate, then p' =calcium chloride, and q' =oxalic acid.

If the experiment is arranged so that hydrochloric acid acts on an excess of calcium exalate and that undiscolved calcium exalate is always present, then at all stages of the change exalic acid and calcium chloride are present in equivalent quantities. Putting the original quantity of hydrochloric acid=1, and that of the explate dissolved = the smaller becomes

the oxidate dissolved =
$$\xi$$
, the equation becomes $k(1-\xi)c = k'.\xi.\xi$; hence $\frac{k}{k'} = \frac{\xi^2}{c(1-\xi)}$, and $\sqrt{\frac{k}{k'}} = \phi = \frac{\xi}{\sqrt{c(1-\xi)}}$

where c stands for the constant chemical mass of calcium oxalate. In this equation k, k, and c are unknown, while can be measured directly. Repeating the experiment with a different said, say nitric acid, a new expression of the orm.

$$\phi_1 = \frac{\xi_1}{\sqrt{c(1-\xi_1)}}$$

is obtained, in which o has the same value as referring to calcium oxalate, which is used in both experiments under the same conditions. Dividing the one equation by the other we get the relative affinities

$$\phi: \phi_1 = \frac{\xi}{\sqrt{1-\xi}}: \frac{\xi_1}{\sqrt{1-\xi_1}}$$
in quantities which can all be direction.

expressed in quantities which can all be directly measured. This method has been used by W. Ostwald and his pupils for determining the relative affinities of various acids, and has given results which agree well with those found for homogeneous colutions. It has great experimental advantages over the physical methods, as the ordinary methods of chemical analysis oan be used. The insoluble, or searcely soluble, salts used were these; zinc sulphide, calcium oxalate, zinc oxalate, barium chromate, cream of tartar, and the sulphates of barium, strontium, and calcium. As a rule the coefficients of affinity thus determined for various salts agree very well amongst themselves, but there are some deviations which are not yet fully explained.

As an example of the method the following numbers are given (J. pr. [2] 28, 493); thesc numbers were obtained by the action of acids on calcium oxalate, a substance lending itself particularly well to these experiments. Experiments were made both with normal and decinormal solutions of acids. Nitrie acid is again put = 1.

Acid	Normal	10 Normal
Hydrochlorie	1.00	0.98
Hydrobromie	0.95	0.99
Nitrie	1.00	1.00
Chlorie	1.04	1.00
Sulphurio	0.70	0.74
Formio	0.0259	0.129
Acetio	0.0105	0.735
Monochloracetic .	0.051	0.213
Dichloracetic	0.183	0.188
Trichloracctic	0.612	0.899
Lactie	0.041	0.133
Succinio	0.0205	0.093
Malio	0.0505	0.121
Tartarie	0.0452	0.141
Citrie	0.0306	0.144

The numbors in the first column, which refer to normal solutions (one gram-equivalent in a litre), agree well with those before obtained by the volumetric method. Along with them is given a second series referring to solutions ten times as dilnte. While the stronger acids cxhibit scarcely any change by the dilution, the values for the weaker acids have increased very considerably, and this the more the weaker are the acids. We shall consider this phenomenon at length later on, and find the genoral law underlying it.

Of further results which have been arrived at by this method one must be noticed as im-portant, viz. that the action of the acids varies as they are present alone or along with thoir neutral_salts (J. pr. [2] 23, 209).

Some such result was to be expected in the

basic soids, however, exhibit no tendency to combine with their neutral salts, and yet they too show a change which in the case of the strong seide, such as hydrochloric and nitric, is an increase in the affinity. This increase is proportional to the quantity of the neutral salt present, and decreases rapidly with increasing dilution. This statement does not, however, hold for all monobasic acids, but only for the strong acids. The weak monobasic acids, on the contrary, are considerably more weakened by the presence of their neutral salts, and this the more the weaker are the acids. These facts are of great importance in the interpretation of experiments undertaken for the purpose of determining the relative affinities of acids by the division of a base between two competing acids. Since in this case the acids always act in presence of their own salts, this condition doubtlessly exerts some influence, making the strong acids appear stronger and the weak acids appear weaker. This shows that too much importance must not be attached to the numerical values obtained by the preceding methods; they certainly give the order of the affinities correctly, but the numerical values doviate from the true value in the sense that the large numbers are too large and the small values are too small. Wo shall see later on that other determinations of the same quantities, which are more likely to give the true values, show deviations in this sense from the above numbers.

Kinetical Methods.

The second way of measuring the intensity of chemical forces is based on determinations of the velocities of the reactions produced by these forces. The theoretical introduction concerning this method has been already considered, and we have seen that many reactions proceed according to a course which agrees well with that calculated from the influence of mass.

This method does not, however, lend itself to direct applications to the majority of the reactions investigated by the statical method. This statement applies particularly to the effects of affinity between acids and bases, because these processes are of too short duration to allow of measurements being accurately made of their velocities. If, however, the magnitudes in question can be measured by kinetical methods, this is because of a general and important principle.

It has been already shown that the nature of the base exerts no influence on the rolative avidities or affinities of the acids which react with the base. If the affinity between an acid a and a base b is designated by f(a, b), then the

following equations hold good: $\frac{f(a,b)}{f(a',b)} = \frac{f(a,b')}{f(a',b')} = \frac{f(a,b'')}{f(a',b'')}$

These equations can be true only if each ex-

pression f(a, b) is the product of two factors one of which depends on the acid only and the other on the base only $f(a, b) = \phi(a) \cdot \psi(b)$.

The affinity between acids and bases is therefore the product of specific affinity-of-efficients. All reactions due to acids and bases s such must, on this view, be proportional among themselves. From this it follows that sentral salts to form said salts. The mono proceeses which, taken by themselves, have

nothing to do with the formation of salts, may be employed for finding numerical values for the affinities which come into play during the formation of salts, provided the roactions in question have been accomplished by the acids and bases only. Determinations of the specific affinity-coefficients of acids and bases are thus of the greatest importance. It will be our task to show first that the above conclusions are verified by experiment, and then to use tho numbors thus arrived at for drawing further deductions.

The first reaction which was used to check the values of the occflicients of affinity of acids determined in the statical way, by means of a kinetical method, was the change of acetamide into ammonium acetate (Ostwald, J. pr. [2] 27, 1), which takes place according to the equation $CH_3CONH_2 + H_2O = CH_3COONH_4$. water only is present the reaction does not preceed to a sensible extent, but when an acid is added the latter exerts a predisposing influence, and the precess takes place to the degree which is possible under the existing conditions of affinities, concentration, and temperature.

By 'predispesing affinity' is usually understood the cause of reactions between certain substances, which reactions could take place, but do not actually occur, without the presence of another substance, which has affinity towards one of the possible products of the reaction. In the case just discussed, water and acctamide do not react unless an acid capable of combining with the ammonia produced (or a base which has affinity for acctic acid) is present. The strange assumption expressed in the name predisposing affinity, viz. that the affinity of the predispesing substance tewards a bedy not yet formed induces the other substances to produce this special body, has been given up, since the molecular theory of Williamson and Clausius, as developed by Pfaundler, gives a much more simplo view of such reactions. When applied to the case just considered, this theory tells us that the atoms which form the molecules of acetamido and water only very seldem get a chance of forming ammonium acctate during the movements and cellisions of the molecules in question, since the forces which tend to retain the original condition of the system are greater than the forces which tend towards decomposition. If, however, a strong acid or base is added, the forces tending towards decomposition are correspondingly increased, and, in many collisions, in which previously no change occurred, decomposition now takes place (v. also Mcndeléeff, B. 19, 456).

The experiments were conducted by keeping equivalent quantities of acetamide and acid for some time at 65° and 100°. The quantity of ammonium salt formed was determined by decomposing it with sedium hypebremite and measuring the volume of nitregen evolved. The reaction takes place in the presence of acids, such as hydrochlor catid, according to the equa-

CH_CONH_2 # H2O + HCl = CH_COOH + NH_Cl. Three different kinds of melecules are therefore acid, undergo an appreciable change of mass relative affinities:-

during the reaction. The water is present in such excess (about 800 H2O to 1 HCl) that the change in its mass is unappreciable. Hence, if there are no secondary reactions, the change will proceed according to equations (3) and (4). An example in which the actual reaction agrees with theory has already been given.

But the reaction is by no means free from secondary changes. Partionlarly (as has been already noticed), the presence of the neutral ammonium salt of the acid added has the effect ef inercasing the strength of strong aoids, and decreasing the strength of weak acids. Consequently when streng acids are used the process is accelerated, compared with its normal value, and the acceleration is the greater the further the change has proceeded. With weak acids, on the other hand, the process is retarded. Owing to the fermatien of acid salts, the polybasic acids are influenced by similar but much more preneunced secendary reactions. All these conditions have to be taken into account in the investigation of the progress of the reaction, as is scen mest conspicuously in the graphical representation given in the original paper. These circumstances are disadvantageous if it is desired to make absolute determinations of the velocity of the reaction, but they are of advantage in the comparison of the kinetical and statical methods, since the results obtained by the latter are also influenced in the same way by similar sources of error.

The time taken to convert half the acetamide into the ammenium salt was taken as the reciprecal measure of the velocity of the reaction. The velocity in minutes was found to be as fol-

Acid	At 650	At 1000	Ratio
Hydrochlorie . Nitrie Hydrobromie .	72·1 75·2 74·0	4·98 5·35 5·14	14·5 14·4 14·4
Trichloracetic Dichloracetic .	112·8 433·7	=	=
Menechloracetic . Formic	4,570 28,950	2,138	13.6
Sulphuric Oxalic	180 1,516	14·1 118·6 929	12·8 12·8
Malio	35,310	7,976	14.7
Citric	44,810	3,088 3,880	14.5
Arsenio	_	4,005	_

In order to make these numbers comparable with the coefficients of affinity as found by the velumetric method they must be referred to IICl = 1, by dividing the times corresponding to the several acids each into that corresponding to hydrechleric acid. The relative velocities of the reaction are thus obtained for hydrochloric acid = 1. Further it must be borne in mind that by theory the ratio of the affinities is equal to that of the square roots of the velocities of the reaction. In the following table I have collected the acids the relative affinities of which are known. always necessary for the reaction. Mereovera Under I. are given the velocities of the reaction, only two substances, acctamido and hydrochlorio under II. their square roots, and under III. the

	Acid	87	ti.	in.
	Hydrochloric .	1.00	1.00	0.98
	Nitrio	0.96	0.98	1.00
•	Hydrobromic .	0.97	0•98	0.05
	Trichloracetic .	0.689	0.80	0.80
	Dichloracctic .	0.166	0.41	0.33
	Monochloracetic	0.0169	0.13	0.07
	Formic	0.00266	0.052	0.039
	Acetic	0.000547	0.0234	0.0123
	Sulphuric	0.428	0.65	0.67
	Tartaric	0.00564	0.075	0.052
	Malic	0.00218	0.0467	0.0282
	Succinic	0.00065	0.0255	0.0145

The numbers in the two last columns agree as well as could be expected. The deviations are in the direction of a greater value for II. than for III. in the case of weak acids. The reason for this lies in the fact that in the enunciation of the equation of velocity no attention was paid to the acetic acid formed in the reaction, by the presence of which the change is accelerated. This action of acetic acid scarcely comes into

play when strong acids are employed. The examination of the action of acids on the changs of acetamide into acetic acid and ammonium salt has established the connection between equilibrium and velocity which is predicted by theory; but the reaction employed was not of a kind to give completely accurate values for the velocity of the change, since too many secondary reactions exert their influence on the primary process. Another reaction etudied by Ostwald (J. pr. [2] 28, 449) lends itself better for this purpose. This is the decomposition of ethereal salts by water in the presence of acids. Aqueous solutions of methyl acetato (or of similar com-pounds) undergo ouly very slow decomposition at ordinary temperatures; if, however, an acid is present the process is greatly accelerated. The acid does not undergo a permanent change, since at the snd of the reaction exactly the same quantity of acid is found as was present at the beginning. It is doubtless by its affinity for the mothyl alcohol that the acid influences the rate of the change. It predisposes in the sense already explained, only the compound which the acid forms with the methyl alcohol cannot exist in the presence of the great excess of water. The chemical process is represented by the equation CH,COOCII, + H,O = CH,OH + CH,COOII. Two substances are required; but the quantity of water is so great that its change need not be taken into account. Equations (1) and (2) must therefore hold good. This conclusion is verified by experiment. Thus for example 10 c.c. of normal hydrochloric acid were mixed with 1 c.c. of mothylacetate and diluted with water to 15 c.c. One c.c. of this solution required for neutralisation 13.33 c.c. of baryta. Owing to the decomposition of the methyl acctate the acidity increased; the results are given in the first table of next column. The numbers in the last line reprosent the results when the decomposition was completed.

In the third column, under x, is given the increase in the number of c.c. of baryta used to neutralise the acid; the values in this column coots of the velocities of reaction; these numerical arealways proportional to the quantity of methyl-bers ought to be proportional to the affinities, if

acetate decomposed. The last value 14:11 gives the quantity a in the equation

Calculating the expression log $\frac{a}{a-x}$ (for simplicity's sake in ordinary logarithms), and dividing it by the time t, # (=the coefficient of velocity) is obtained; the value of k is given in the last column; it is nearly a constant.

		x	£
After 14 minutes	14.25	0.92	0.00209
34 "	15.47	2.14	0.00211
59 "	16.85	3.52	0.00212
89 ,,	18.24	4.91	0.00209
119	19.48	6.15	0.00209
159 ,,	20.92	7.59	0.00211
199 ,,	22.15	8.82	0.00214
239 ,,	23.10	9.77	0.00214
299 ,,	24.21	10.88	0.00214
399 ,,	25.46	12.13	0.00214
539 ,,	26.42	13.09	0.00213
∞ ″	27.44	14.11	

The same method was used for determining the velocity of decomposition of methylacetate by many other acids; the coefficients, referred to HCl = 1, are collected in the following table:-

Hydrochloric 1-00 1-00 Hydrochloric 0-98 0-99 Hydriodic 0-96 0-98 Hydriodic 0-96 0-98 Hydriodic 0-92 0-96 Chloric 0-92 0-96 Chloric 0-94 0-97 Sulphuric 0-517 0-739 Methylsulphuric 1-00 1-00 Ethylsulphuric 0-98 0-99 Propyleulphuric 0-98 0-99 Isobutylsulphuric 0-97 0-98 Ethylsulphonic 0-98 0-99 Ethylsulphonic 0-98 0-99 Benzenesulphonic 0-98 0-99 Benzenesulphonic 0-99 0-99 Formic 0-0131 0-115 Acetic 0-0345 0-0551 Brityric 0-00345 0-0551 Brityric 0-0039 0-0551 Isobutyric 0-00299 0-0551 Isobutyric 0-00298 0-0518 Monochloracetic 0-430 0-208 Dichloracetic 0-682 0-826 Lactic 0-00901 0-949 Hydroxyisobntyric 0-0021 0-0960 Trichlorolactic 0-0969 0-263 Pyruvic 0-067 0-259 Oxalic 0-1746 0-430 Malonic 0-0287 0-169 Succinic 0-00196 0-0704 Malic 0-0115 0-1279 Citric 0-0105 0-1279 Citric 0-0105 0-1279 Citric 0-0105 0-1279 Citric 0-0105 0-1279 Condend 0-0127 0-1279 Condend 0-0287 0-1515 Condend 0-0127 0-1279 Condend 0-0127 0-1279 Condend 0-0127 0-1279 Condend 0-0127 0-1279 Condend 0-0287 0-1515 Condend 0-0127 0-1279 Condend 0-0127 0-1279 Condend 0-0287	∆cid		I.	II.
Hydriodic 0.96 0.98 Nitric 0.92 0.96 Nitric 0.92 0.96 Chlorie 0.94 0.97 Sulphuric 0.517 0.739 Methylsulphuric 1.00 1.00 Ethylsulphuric 0.98 0.99 Isobutylsulphuric 0.98 0.99 Isobutylsulphuric 0.96 0.98 Isobutylsulphuric 0.96 0.98 Isobutylsulphuric 0.96 0.98 Isoamylsulphuric 0.96 0.99 Isethionic 0.98 0.99 Isethionic 0.98 0.99 Isethionic 0.99 0.99 Isethionic 0.90 0.99 Isethionic 0.90 0.95 Isobutyric 0.00345 0.0587 Propionic 0.00304 0.0551 Isobutyric 0.00268 0.0518 Monochloracetic 0.430 0.208 Dichloracetic 0.682 0.826 Lactic 0.00901 0.0949 Hydroxyisobntyric 0.00921 0.0960 Trichlorolactic 0.0267 0.258 Pyruvic 0.067 0.258 Pyruvic 0.0287 0.169 Succinic 0.00196 0.7046 Malio 0.01181 Tartaric 0.0296 0.1515 Racemic 0.0517 Carlorionic 0.00196 0.7046 O.01515 0.05151 O.0296 0.1515 O.0297 0.1515 O.0297 0.1515 O.0298 0.0298 O.0298 O.0298 O.0298 0.0298 O.0298 O.0299 O.0290 O	Hydrochlorie .		1.00	1.00
Nitrio	Hydrobromio .		0.98	0.33
Chlorie	Hydriodic		0.96	0.98
Sulphurio	Nitric		0.92	
Methylsulphuric 1.00 1.00 Ethylsulphuric 0.98 0.99 Propyleulphuric 0.98 0.99 Isobutylsulphuric 0.96 0.98 Isoamylsulphuric 0.98 0.99 Isoamylsulphuric 0.98 0.99 Isethionic 0.98 0.99 Benzenesulphonic 0.99 0.99 Formic 0.0131 0.115 Acetio 0.00345 0.0587 Propionic 0.00304 0.0557 Propionic 0.00299 0.0551 Isobutyric 0.00298 0.0518 Monochloracetic 0.0430 0.208 Dichloracetic 0.682 0.826 Iactic 0.00901 0.0440 Hydroxyisobntyrie 0.00921 0.0960 Pyruvie 0.067 0.253 Oxalio 0.1746 0.430 Malonic 0.0287 0.169 Succinic 0.00181 0.1515 Racemic 0.1515 </td <td>Chloric</td> <td></td> <td>0.94</td> <td></td>	Chloric		0.94	
Ethylsulphuric	Sulphuric		0.517	
Propylealphuric 0.98 0.99 1sobutylsulphuric 0.96 0.98 0.99 1soamylsulphuric 0.96 0.98 0.99 1sethionic 0.98 0.99 1sethionic 0.99 0.95 0.99 0.95 0.99 0.95 0.99 0.95 0	Methylsulphuric		1.00	
Propyleulphuric	Ethylsulphuric.		0.99	
Isobutylsulpluric 0.97 0.98 Isoamylsulpluric 0.96 0.98 0.99 0.99 Isethionic 0.98 0.99 0.99 Isethionic 0.98 0.99 0.99 0.90			0.98	0.99
Isoamylsulphurio 0.96 0.98 Ethylsulphonic 0.98 0.99 0.99			0.97	
Ethylsulphonic . 0-98 0-99 Isethionio . 0-98 0-99 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9551 0-9029 0-9562 0-9552 0-9522 0-9552			0.96	0.98
Isethionic 0.98 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.91 0.115 0.0345 0.0587 0.00346 0.0551 0.00346 0.0551 0.00346 0.0551 0.00346 0.0551 0.00346			0.98	0.99
Benzenesulphonic 0-99 0-99 Formic 0-0131 0-115 0-0587 0-058			0.98	0.99
Formic . 0-0131			0.99	
Propionic 0 00304 0 0551			0.0131	
Butyrie	Acetio		0.00345	0.0587
Butyric 0-00299 0-0551 Isobutyrio 0-00268 0-0518 Monochloracetic 0-0430 0-208 Dichloracetic 0-2304 0-480 Trichloracetic 0-682 0-826 Lactic 0-00901 0-0949 Hydroxyisobutyric 0-00921 0-0960 Trichlorolactic 0-069 0-263 Pyruvic 0-067 0-259 Oxalic 0-1746 0-430 Malonic 0-0287 0-169 Succinic 0-01181 0-1086 Tartaric 0-2296 0-1515 Racemic 0-0058	Propionie		0.00301	
Monochloracetic 0.0430 0.208 0.208 0.2304 0.480 0.2304 0.480 0.2304 0.480 0.682 0.682 0.682 0.682 0.682 0.682 0.682 0.682 0.682 0.682 0.682 0.0921 0.0949 0.0949 0.0960 0.263 0.067 0.259 0.263 0.0716 0.430 0.0716 0.430 0.0987 0.0987 0.0987 0.0987 0.0987 0.0918 0.0181 0.0181 0.0181 0.0181 0.1515 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.0287 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.1515 0.0296 0.029			0.00299	
Monochloracetic 0.0430 0.208 Dichloracetic 0.2304 0.480 0.248	Isobutyric		0.00268	0.0518
Trichloracetic . 0-682 0-826 Lactic . 0-00901 0-0949 Hydroxyisobntyrie 0-00921 0-0960 Trichlorolactic . 0-689 0-263 Pyruvio . 0-667 0-259 Oxalio . 0-1746 0-430 Malonic . 0-0287 0-169 Succinic . 0-00196 0-0704 Malio . 0-01181 0-1086 Tartaric . 0-296 0-1515 Racemic . 0-682 0-0949 0-0096 0-1515 Racemic . 0-682 0-826 0-00961 0-0096 0-1515			0.0430	0.208
Lactic 0 00901 0 0949 Hydroxyisobutyrie 0 00921 0 0960 Trichlorolactic 0 069 0 263 Pyruvic 0 067 0 259 Oxalic 0 1746 0 430 Malonic 0 00196 0 0704 Succinic 0 00196 0 0704 Malic 0 01181 0 186 Tartaric 22296 0 1515 Racemic 0 0296 0 1515	Dichloracetic .		0.2304	
Hydroxyisobntyrie 0.00921 0.0960 0.263	Trichloracetic .		0.682	
Trichlorolactic . 0.069 0.268 Pyruvic . 0.067 0.259 Oxalic . 0.1746 0.430 Malonic . 0.0287 0.169 Succinic . 0.00196 0.0704 Malic . 0.01181 0.1086 Tartaric . 0.2296 0.1515 Racemic . 0.0287 0.1515	Lactic		0.00901	
Trichlorolactic . 0.969 0.263 Pyruvio 0.067 0.259 Ovalio 0.1746 0430 Malonic 0.0287 0.169 Succinic 0.00196 0.0704 Malio 0.01181 0.1086 Tartario 0.2296 0.1515 Racemic 0.0296 0.1515	Hydroxvisobntyrie		0.00921	0.0960
Oxalio 0.1746 0.430 Malonic 0.0287 0.169 Succinic 0.00196 0.0704 Malic 0.01181 0.181 Tartario 0.2296 0.1515 Racemic 0.0296 0.1515	Trichlorolactio .		0.069	
Oxalio 0.1746 0.430 Malonic 0.0287 0.169 Succinic 0.00196 0.0704 Malio 0.01181 0.1086 Tartaric 0.2296 0.1515 Racemic 0.0296 0.1515	Pyruvio		0.067	0.259
Succinic 0.00196 0.0704 Malio 0.01181 0.1086 Tartario 0.2296 0.1515 Racemic 0.0296 0.1515			0.1746	
Malic 0-01181 0-1086 Tartario 2296 0-1515 Racemic 0-0296 0-1515	Malonic		0.0287	
Tartario	Succinic		0.00196	0.0704
Racemic 0 296 0 1515				
	Tartaric			
Citrio . 0.015 0.1279	Racemic		0.01596	0.1515
	Citrio	٠	0.012	0.1279

In the second column I have given the square

the determination of the affinities were free from the influence of all secondary reactions. Comparing these numbers with those already ob-tained, the same order of affinities appears, but in this case the values are more nearly equal. This agrees entirely with what had been expected; for it has been often emphasised that, owing to the presence of neutral salts, the statical methods make the strong acids appear too strong, and the weak acids appear too weak. The numbers found in the present case can therefore be justly considered as approaching nearer to the true coefficients of affinity than the previous values.

It is of special interest that a reaction such as the catalysis of methylacetate, which is only very remotely connected with the process of the formation of salts, is yet doubtlessly hrought about by that very property of acids which produces the latter class of reactions. This leads to the conclusion that the numerical values of all reactions exhibited by acids as such depend on that one property which till now has been somewhat vaguely termed the strength of the

Acid	I,	II.	111,
Hydrochloric	1.00	1.00	1.00
Hydrobromic	1.114	1.05	0.99
Nitrio	1.000	1.00	0.96
Chlorio	1.035	1.02	0.97
Sulphuric	0.536	0.732	0.739
Ethylsulphuric .	1.000	1.00	0.99
Isethionic	0.918	0.96	0.99
Ethylsulphonic .	0.912	0.95	0.99
Benzenesulphonic .	1.044	1.02	0.99
Formio	0.0153	0.124	0.115
Acetic	0.00100	0.0632	0.0587
Isobutyrio	0.00335	0.0579	0.0518
Monochloracetic .	0.0484	0.220	0.208
Dichloracetic	0.271	0.521	0.480
Trichloracetic .	0.754	0.868	0·826
Glycolio	0.01308	0.114	
Lactio	0.01066	0.103	0.0949
Methylglycolic .	0.01815	0.135	
Ethylglycolio	0.01372	0.117	
Methyllactio	0.01390	0.118	
Diglycolic	0.0267	0.163	
Pyruvio	0.0649	0.255	0.259
Glycerio	0.01715	0.131	
Oxyisobutyrio .	0.01062	0.103	0.0960
Oxalio	0.1857	0.430	0.430
Malonic	0.0308	0.175	0.169
Succinio	0.0545	0.0738	0.0701
Pyrotartaric	0.0107	0.103	
Malio	0.0127	0.113	0.109
Citrio	0.0173	0.131	0.128
Phosphoric	0.0021	0.249	
Arsenic	0.0481	0.219	

In order to verify this assumption Ostwald 4J. pr. [2] 29, 385, [1884]) investigated another process which is not connected with the formation of salty, via., the inversion of cane sugar. The reguion proceeds, as in the case of methylacocate, by the addition of water— $C_{12}H_{12}O_{11}+H_{2}O=2C_{6}H_{12}O_{6}$ —without an apparent intervention of an acid. Yet it takes place only in the prosence of acids. It is a catalytic reaction in the same sense as that already con Lnormal solutions; under III. the velocities of

sidered. As again only one substance, the sugary undergoes thange, the same equation holds good.

As an example has already been given which shows that the process is represented by equations (1) and (2), the velocities of inversion are now given directly (v. table in last column).

In the second column are given the square rocts of the velocities of inversion, in the third column the corresponding values for the catalytic change of methylacetate. The agreemont is evidently sufficiently great to prove the identity of the causes which produce the inversion of cane sugar and the decomposition of methyl acetate. This agreement also forms the complete experimental verification of the assumption that there exist constants of specific activity, which numerically determine all the manifestations of affinity exerted by acids as such. The numbers found for methylacetate and for cane sugar represent these constants with great oxactitude. The process of the inversion of cane sugar had been already used by Löwenthal and Lenssen (J. pr. 85, 321, 401) for the purpose of measuring constants of allinity. These chemists did not, however, deduce the constants of inversion from their experiments, although the theory of inversion had been established long before by Wilhelmy (P. 81, 413), nor did they give a proof of the fact that there are other chemical reactions which proceed according to a course analogous to that observed by them in the inversion of cane sugar.

Besides the chemical methods for the determination of the affinities of acids, there is yet another method which, by means of physical measurements, allows very accurate determination of these values to be made. It has been proved beyond doubt that the electrolytic conductivities of acids are closely connected with their chemical properties; so that this conductivity is proportional to the velocity of the reactions produced by the acids. As the electrolytic conductivity can easily be measured to a high degree of accuracy, we have here a method of much importance for the solution of the problems connected with affinity. The oxistence of this relation was first recognised and enunciated by W. Hittorf (W. 4, 391), who had, however, almost no measurements at his disposal. Arrhenius (Bijh. K. Svensk. Vet. Ak. Hand. 8, Nos. 13, 14 (1884)) developed a theory of the chemical changes among electrolytes, starting with the supposition that the power of conducting electrolysically and the power of participating in chemical reactions were idontical.

This theory leads to equations which agree with those of Guldberg and Waage. Finally W. Ostwald has considerably increased the somewhat scanty material available for comparisons between the power of inducing chemical reactions and electrolytic conductivity. Ho fully proved the proportionality between the velocity of the reactions induced by an acid and the electrolytic conductivity of the acid (J. pr. [2] 30, 93; ib. 30, 225 [1884]; ib. 31, 433; ib. 32, 300 [1885]). The following table shows this 300 [1885]). The following table shows this agreement. Under I. are given the electrolytic conductivities for normal solutions; under II: the velocities of inversion of oane sugar for semisolutions; all the numbers are referred to hydrochloric sold = 1. The numbers are referred to hydrochloric sold = 1.

7 14 4						
Acid	I,	ii.	III.			
Hydrochloric .	1.002	4.00	1.00			
Hydrobromic .	1.01	1.11	0.98			
Hydriodic .	1.01		0.96			
· Nitric	1.00	1.00	0.93			
Sulphuric	0.65	0.73	0.74			
Formic	0.0168	0.0153	0.0131			
Acctic	0.00424	0.004	0.00345			
Monochloracetic.	0.049	0.0484	0.0130			
Dichloracetic .	0.253	0.271	0.230			
Trichloracetic .	0.623	0.754	0.682			
Glycolic	0.0134	0.0131				
Methylglycolic .	0.0176	0.0183				
Ethylglycolic	0.0130	0.0137	_			
Diglycolic	0.0258	0.0267	_			
Propionic	0.00325		0.00304			
Lactic	0.0104	0.0107	0.0000			
Oxypropionic .	0.00006	0.0080				
Glyceric	0.0157	0.0177	_			
Pyrnvic	0.0560	0.0649	0.0670			
Butyric	0.00316	_	0.00800			
Isobutyrie	0.00311	0.00335	0.00268			
Oxyisobutyric .	0.0124	0.0106	0.0092			
Oxalio	0.197	0.186	0.176			
Malonic	0.0310	0.0308	0.0287			
Succinio	0.00581	.0.0055	0.0050			
Malic	0.0134	0.0127	0.0118			
Tartaric	0.0228	_	0.0230			
Racemic	0.0228		0.0230			
Pyrotartaric .	0.0108	0.0107				
Citric	0.0166	0.0173	0.0163			
Phosphoric	0.0727	0.0621				
Arsenic	0.0538	0.0481	_			

The agreement of the numbers in the three columns is evident, and proves the truth of the assertion made above.

In order to understand the rolation between conductivity and the power of taking part in chemical changes, we must go back to the theory of Clansius and Williamson. According to this theory the molecules of the electrolytic substances are continually interchanging their constituent atoms. These atomic exchanges generally take place to an equal amount in all directions; but when an electric current is passing they are so influenced that the electropositive or basic constituents go to the one side, and the electronegative or acid constituents to the other side, each constituent separating from the solution on one of the electrodes. This motion of the constituents occurs to a greater extent the greater the difference of potential between the electrodes, i.e. the greater the electromotive force. The change proceeds according to Faraday's law of electrolysis, which states that the quantity of electricity passed is proportional to the equivalents of the parts of the molecules separated out. Put into a slightly different form, this means that each electrified atom, or group of atoms, conveys the same quantity of electricity quite independently of its. nature.

Since the electric current only exerts a directive influence on the electrolyte, but does not relation is manifested:—
You. I.

itself induce the socion, the conducting power of substances depends antirally on the power of interchanging their ions. But on that same power depends also the velocity of the chemical changes produced by these substances; hence it follows that the velocities of the reactions must be proportional to the conductivities of the reacting substances. The experimental proof of this proportionality is in itself an important point in favour of the theory of Williamson and Clausius. (In Faraday's works we also find views which agree in the chief points with those explained above.) For the experimental details of the method the reader must be referred to the papers of Kohlrausch, Arrhenius, and Bouty. A short account of the conceptions and definitions used will, however, be given here, since the assumptions generally used in physics proper do not lend themselves well for our purpose.

Imagine a vessel having the form of a parallclopiped, the two parallel sides of which form the electrodes, and imagine the distance between the two to be equal to unit length. Into this vessel we imagine a quantity of the electrolyte to be placed, either by itself or in solution, such that its weightin grams is numerically equal to its molecular weight. Let us further suppose that the electromotive force between the two electrodes is unity; then the quantity of electricity passed through in unit time represents the molecular conductivity. Since equal quantities of electricity are conveyed by each electrolytic molecule, the total quantity of electricity passed is proportional to the number of double exchanges which takeplace in unit of time in one molecular weight of the substance

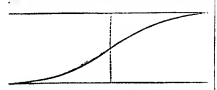
The electrolytic conductivity can be very early and accurately determined. The possibility of solving a great many problems connected with the values of affinities is thus presented. W. Ostwald has specially investigated the influence of dilution, and has established the laws which hold for it. The simplest relations are found for the strong monobasic acids, hydrochloric, hydrobromic, hydriodic, nitric, chloric, and perchloric. For normal solutions all these acids have nearly the same conductivity, and this increases by about 10 per cent. to 12 per cent, with increasing dilution, gradually approaching a maximum value, which in the units used by Ostwald was equal to 90. Sulphocyanic and bromic acids approximate to the acids named above.

The other monobasic acids, which are weaker, and which therefore have smaller conductivities than those already mentioned, exhibit a greater change in conductivity with igercasing dilution, the weaker they are, and they all do this according to the same law. This law states that the dilutions at which two acids have the same molecular conductivity always bear the same ratio to each other.

Thus, when measured in the units mentioned above, formic acid of the dilution 2—i.e. H₂CO₂=46 grams, in two litres of the solution—has the molecular conductivity 1%6; butyric acid reaches the value 181 only at 32 litres dilution. On further dilution the following relation is manifested:—

PORMIO APID		Вотчило Астр	
Bilation	Conductivity	Dilution	Conductivity
4	2.47	64	2.56
6	8.43	128	8.50
16	4.80	256	5.01
82	6.63	512	7.02
64	9.18	1024	9.74
128	12.6	2048	13.4
256	17.0	4096	18.0
512	22.4	8192	23.8
1024	29.0	16384	31.5

Butyrio acid and formio acid have always nearly the same molecular conductivity when the former is sixteen times as dilute as the latter. The same holds good for dilute solutions of all the monobasic acids. Plotting a curve, with the molecular conductivities as ordinates and the logarithms of the dilutions as abscisses, we find it to have the shape indicated in the annexed figure. In this, the



logarithms are not referred to the base 10, but to the base 2, since in Ostwald's experiments the dilutions increase as the powers of two. They are the exponential powers, p, of the dilution v, $v = 2^p$.

The curve appears to be symmetrical about two lines at right angles to each other, and has a point of contrary flexure when the conductivity reaches the value 45. The equation to the curve can be approximately expressed—using the given units—by the empirical formula

tan.
$$m = \left(\frac{v}{v_0}\right)^{-4518}$$

where m is the molecular conductivity; v_o the dilution (in litres per gram-equivalent) for which the conductivity is 45; and v any dilution for which the conductivity is to be calculated. The same curve holds for all acids if the abscissa v_o is chosen properly. The value v_o is characteristic for each acid. For the above-mentioned strong acids, it is found in high concentrations:—for iodic acid, at 2.8 litres approximately; for hypophosphorous acid, 8 litres; for dichloracetic acid, at 10 litres; for monochloracetic acid, between 400 and 500 litres; for formic acid, at about 1000; and for flutric feid, at about 1000; and for flutric feid, at about 1000 litres.

httes; for forme and, at about 1000; and for butyric acid, at about 1000 litres.

The influence of the dilution on the relative affinities varies welly for the different acids, as has already beginner. It seemed therefore very donbtful we there much importance could be attached to these values as natural constants on which the action of the acid as such depends. In the law of dilution, as enunciated above, there has been found the proof that we

are dealing with important and characteristic values, with values which do not alter with the nature of the reaction induced by the acid, and which are related to the dilution in a perfectly fixed manner. The general truth of this law for the case of chemical reactions has been separately proved by Ostwald (J. pr. [2] 31, 307). The values of affinities sought by men of science in the last century have thus been found by means of a method which had been evon then indicated by the famous opposer of the old theory of affinity.

It has already been mentioned that the above law of dilution holds primarily for monobasic acids. Polybasic acids behave differently according to their constitution. Some dibasio acids, such as phosphorous, selenious, &o., in which the second hydrogen atom is of the nature of a weak acid (this is shown by the alkalino reaction of their normal salts), behave on dilution at first exactly like monobasic soids, the conductivity being referred to molecular and not to equivalent weights. Hence electrolysis of these solutions takes place at first according to the type H | HR". It is only on reaching very great dilutions that the second hydrogen atom begins to participate in the reaction. Dibasic acids whose normal salts are neutral behave differently. It is true that they, too, conduct at first according to the type H | HR"; but the second hydrogen atom exerts its influence even in moderately dilute solutions. The conductivity increases much more rapidly than in the case of monobasic acids, and approaches a maximum which is double that observed for monobasic acids. Oxalic acid may be taken as a typical acid of this class. In the ease of very strong dibasic acids, such as sulphuric acid, it is the last part of the phenomenon just described which becomes prominent. Even for a concentrated solution the molecular conductivity exceeds the maximum of mone. basic acids, and rapidly approaches a value double that found with these acids. Henco conduction takes place from the beginning, for the most part, according to the type H2 | R".

ostwald has recently (J. pr. [2] 32, 300) examined the conductivity of a number of acids, and has shown that the relations already stated hold in all cases. He has also established many relations between the conductivities of acids and their chemical constitution. This investigation opens up the possibility of drawing many inferences concerning the action of chemical forces. The following tables exhibit an abstract of the measurements of the molecular conductivities of various acids for the dilutions of 4, 32, and 256 litres:—

	-		
Acid a	4 iitres	32 litres	256 litres
Hydrochloric HCl .	80.9	87.0	89.2
Hydrobromic HBr .	83.4	87.9	89.6
Hydriodic H1	83.2	89.6	89.7
Hydrofluorie HF	6.54	13.14	80.8
Hydrocyanic HCN .	0.077	0.108	
Sulphocyanic HSCN.	79.3	84.2 0	86.5
Sulphydric H.S		0.214	
FerrocyanioH, Fe(CN),		205.9	250.7

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ane some which do not contain oxygen show great differences: HCl. HBr, HI, form a group of strong soids, while HF is much weaker; HCN can searcely be called an acid; H₂S is slightly stronger; but if cyanogen combines with sulphur and hydrogen sulphocyanic acid is formed the strength of which approaches that of hydrochloric acid. Ferrocyanic acid is also a strong acid, though made up of the neutral iron cyanuce and the weakly acid hydrogen cyanide.

∆ cid	4 litres	32 litres	256 litres
Nitrie HNO ₃ Chlorie HClO ₃ Perchlorie HClO ₄ Bromio HBrO ₃ Iodie HIO ₃ Periodie H ₄ IO ₄	 80·4 80·2 82·2 	86·3 85·3 88·1 79·4 72·3 49·2	88·4 88·7 89·9 80·3 81·8 70·7

HNO₃, HClO₃, HClO₄ closely follow the halogen hydraoids; HBrO₃ is weaker; HIO₃ is still weaker; and H₄IO₆ shows this decrease in strength to a marked extent. A similar relation is shown by the acids of phosphorus.

Acid	4	32	256
	litres	litres	litres
Hypophosphorous H ₃ PO ₂ Phosphorous H ₃ PO ₃ Orthophosphoric H ₃ PO ₄	34.29	62·1 56·96 31·41	

Here too the acids become weaker as the amount of oxygen increases. The opposite relation is shown by the acids of sulphur and selenion.

Acid	•	4 litres	32 litres	256 litres
Sulphurous H.SO ₃ Sulphuric H ₂ SO ₄ Dithionic H ₂ S ₁ O ₃ Tetrathionic H ₂ S ₄ O ₃ Selenious H ₂ SeO ₃ Selonic H ₂ SeO ₄		19·19 96·4 — — 9·74 103·2	41.6 116.3 166.4 170.6 21.73 127.0	66.5 150.6 178.0 181.5 45.11 157.9

The strength increases with increase of oxygen as well as with increase of sulphur.

As rogards organic acids, the members of the acetic series are weak acids, and the strength decreases as we ascend in the homologous series.

Acid	4	32	256
	litres	litres	litres
Formic HCO.H Acctic CH.4CO.H . Propionic C.H.4CO.H . Butyric C.4H.5CO.H . Isobutyric C.4H.8.CO.H .	2·47	6·63	17:0
	0·755	2·12	5:64
	0·601	1·77	4:92
	0·604	1·81	5:01
	0·608	1·81	4:95
Valeric C,H,CO.H	0.615	1·87	5·16
Caproic C,H,1CO.H		1·70	4·78

Appreciable differences are shown in the first three members only.

When chlorine, bromine, iodine, or exanogen, is substituted for hydrogen, the acid a wacter increases.

Mold	litres	32 Hitres	256 litres
Monochioracetic CH, COCO, H Dichloracetic CHCl, CO, H Trichioracetic CCl, CO, H Monobromacetic CH, BrCO, H Cyanacetic CH, ONCO, H	6.98 84.6 65.9 10.6	17·3 50·8 75·0 16·1 25·8	87·8 76·2 79·1 36·8 57·8
*Bromopropionio CH_CHBrCO_H *Iodopropionio CH_ICH_CO_H	7·87 1·534	17·8 4·385	36·2

The chlorinated acids are seen to increase in strength as the amount of chlorine increases that even trichloracetic acid does not reach the value for hydrochloric acid. The substitution of bronnine, or cyanogen, for hydrogen acts in the same way as the substitution of chlorine; the action of cyanogen is much more marked than that of bromine. Introduction of the group OH for H in the fatty acids increases the strength of the acids, although not to so great an extent as is noticed in the preceding table.

Acid	4	82	255
	litres	litres	litæs
Glycolle CH_OHCO_H	2·08	5·77	15.09
	3·65	9·83	24.51
	1·94	5·49	14.42
#Uxypropionic CH_OHCH_CO_H	0:896	2·63	7·18
Trichtorolactic CCI_CHOHCO_H	11:0	27·7	54·8
Pyruvic CH_COCO_H	9:05	22·1	43·8
Glyceric CH_OHOHOHOO_H	2:46	6.87	17·9
=Oxybutyric C_H_CHOHCO_H	1:44	3.99	10·08
=OxybutyricCH_CHOHCH_CO_H	1:31	3.4	7·70
Oxylsobutyric (CH_)_COHCO_H . Methoxyacetic CH_OCH_CO_H . Rahoxyacetic CH_OC_H_CO_H .	1:98	5·07	12.81
	2:99	8·29	20.75
	2:40	6·94	17.98
Diglycolic O(CH_CO_H). Thiodiglycolic S(CH_CO_H).	5.00	13·78 11·73	83·58 28·22

The influence of the relative positions of the Oil groups appears to be of importance. In the cases of the isomeric lactic acids and the explutyric acids, that one is the strongest in which the OH is nearest the COOH group. Pyrnvic and glyceric acids seem to obey the same law, The following members of the exalic acid series have been investigated:

Acid	4 litres	32 litres	255 htres
Oxalic (CO,H), Malonic CH,(CO,H), Succhie C,H,(CO,H), Succhie C,H,(CO,H), Pyrotartaric C,H,CH,(CO,H), Pyrotartaric C,H,CH,(CO,H), Mathylmalonic CHCH,(CO,H), Kthylmalonic CHC,H,(CO,H), Sub-ric C,H,(CO,H), Sub-ric C,H,(CO,H), Malic C,H,(CH,(CO,H), Tartaric C,H,(CH,(CO,H), Lacenic C,H,(CH,(CO,H),	35-82 6-31 1-30 2-29 	61·4 16·8 3·72 12·91 5·74 12·14 15·18 ————————————————————————————————————	79-8 37-7 10-03 30-8 13-19 29-59 85-96 6-99 5-5-8 22-28 36-15 38-19
Succharic Calla(OH)a(COaH)a Mucic Calla(OH)a(COaH)a	Ξ	12.14	29.78 16.89

· By extrapolation.

As the distance between The two carboxyls increases the dibasic acids been acrapidly weaker; schacic acid is searcely stronge than the higher acids of the acotic series. Of the two isomerides, succinic acid and isosuccinic or no hydralonic acid, the latter has the carboxyls nearer together, and is therefore the stronger. Tartaric acid and acemio acid show no difference; hence the latter when in solution is not a compound of

right-handed and left-handed tartario sold, as is the case in the crystalline form, but it is rather a mixture of the two.

The derivatives of benzolo acid are of special interest owing to the conditions under which isomerism occurs in this series. The following have been investigated—

Acid	32 litres	256 litres
Benzolo C, H, CO, H Oxybenzolo [1:2] O, H, OHOO, H [1:3] [1:4] Nitrobenzolo [1:2] O, H, NO, CO, H [1:4] [1:4] [1:4] [1:4] Chlorobenzolo [1:2] C, H, CHOO, H	4:31 2:39	9·95 33·02 11·21 8·65 54·34 20·83 22·0* 32·54 15·13 12·7*
Bromobenzoio [1:2] C _e H _e BrCO _e H .	=	38·89 14·8

· By extrapolation.

The substituting radiclo always exerts most influence on the strength of the acids when it is in the ortho position. There is little difference between the meta and para positions. It is strange that para-exybenzoic acid should be weaker than benzoic acid itself, as in all other cases the introduction of OH increases the strength of the acid. This points to the conclusion that the affinity-forces are of the nature of vector quantities, that is, that they are directed forces the resultants of which cannot be put as simply equal to the sums of the components. The other acids of the aromatic (or benzenoid) group for which measurements have been made are as follows:—

		Acid		32 litres	256 litres
	midobenzene s	C _a H	NILSOLE	4.55	16.13
		C _a II.	$_{ m NH,SO,H}$	10.81	26:36 1:02
M	ononitropheno "	1 [1:2] C ₄ [1:3]	,,	0·14 0·177	0.26 0.41
Di	nitrophenol []	1:2,1:4]C,H	HO.(.ÖZ).	· =	10·95 79·7
A1	nisio C.H.OCE	"CO"II ["CO"H	: : :	3.51	5·00 9·1
Pi	enylgiyeolie (nenoxyacetie (CHOHOLL'S	Uall	9·02 12·57	22:75 29:55
Pl	thalic [1:2] C , [1:3] trophthalic C	, H (СО H)		38·62	35.23 20.0 66.57
N	trophthalic C	" 9H³(∇Ω°) ((29.22	\$57.50 25.00

The introduction of NH₂ into benzene sulphonio acid, which is nearly as strong as hydrochloric acid, is accompanied by the production of a much weaker acid. The meta-acid is weaker than the para-acid. The three nitrophenols show the gradation orthe, para, meta, in the same way as the nitrobenzoic acids. The phenols rapidly increase in strength with the number of NO₂ graps they contain. Another point of interest is the difference exhibited by the isomeride acid, phenylglycolic acid, and phenotyacetic acid. Orthophthalic acid greatly exceeds metaphthalic acid in strength; while of its two nitro derivatives, the a compound, in which the NO₂ group is adjacent to the calboxyl, proves itself superior to the B derivative in which there is a greater distance between the

NO, and the COOH groups. As regards the unsaturated soids, the following numbers show that, they are the stronger the less hydrogen they contain.

Acid "	4 litres	32 litres	256 litres
Acrylic C _a H _a CO _a H Crotonic C _a H _a CO _a H Funarlo Maleic C _a H _a CO _a H .	1·25 0·728 17·46	3·46 2·15 13·52 39·15	9-20 5-88 32-5 65-49
Citraconio CaHaCOaH	9·56 1·92	24.05 5.33 11.93	49.67 14.66 29.45 8.08
llydrociunammic C _a H _a C _b H _a CO _a H Cinnaminic C _a H _a C _b H _a CO _a H Phenylpropiolic C _a H _a C ₂ CO _a H Hydrosorbic C _a H _a CO _a H	1111	2·26 — 27·66 2·29	7·56 52·0 6·29
Sorbie C ₂ H,CO ₂ H. Bromecinnamnie C ₂ H ₂ C ₄ HBrCO ₂ H Bromecinnamnie	_	_	8·70 62·70
C ₆ H ₂ C ₃ HBrCO ₆ H Mecoute C ₂ HO ₂ Oll(CO ₂ H) ₂ . Quinte C ₆ H ₂ (Oll) ₂ CO ₂ H Camphoric C ₆ H ₆ C ₆ H ₆ CO ₂ H .	=	102·1 7·81	33·1 141·5 19·92 s·07

On comparing acrylic with propionic acid, crotonic with butyrio acid, fumaric and maleiq with succinic acid, and citraconic, itaconic, and mesaconic with pyrotartaric acid, it is evident that for each withdrawal of H., the acid becomes stronger. These relations become very conspicuous when we compare hydrocinnammic with cinnammic and phenylpropiolic acid, and hydrosorbic with sorbic acid. As regards the peculiarities of the dibasic unsaturated acids, they can only bo just mentioned. Meconie acid, which stands by itself, is conspicuous by its great strength. It is the strongest of all acids which consist of carbon, oxygen, and hydrogon only, and it approaches very near to sulphurio acid. There is evidently some connexion between this property and the small amount of hydrogen this acid contains.

The introduction of amidogen and similar radicles makes the acids weaker.

Acid	8 litres	32 litres	256 litres
Amidoacetic CH2NH2CO.H .	0.236	0.257	0.32
Ilippurio, CllaNII(C.H.CO)CO.H	-	8.68	17:88
Acetylamidoacetic CH_NH(CH_CO)CO_H Oxamic CONH_CO_H	21:07	6.88 35.88	17·76 82·26
Oxalurio CO(NHCONH ₄)CO ₂ H Parabanic (CONH) ₂ CO	43.35	57·03 48·23	74·28 53·98

The introduction of the NH₂ group into acetic acid is attended with a great reduction of the strength of the acid. This cid is considerably less weakened when one of the hydrogens in the NH₂ group is replaced by the negative radicle benzoyl or ucetyl. Oxalic acid is also weakened by introducing the NH₂ group. On the other hand the introduction of the urea residue (NHCONH₂) into oxalic acid only slightly decreases the conductivity of the oxalic acid Parabanic acid does not contain the carboxyl group, nor does it behave at all like an acid, since its conductivity increases but slightly with dilution.

The conditions of affinity among soids form the best-known part of the theory of affinity.

Our knowledge concerning the bases is much more scanty. From the fact that the relative affinities of agids are independent of the nature of the bases, we can infer that the relative affinities of the bases must also be independent of the nature of the acids (J. pr. [2] 16, 422). It is highly probable that the same laws hold for bases as for acids.

It must, however, be admitted that there is great lack of experimental data in this department. Some experiments of Menschutkin (C. R. 96, 256), who drew from them the conclusion that Bertholict's law of the influence of mass does not hold, prove only that under the conditions of the experiment the relative affinities of the bases compared-potash to aniline, to trimethylamine, and to ammonia, in alcoholic solutions - are very different. The same fact was proved by some thermo-chemical experiments of Berthelot.

Nor have many experiments been made by the kinetical method. Warder (B. 14, 1361) who first applied this method to bases, measured the velocity of decomposition of ethylic acetate. An investigation made by Reicher (Van't Hoff. . Dyn. chim. 107) in connexion with Warder's work, showed that the velocity of the reaction is nearly the same for potash, soda, and baryta, the olectrical conductivities being also nearly the same.

Ostwald (J. pr. [2] 33, 352) has recently investigated the electrical conductivities of some The alkalis KOII, NaOH, LiOII, arc strong bases; they have nearly the same conductivities. TIOH acts as an alkali. The conductivities of CaO,H., SrO,H., and BaO,H., referred to masses equivalent to NaOII, &c., are the same as those of NaOII, &c.; the molecular conductivities of CaO₂H₂, &c., are, however, double those of NaOH, &c. Ammonia is a weak baso; its conductivity is much influenced by dilution; the variations in the molecular conductivity follow the same law as was observed for acids. Substitution of H or H, iu NII, by CH3, C.H5, and other alkyl groups, increases the strength of the base; but N(CH,), and N(C,H,), are weaker bases than NII(CH_3)₂ and NII(C_2H_5)₂. These bases all follow the same law of dilution. The non-volatile ammonium bases, e.g. NMe,OH -and also the base (C2H3)3S(OII)2-exhibit conduotivities nearly the same as those of the alkalis; guanidine is a little weaker, but belongs more to the animonium bases than to the derivatives of ammonia.

In a memoir not yet published, Ostwald shows that those conclusions are confirmed by kinetical experiments on the saponification of cthylic

Besides the investigations which have led to numerical values for certain constants of affinity, there are numerous others from which such quantities cannot be deduced, because the resotions invostigated were too complex. To this class belong principally the works of Borthelot and P. de St. Gilles on etherification, and the investigations of Menschutkin (collected in A. Ch.

[6] 80, 81) on the same subject.
The importance of the first of these in chemical dynamics has already been emphasised The latter have brought out interesting conno

chemical constitution of the acids used. The values obtained do not, however, lend themselves to the determination of coefficients of affinity; and the investigations themselves cannot therefore be considered here at greater length. The work of Horstmann (B. 12, 64), and of Dixon (T. 1884. 617), on incomplete combustion can likewise only be mentiofied.

Little attention has as yet been directed to the investigation of the influence of temperature on the velocities of reactions, and on the constants of affinity. For the case of the inversion of cane sugar there are the investigations of Wilhelmy, Urech, and Spohr; for the velocity of etherification there are those of Berthelot and P. de St. Gilles, as well as those of Menschutkin. The whole question has been thoroughly investigated by Van't Hoff. By applying the dynamical theory of heat ho finds that the equation for the relation between the temperature and the velocity of the reaction, k, must be of the form

$$\frac{d \log k}{d\tau} = \frac{A}{\tau^2} + B$$

where r is the absolute temperature, and A and B are constants.

Van't Hoff has also shown that for several reactions the observed facts can be very well represented by such a formula. For details the book of Van't Hoff which has been mentioned must be consulted.

Berthelot and P.de St. Gilles have found that the chemical equilibrium of etherification is independent of the temperature within a wide range. Ostwald established the same generalisation for the relative affinities of various acids. Van't Hoff shows by the help of the dynamical theory of heat that this will occur when the reactions producing equilibrium do not give an appreciable thermal effect as their resultant. Moreover, when this is not the case, with a fall of temperature the equilibrium shifts in favour of that reaction which produces more heat than the reverse one (l. c. 167).

We have now arrived at the point where we must take up the problem concerning the rolation between affinity and production of heat. As soon as it had been recognised that the thermal action accompanying a chemical reaction was the measure of the chemical energy used up therein, an attempt was made to apply this to the question of affinity.

In 1854, J. Thomsen enunciated the following laws (P. 92, 34). The magnitude of the force evolved in the formation of a compound is equal to the quantity of heat produced. When a compound is decomposed by another body the reason for this is that the stronger admittes satisfy them. selves; hence decomposition must be accompanied by an ovolution of force. Since chemical force when liberated under ordinary conditions generally manifests itself as an evolution of heat, it follows that every simple or complex effect of a purely chemical nature is accompanied by production of heat.

This deduction, plausible though it seems at first sight, is not correct. Heat do's not measure forces but quantities of energy; hence the ohemical production of heat does not tell us any. thing concerning the intensity of ohemical forces: tions between the reactions observed and the it is only the product of their (mean) value (#: We imagine them to be forces of attraction) into the space passed over by the atoms that is a quantity of energy, and as such is measurable by thermal methods. Since we know nothing about the spaces passed over by the atoms, and, moreover, cannot assume that the spaces are the same for all compounds, no conclusion regarding the chemical forces can be accurately drawn from measurements of the quantities of heat

produced in ohemical reactions.

To point out the old mistake would have been uncalled for, were it not that Berthelot has of late years enunciated an analogous erroneous 'law,' which he has defended with great warmth. It is the more necessary to submit this theory to criticism, as, owing to the great prestige which the renovator of this old mistake enjoys - a prestige he owes to his excellent experimental researches-the theory is surrounded as it were by a halo which has deterred many from closely examining it. Bertholot formulates his law as follows: 'Tout changement chimique accompli sans l'intervention d'une energie etrangero tend vors la production du corps, ou du système de corps, qui dégage le plus de chaleur.' And further: 'Toute réaction chimique susceptible d'être accomplie sans le concours d'un travail preliminaire et en dehors de l'intervention d'une energie étrangère à celle des corps présents dans le système, se produit nécessairement, si elle dégage de la chaleur.'

By a vague connexion with general dynamics, Berthelot ealls this the principle of maximum work. He assumes it to follow as a natural consequence from the dynamical theory of heat. This view is erroneous. There is no snell thing as a law according to which a dynamical system is in equilibrium when the greatest possible quantity of its potential energy has been changed into actual energy; but this would be the dynamical analogue of the so-called chemical

There does, however, exist a law in the dynamical theory of heat according to which a system is in equilibrium when it has attained to the maximum entropy. This function, which was first introduced by Clansius, is, like 'the onergy of a system, entirely dependent on the condition of the system; it is defined by the equation,

 $dS = \frac{dQ}{T}$, where S stands for entropy, Q for a

quantity of heat given to the body, and T for tho absolute temperature. Horstmann (A. 170, 192), was the first to apply this law to chemical phenomena. The investigation has also been carried out very fully by Willard Gibbs. Unfor. tunately, the law is of very limited application. The integration can only be accomplished if the substances experimented with are perfect gases: Horstmann has shown that the law of entropy then leads to the same result as is attained by applying Guldberg and Waage's law of the influence of mass. This law has thus been supplied from the theoretical side with a valuable confirmation.

As far as we can tell, the law of entropy does not generally lead to reactions which are quite completed on one side, but rather to conditions of chemical equilibrium botween opposite processes. According to Van't Hoff (l.c. p. 153), these processes vary with the temperature, if they are accompanied by a positive or negative production of heat, and the law is that the equilibrium shifts the more in favour of the positive thermal production the lower is the temperature. It is, however, only at absolute zero that the reaction would take place in one direction only (if at this temperature chemical reactions are at all possible); and it is only for this limiting case that a law of maximum thermal offeet would hold good.

This is all that the law of the maximum thermal effect really contains; it is a limiting case from which the actual conditions differ the more the higher is the temperature. Since the temperature at which ordinary chemical reactions occur is not very high, the reactions accompanied by production of heat preponderate. This had been already noticed by Thomsen, and the approximation to truth contained in the law

we certainly owe to him.

What Berthelot has added refers to the cases of chomical equilibrium which have been established beyond doubt, and which, according to the principle of maximum work, ought not to occur; this law asserts that because one of two reciprocal roactions is attended with production of heat that one ought to take place exclusivoly. It is Berthelot's endeavour to reduce all reactions in which chemical equilibrium has been observed to cases of partial dissociation, wherein the masses of the reacting bodies do not act as wholes. To accomplish this, he is obliged to call reactions of a purely chomical nature, dissociations; for example, the decomposition of acid sodium sulphate in aqueous solution, a reaction brought about by the affinity between sulphuric acid and water. The whole explanation reselves itself into reasoning in a circle. It need searcely be said that an explana. tion of this kind cannot account for the laws by which the chemical equilibrium, the velocity of ehemical reactions, and the electrical conductivities of the reacting bodies, are connected.

There is no doubt that, with the possibility of a more general application of the laws of entropy to chemical reactions, thermochemical data will become important and fundamental means for the investigation of the relations of affinity. Moreover, there is little doubt that Bergmann's theory of affinity, revived in a thermochemical form, is not the solution of the problem, and that, in spite of its modern appearance, it can as little keep its ground against Berthollet's far-reaching views as it could

in its older form.

Of all the great old-standing problems of chomistry, that of chemical affinity has been least developed. The general relations and laws given in this article refer only to a limited number of substances, and to a limited number of reactions; many parts of the question have not yet been investigated at all. Great and important progress has, however, been achieved by Berthollet's enunciation, and Guldberg and Waage's rational formulations of the law of active masses. It must, however, be admitted that there are some reactions which seem to contradict this law, and which cannot be explained by it when taken in ets simple form. It is not necessary to reject the w on this account, as has been done by some. The actual conditions of each experiment we make

are so complex that we are not able to completely. apply the law of the influence of mass. must content ourselves with an approximation which does not always lie within the limits of experimental errors. The motions of the stars cannot yet be represented in strict accordance with the law of gravitation; yet the first approximation is sufficient to remove any doubts as to the validity of the law. The law of the influence of mass in its simple application is also only true to a sufficient approximation in those cases in which the effects considered are of great magnitude as compared with those neglectel. (In connexion with affinity v. PHYSICAL METHODS; section Optical.)

AGAR-AGAR or Bengal Isinglass.

A vegetable gum obtained in China from seaweeds: Eucheuma spinosum, sphærococcus lichenoides, spinosus, and tenar. Transparent colourless strips, almost completely soluble in water: forms a large quantity of thick, tasteless, and odourless jelly. Dilute H2SO, forms galactose, characterised by its conversion into galactonio acid by Br₂ and Ag₂O. This galactose is formed from a carbohydrate, C, H, O, present in the agar-agar (Bauer, J. pr. [2] 30, 367).

AGARICIC ACID C₁₆H₃₆O₅aq. [139°] (J.); [145·7°] (F.). S. 8 at 15°. Obtained, together with agaric resin, from the larch-fungus (Boletus Laricis) by extraction with dry ether (Fleury, C. R. 70, 53) or with 90 p.c. alcohol (Jahns, Ar. Ph. [3] 21, 221, 260).

Four-sided, silvery plates (from 90 p.c. alcohol at 50°) or prisms (from dry alcohol). V. sol. hot glacial HOAc or oil of turpentine, m. sol. chloroform or ether, v. sl. sol. benzene or cold water. Swells up and dissolves in beiling water but crystallises out again on cooling. Oxidised by HNO3 to butyric and succinic acids.

Salts.—Amorphous, insoluble pps. The neutral salts, M.C.,cH₂₅O₅ lose H₂O at 120° becoming M₂C₁₆H₂₆O₄.—NH₄HA".—Na₂A" (at 120°).—K₂A".—BaA".—Ag₂A": gelatinous pp. Hot alcoholic solutions give, with AgNO₃, a pp.

of Ag₂C₁₆H₂₆O₄ (Jahns).

AGARICIN. The fly-agaric (Agaricus albus) yields to alcohol a non-nitrogenous erystalline powdor having a sweet taste with bitter pungent after taste; slightly soluble in water, insoluble in ether; decomposed by boiling with dilute acids, or by contact with saliva, yielding a substance which exerts a slight reducing power on alkaline copper-solution (Schoonbroodt, J. 1864, 613). According to Jahns (J. 1883, 1400) it is identical with agaricic acid. AGARIC RESIN v. AGARICIC ACID.

Red, amorphous, solid; melts at 89.7°; dissolves in absolute alcohol, ether, wood-spirit, and ohloroform, but is insoluble in water, benzeue, and CS₂; slightly bitter; dissolves in alkalis. Na salt precipitated by alcohol in flocks changing in 24 hours into long needles. Forms precipitates, mostly crystalline with metallio salts (Fleury, C. R. 70, 53).

AGARICUS. A genus of Fungi.

Many fungi, especially the agaries, contain an amount of nitrogen exceeding that in peas and beans, varying from 3.19 p.c. to 7.26 p.o. (Schlossberger a. Döpping, A. 52, 106).

The solid tissue consists of cellulose. Agaric f

starch. Many agaries contain fumarie seid, sometimes associated with malic or citric acid. Agaricus bulbosus and A. integer yield stallisable hydrochlorides and platingcrystallisable chlorides of basic bodies (Thörner).

Aguricus ruber or sanguineus contains a colouring matter, ruberine, soluble in water and in alcohol. It is rose-red by trausmitted light, having two absorption bands in the green, but it exhibits strong blue fluoreseence. Dilute HCl extracts an alkaloid, agarythrine, from the fungus; this alkaloid is converted by oxidising agents into a red substance, possibly ruberine (T. L. Phipson, C. N. 46, 199).

Agaricus integer, contains an acid, with following properties: white needles [70°], very soluble in ether, benzene, CS., C11Cl., hot alcohol and acetic acid, insoluble in water, and cold alcohol and acetic acid.—A', Pb: insoluble white pp. [114°]. The alkaline salts are sparingly soluble in cold water, and the salts of the heavy metals, insoluble (Thörner, B. 12, 1635)

Agaricus atramentosus yields to boiling ether a dioxyquinone C₁₁H₆O₂(OH)₂. Dark brown metallic-shining lamine, dissolving with yellow colour in alkalis, insoluble in water, ether, light petroleum, benzene, chloroform and CS2. Sublimes with great difficulty in yellow microscopia tablets. It is reduced by boiling its alcoholic solution with zinc-dust, the resulting colourless liquid becoming yellow green again on exposure to the air. The ammonium salt is a green crystalline powder, dissolving readily in water with violet colour, nearly insoluble in boiling absolute alcohol.—Ba salt: dingy flesh coloured erystallino precipitato (Thörner, B. 11, 533). The diacetyl derivative $C_{13}H_{12}O_4 = C_{11}H_{\bullet}O_2(OAc)_{\bullet}$ forms small reddish yollow tablets.

Boletus Laricis contains besides agaricio acid and (25 p.c. of) agaric resin also 3 to 5 p.c. of a neutral body, which crystallises in needles, [272°], and may be sublimed (E. Jahns, J. 1883, 1400). H. W.

AGAVE. Well preserved juice of Agave americana, sp. gr. 1.046 at 15° was found by J. Boussingault (A. Ch. [4] 11, 447) to contain in 1000 parts: 26:45 levulose, 61:71 saccharose, 3.53 malic acid, 5.45 gum, 10.13 albumin, 0.06 ammonia, 6.21, inorganic salts, and 886.46 water.

AGE or AXIN. The fat of Coccus Axin growing in Mexico, consists of the glycerides of lauric

and axinic acids (Hoppe, J. 1860, 324). H.W. AGGREGATION, STATES OF.—In this articlo the differences between the properties of bodies in the solid, the liquid, and the gaseous, condition, are looked upon as due to differcuces in the state of aggregation of those small particles, of which, according to the molecular theory of the constitution of matter, all bodies are composed. According to this theory, our power of subdividing matter cannot be carried beyond a certain limit, whatever means-chemical, physical, or mechanical-we employ. In other words, the theory asserts that the largest quantity of a body which we cannot subdivida by any means in our power is of finito size; it is called the atom of the substance of which the body is composed. Each elementary body has its peculiar atom, and the union of atoms of sontain mannite and fermentable sugar, but no different kinds forms the smallest quantity

which can exist of a compound substance; this, however, cannot, in accordance with the definition, be called an atom, since, by the nature, of the case, it can be divided by chemical, and often even by physical, means. Though matter can be divided down into atoms by chemical means, yet we have reason to believe that when only physical processes are going on the sub-division of matter is not in general carried so far; and that just as in an army, though the unit is the individual soldier, yet for military purposes the soldiers forming a regiment always act together, so in matter, groups of atoms, called molecules, remain together for a considerable time. The molecule, however, is a very much less definite thing than the atom, and it must not be assumed without proof in each case that the term has always a definite meaning, or that there may not in the same body be molecules consisting of vory different numbers of atoms. There is strong evidence, too. that, in some cases at any rate, the molecule does not always consist of tho same atoms; the molecule after a time seems to break up and the constituent atoms find fresh partners. In some cases, however, such as those of the permanent gases, we have reason to believe that the number of molecules which consist of the same number-say n-of atoms, is enormously greater than the number of those consisting of any other number of atoms. If. however, we raise the temperature, then, in the case of some gases at any rate, dissociation sets in; that is, there are now a considerable number of molecules in which the number of atoms is lesa than n; this is shown by the abnormally smell densities of such gases at high temperatures. On the other hand, the donsity of a vapour near its point of condensation is often abnormally great, es in the case of acetic, formic, and monochloracetio, acid; a part at any rate of this increase in donsity would seem to be due to the formation of molecules consisting of a greater number of atoms than those formed when the temperature was raised far above that of the point of condensation.

According to the molecular theory of matter, the difference between the molecular constitution of bodies in the solid, liquid, and gascous, state is that in the solid state the molecules oscillate about a position of equilibrium and never get far from their original position in the body; in the liquid state the molecules are supposed not to oscillate about positions of equilibrium, but to be comparatively free to move in any direction; they cannot, however, move far without coming under the influence of other molecules, so that their courses are constantly being changed and do not bear any approximation to straight lines; in the gaseous state the molecules are so far apart that for the greater part of the time they are describing straight lines, the time during which they are under the influence of other molecules being an exceedingly

small frection of the whole time.

We must be careful to remember that there is no evidence that the molecules in the liquid or solid state consist of the same number of atoms as those of the same substance in the tion is quite appreciable in the case of all but gaseous state; but that on the contrary it seems, the most permanent gases. Maxwell investimest probable that in the solid and liquid states gated the distribution of velocity among the the molecules are systems whose comploxity is molecules of a gas, and showed that when the

not only very different from the molecules in the gaseous state but that these molecular aggregations vary very much in complexity among themsalves. These molocular aggregations are probably not permanent but are continually breaking up and their constituents changing partners; this breaking up and re-formetion of the molecular aggregations would produce the same effect as the collisions between the molecules of a gas, that is, it would tend to equalise the distribution of momentum and energy, so that it would make the substance possess viscosity, and he able to conduct heat. In fact, the collision between two molecules of a gas is the formation and breaking up of a molecular aggregation, and the difference between this case and that of a solid or a liquid is that the ratio of the time the molecular aggregation lasts to the time which clapses between the formation of suocessive aggregations is much smaller in the case of the gas than in that of the liquid or solid. The simplest state of aggregation we can imagine is one where the molecule and the atom are identical, that is, where the molecule consists of only one atom; this easo is realised by a monatomic gas such as mercury, and possibly hy all gases when the temperature is sufficiently high. The properties of matter in this state have not been investigated with special regard to the differences between this and more complex states of aggregation; Schuster (Pr. 1885), however, has shown that the phenomena of the cleetric discharge through mercury vapour are quite different from those occurring in a ges whose molecules are polyatomic.

In the case of most elementary gases the molecules consist generally of two atoms, and this case has received by far the largest amount of attention both from the experimental and the theoretical point of view. The most important results of the kinetic theory of gases from the chemical point of view are: -first, Avogadro's law, which states that in equal volumes of all gases at the same temperature and pressure there are the same numbers of molecules. From this it follows at once that, as long as all the molecules consist of the same number of atoms, the ratio of the molecular weights of two gases is tho same as the ratio of their densities. It must, however, be clearly understood that this result is only true for perfect gases, that is, for gases in which the pressure is produced entirely by the striking of the molecules against the sides of the vessel containing the gas, and not at all by the force between the molecules. If a gas obeys Boyle's law it is a perfect gas for this purpose, and wo may apply Avogadro's law to it; this law is not, however, applicable when Boyle's law does not hold. If the departure from tho law be slight, and if δp be the deviation of pressure from that given by Boyle's law, then the number of molecules in unit volume will equal the number in the same volume of a perfect gas at the same temperature and pressure multiplied

by $\left\{1-\frac{\delta p}{p}\right\}$, where p is the pressure. This correc-

gas was in a steady state the molecules could not all be moving with the same velocity; he gave (P.M. [4] 19, 22) a formula which tells how many molecules there are whose velocities are between any assigned limits. We shall here, however, only give a few numbers calculated from that formula. To take the case of oxygen at 0°C., about \(\frac{1}{3} \) the molecules are moving with velocities of between 300 and 600 metres per second, about $\frac{1}{4}$ between 300 and 100, only about $\frac{1}{100}$ with velocities less than 100 metres per second, and not $\frac{1}{1000}$ part with velocities greater than 1,200 metres per second. velocities with which the molecules of the same gas are moving at different temperatures are proportional to the square roots of the absolute temperatures; thus the distribution of velocity among the molecules of oxygen at 273°C. would be got by multiplying by $\sqrt{2}$ all the velocities at 0°C. The velocities with which the molecules of different gases are moving at the same temperature are inversely proportional to the square roots of their molecular weights; thus, for example, the velocities of the hydrogen molecules are on a scale four times as great as that of the oxygen molecules.

We can estimate by the methods of the kinetic theory of gases (see Meyer, Die Kinetische Theorie der Gase) the number of molecules in a cubic centimetro of the gas, and the diameter of the molecule, if the molecule is looked on as a hard elastic sphere; or if the moleenle be considered as a system, we can estimate the distance between two molecules when their paths become appreciably curved. We find as the result of such calculations that there are about 21 trillion molecules in a cubic centimetre of gas under the pressure of 760 mm. of mercury at 0°C; so that the mean distance between the molecules is between 3 and 4 millionths of a millimetre, or about 3.5 x 10-7 centimetres; tho diameter of the molecule is probably between 1×10^{-7} centimetres and 3×10^{-9} centimetres, or between 1 and 110 of the mean distance between the molecules. Another quantity which it is important to know is the mean distance through which the molecule passes between two collisions; this is called the mean free path of the molecule, and it is inversely proportional to the density. For hydrogen at the pressure of 760 mm. of mercury the mean free path is about 1.8 × 10-5 centimetres; at the pressure of 1 mm. the free path is about in of a millimetre; and at a pressure of a millionth of an atmosphere about 18 centimetres. When the free path is comparable with the dimensions of the vessel in which the gas inclosed, the gas can exhibit phenomena of a different character from those shown when the free path is indefinitely small compared with the dimensions of the vessel. The radiometer exhibits effects of this kind, and Crookes has called a gas rarefied so much as to show rotation in a radiometer, matter in the fourth or ultra gaseous state. But this is using the word state in a different sense from that in which it is used in the phrases solid, liquid, and gaseous, states; for these states do not depend upon anything but the matter itsel while the ultra-gaseous state depends upd i the ratio of the free path to the other lengths

involved; if we increased all the lengths proportionately to the ratefaction, the gas would not show any of those properties which characterise the so-called ukra-gaseons state, while, on the other hand, if we experimented with small enough instruments we could get all the ultragascous effects manifested by a gas at the atmo-

spherie pressure.

The distribution of energy among the molecules is of much chemical interest. It seems, however, that in one respect the results of theory have been misinterpreted; it has been said that because iodine, for example, is dissociated at a temperature a little over 600°, and since in iodine at any temperature there are some molecules possessing the same amount of energy as those which are split up at 600°, that therefore these molecules ought to be split up, and if any substance were present capable of combining with free iodine the whole of the iodine would ultimately combine with this substance. Now although there are apparently no experiments which may be called secular to say whether or not this would ultimately happen. yet it is certain that it does not happen so quickly as theory would indicate, if every molecule which possessed the same kinetic energy as that possessed by the average molecule at 600° were straightway dissociated and entered into combination with the other substances present; there seems, however, to be no reason why this should be the case, for though one molecule at 0° may have the same energy as one at 600° yet dissociation must depend upon the surround. ing molecules as well as upon the molecule itself. Now the molecule at 600°, though it possesses at any instant the same energy as one at 0°, is yet surrounded by molecules which are moving very much faster than itself, and whose energy is much more nearly equal to its own, so that it is not so likely to lose its energy by collision with other molecules as the molecule at 0° which is surrounded by molecules with much less energy than itself. For this reason the tendency to dissociate will be very much greater at 600° than at 0°, and a molecule at the former temperature may dissociate while the latter may lose its energy before this can happen.

The distribution of energy affects the specific heat very much; so that if we know the value of the specific heat we can tell a good deal about the energy of the molecule, as the following theoretical investigation will show. Let us begin with the case of a gas the molecule of which is of any degree of complexity, measured by the number of degrees of freedom, p. There is a theorem due to Boltzmann which etates that the mean energy corresponding to each degree of freedom is the same, so that the mean

total energy of the molecule is $\frac{p}{3}$ times the mean

energy due to the translatory motion of the centre of gravity. Though there is very strong evidence against the truth of the theorem in this form, and the mathematical proof of it is nnsatisfactory, yet a very special case of it is probably true, viz. that if we have a molecule consisting of n atoms approximately symmetrically arranged (that is, if the distance between a particular pair of atoms is not always very much less than the distances between the other pairs), then the ratio of the mean total kinetic energy of the moleculs to the energy due to the translatory motion of its centre of gravity is proportional to n, the number of atoms in the molecule.

Let the ratio of the total kinetic energy to the translatory energy of the centre of gravity, which by the kinetic theory of gases is measured by θ the absolute temperature, be βn . Then the total kinetic energy in the gas $-\Sigma \beta n\theta = \beta n'\theta$, when n' is the number of atoms in the gas. If all the atoms be of the same mass, m, and the quantity of gas be the unit of mass, then n'm = 1, so that the total kinetic energy in the gas $\frac{\beta \theta}{m}$, which, if β be the same for all elementary

gasss, is inversely proportional to the atomic weight of the gas. If the gas had been a compound such that each molecule consisted of a atoms of mass m, b of mass m', c of mass m'', and so on, then for unit mass of the gas,

$$\frac{N'(ma + m'b + m''c)}{(a + b + c)} = 1,$$
energy in unit mass

so that the energy in unit mass

$$= n'\beta\theta$$

$$= \frac{\beta\theta(a+b+c)}{ma+m'b+m''c}.$$

Now a+b+o is the number of atoms in the molecule, and ma+m'b+m''c is the mass of a molecule, so that the energy of unit mass

 $=\frac{\beta\theta(\text{number of atoms in the molecule})}{\text{mass of the molecule.}}$ Let us first suppose that all the energy in the gas is kinetio, then the energy in unit mass of the gas at temperature θ is $\frac{\beta\theta}{m_1}$ so that the specific heat is

 $\frac{\beta\theta}{m_i}$ or the product of specific heat into the mass of an atom, which is called the atomio heat, is equal to β , and, as experiment shows, does not vary much from gas to gas. For a compound, we see from the expression given above for the energy, that the product of the specific heat into the mass of a molecule equals β (number of atoms in the molecule), so that for all perfect gases, simple or compound, the product of the specific heat into the mass of the molecule = β (number of atoms in the molecule). We may remark that with our assumptions the ratio of the specific heat at constant pressurs to that at constant volume

=1+ $\frac{2}{8\beta}$ $\left(1+\frac{\delta p}{p}\right)$ number of atoms in the molecule,

when δp is the deviation of the pressure from that given by Boyle's law. The experimental results show that for most perfect gases, simple or compound, the molecular heats are constant, showing that β is constant, for such gases, or that the whole kinetic energy is proportional to the product of the number of atoms in the molecule and the energy due to the translatory motion of the centre of gravity. There are, however, some simple gases, such as chlorine and bromine vapour, whose atomic heats are much higher than the value given by the above rule. These gases are, however, easily liquefied, and so when heat is applied, work is done in altering the molecular state as well as in raising the temperature; this will produce an effect

equivalent to increasing B, and will therefors explain the large value of the atomic heat. We should expect a large value for this quantity too if the gas were dissociating. There are some compound gases, on the other hand, such as ammonia, ethylens, and marsh gas, whose molecular heats are too small to agree with the above rule, if we suppose that the number of different systems in the molecule is the same as the number of atoms indicated by the chemical formula of the gas. If, however, two or more atoms always remain close together, they will for our purpose count as one atom, as it is only when the molecules are approximately symmstrically arranged that we can assume that the total energy is proportional to the number of atoms. The total energy is proportional to the number of distinct systems; and if a group of atoms always remain close together they only count as one system, however many atoms there may be. If, for example, the atoms in a radicls always remain together, the radicle, for this purpose and in the formula for the ratio of the specific heats, will only count as one atom. Ws may therefore regard those compounds which have too small atomic heats, as consisting of but few separate systems, though there may be a great number of atoms in the molecule.

The determinations by Dulong and Petit and others of the specific heats of elementary bodies in the solid state show that for these bodies the atomic heat is approximately constant, while Kopp's experiments on the specific heats of compound solid bodies show that for many such solids the product of the mol. w. and the specific heat is proportional to the number of atoms in the molecule, just as for gases. The expression for the kinetic energy of unit mass of a solid will probably be of the same form as that which we found for a gas; for this only depends upon the assumptions that the absolute temperature is proportional to the mean energy due to the translatory motion of the centre of gravity of the molecules, and that the ratio of the mean total kinetic energy of the molecule to the mean energy due to the translatory motion of the centre of gravity is proportional to the number of atoms in the molecule. These assumptions will probably hold for the solid and liquid as well as for the gaseous state. We must re-member that when heat is applied to a solid or liquid, work is done in altering the molecular configuration as well as in increasing the kinetio energy of the molecules. All solids and liquids appear to be able to get into a condition in which the specific heat does not alter with the temperature, and it is in this condition that the atomic heat is constant. Now if the specific heat is independent of the temperature, the work spent in altering the molecular configura-tion must bear a constant ratio to the work spent in increasing the kinetic energy; and if the atomic heat is constant this ratio must be the same for all substances; so that Dulong and Potit's experiments show that when heat is applied to a solid or liquid it is divided between the energy of molecular configuration and the mean kinetic energy, in the same proportion for all substances; and since for many substances, such as iodine, bromine, msrcury, &c., the specific heat in the solid state is twice that in

the gaseous, it is equally divided between these two forms of energy, a result which purely dynamical considerations would also lead ne to regard as the most probable one. The specific heats of liquide seem to be more irregular than those of either solids or gases, but the bodies for which this is the case are those whose melting and boiling points are comparatively close togother, and we may suppose that the nature of the molecular configuration alters with each change in temperature, and this makes the specifio heat abnormally large. The specific heats of those substances which exist in the fluid etate through wide ranges of temperature seem to ho the same in the solid and fluid states.

Whon the specific heat of a solid compound is much smaller than the number of atoms in it would lead us to expect, we may, just as in the case of a gas, conclude that two or more atoms always remain close together in the molecule. It is important to notice, however, that tho specific heat cannot give us any information about what we may call the molecular aggregation of the solid or liquid, that is it affords no information as to whether the molecules are isolated or form groups, for if we suppose the molecules to unite and form more complex ones the atomic heat would remain the same ac long as the energy was equally divided among the atoms or radicles forming the molecules.

CHANGE OF STATE.—GASEOUS TO LIQUID.

By the application of great pressure accounpanied when necessary by intense cold, all gases have been liquefied, and during this process they pass through all intermodiate states, so that at some stage of the process it is impossible to tell whether the substance is a gas or a liquid. It is found that there is for each gas a temperature above which it cannot be liquefied by the application of the most intense pressure, so that at a temperature higher than this the substance can only exist as a gas. This temperature is called the critical temperature, and Andrews has proposed to call a substance at a temperaturo higher than its critical temperature a gas, and one which though in a gascous condition is yet at a temperature lower than the critical tempsratnre, a vapour. Van der Waals (Continuität des Gasformigen und flüssigen Zustandes, 87), and Clausius (W. 9, 1880), have shown how to calculate the critical temperature from the difference between the pressure of the gas in any state and that given by Boyle's law.

We shall here confine ourselves to showing, by general reasoning, that a critical temperature must exist. When a body is in the liquid state the ratio of the work required to separate the particles to an infinite distance to the kinetic energy of the molecules must exceed a certain limit, for the substance will behave as a liquid or a gas according as the forcee between the molecules are or are not able to change their kiuetic energy appreciably in the interval from one collieion to another. The molocules will change their kinetic energy appreciably if the ratio of the alteration in the mutual potential energy of the molecules to their initial kinetic energy is finite, but for this to be the case the ratio of the work from the liquid into the gaseous state. In the

distance must be finite, so for a body to be in the liquid condition this ratio must exceed a certain quantity, say, R. Let v be the work required to separate one of the molecules from the remainder, and I the kinetic energy of the translatory motion which is proportional to the absolute temperature; then the substance will behave like a liquid if $\frac{\mathbf{v}}{\mathbf{r}}$ be greater than \mathbf{r} , but like a

gas if $\frac{\mathbf{v}}{\mathbf{r}}$ bo less than this quantity. Now \mathbf{v} cannot be greater than the work, v', required to separate the molecules when they are quite close together, so that when $T > \frac{V'}{R}$ the substance

will always behave like a gas. Now T is proportional to the absolute temperature, so that when the absolute temperature exceeds a certain value the substance will always beliave like a gas, that is, it cannot be liquefied. This shows that there must be a 'critical temperature,' and it also shows that the critical temperature is proportional to the work required to scharate the molecules; a measure of this will be the amount of heat required to convert the substance from a liquid into a gaseous etate under infinite pressuro. We can take as a practical measure the latent heat of the substance. The mean kinetic energy of the translatory motion equals the absolute temperature, so that if h be the latent heat, θ the critical temperature, nv' will be proportional to h, where n ie the number of molecules in unit mass, and T is proportional to 0, so that since $\frac{\mathbf{v}'}{\mathbf{r}}$ is constant, we should expect

to find that the critical temperature multiplied by the number of molecules in unit mase-or, what is proportional to it, the reciprocal of the molecular weight - ought to be related to the latent heat so that when one is great the other is great also. The following table will show that this condition is approximately fulfilled:-

Substance	Absolute- criti- cal tem- pera- ture	Critical tempera- ture divided by mol. w.	Latent heat
Alcohol C.H.O Acetone C.H.O Acetone C.H.O Carbon disulphide CS2 Benzene C.H. Methyl acetate C.H.O2 Ethyl formato C.H.O2 Sulphurous oxide SO2 Ether C.H.O Ethyl acetate C.H.O	510 505 546 558 503 503 429 468	11·1 8·7 7·3 7·17 6·8 6·8 6·7	209 140 105 109 110 105 94
Chiloroform CHCl, Carbon tetrachlorideCcl,	513 533 557	5·83 4·51 3·6	105 67 52

PASSAGE FROM THE LIQUIN TO THE SASEOUS STATE.

Though it requires the application of pressure and cold to make a substance pass from the gaseous to the liquid state, yet the substance will always to a limited extent pass of itself required to separate the molecules to an infinite | space over a liquid in the equilibrium condition

there is always a quantity of the vapour of the liquid, the quantity of vapour in unit volume depending only on the nature of the liquid and the temperature; in other words, the vapour exerts a definite pressure called the vapour pressure (often erroneously the vapour tension). If we have a quantity of liquid in a vessel furnished with a piston the liquid will evaporate until there is a certain quantity of vapour in each unit of volume above the liquid; if we depress the piston so that this volume diminishes by w then a quantity of vapour equal to that in volume v will condense; if the piston be raised again the vapour will be re-formed. In this way wo can have a continual transference from the gascous into the liquid state and back again. In this process, however, we have only matter in these two states and have no continuity of state from the gascous to the liquid as we had in the process by which the permanent gases are liquefied. The vapour pressures of different liquids vary enormously; thus for sulphurio acid the vapour pressure is so small as to be almost inappreciable; for sulphuric acid, mixed with its own volume of water, it is about one-eighth of a mm. at 15°C.; for water at the same temperature it is about 12.6 mm.; for alcohol, 32 mm. The vapour pressuro always increases as the temperature rises, but until the temperature reaches a certain value depending on the pressure, the liquid which evaporates is always that on the surface, and none of the liquid in the interior passes into the gareous condition. When, however, the vapour pressure becomes greater than tho pressure on the surface of the liquid, the bubbles of vapour which form on the sides will be at a pressure equal to or greater than the pressure in the surrounding fluid, and so will expand and be able to reach the top without con-When this takes place, i.c. when densing. portions of the liquid not on the surface are converted into gas, the liquid is said to boil. The temperature of the boiling-point will increase with the pressure; it cannot, however, even by the application of an infinite pressure, be raised above the critical temperature of the substance. Bodies which have small vapour pressures at ordinary temperatures have high boiling-points; but it does not follow that because the vapour pressure of one substance is at some temperature greater than that of another its boilingpoint will be lower; for example, at 15°C. the vapour pressure of carbon tetrachloride is greater than that of methyl alcohol, though its boiling point is higher. According to the molecular theory, some of the molecules manage to escape from the liquid, we may suppose because they are moving so fast as to be able to escape from the attraction of the other molecules; on the other hand, some of the molecules of the vapour strike the surface and to caught by the moleonles of liquid. When things are in a state of equilibrium as many molecules escape from the liquid as are caught by it.

Although the vapour densities of many substances have been determined, few experiments seem to have been made on the rate of evaporation and condensation. The knowledge of these rates would very much increase our knowledge of the constitution of fluids.

The forces between the molecules in the

liquid state must be sensible, otherwise we should not be able to spend work upon a liquid without increasing the kinetic energy, as we do when we convert water into steam at the same temperature. The latent heat may be taken as a measure of the potential energy lost by the transition from the gaseous to the liquid state. In a fluid the potential energy of the molecular configuration seems to depend only on the mean distance between the molecules; for the fluid resists anything tending to diminish its volume, but does not resist anything tending to change its shape. When the fluid is in such a condition that its specific heat is independent of the temperature then any increase in the kinetic energy must be accompanied by a proportionate inorcase in the potential energy of the molecular configuration. Now, if the forces between the molecules are large it will require a smaller increase in the distance between them to increase the potential energy by a given amount than if they were smaller; so that for a given increase in the kinetic energy, that is, a given rise in temperature, the increase in volume will be less when the forces between the molecules are great than when they are small, so that the coefficient of expansion will be small when the forces between the molecules are great, but when the forces between the molecules are great the fluid is incompressible and the product of the mol. w. and latent heat is large; so that we should expect a small coefficient of expansion, incompressibility, and large latent heat for equal volumes to go togethor, and we find by the following tables of these quantities that this scems to be the

	Product of latent heat and mol.w.	CTCHE OI	Com- pressi- bility
Water, H ₂ O. Benzene, C.H ₄ Acetone, C.H ₄ Chioroform, CHCl. Curbon tetrachloride, CCl. Ether (C.H ₂) ₂ O. Curbon bisulphide, CS ₂	11,088 8,502 8,120 7,906 7,904 6,956 6,840	*000065 *00138 *00172 *00140 *00140 *0021 *00146	4·61 10·8 6·26

A peculiarity of water is that it is denser at 4°C, under atmospheric pressure than at any other temperature under the same pressure; we may perhaps suppose that this is due to something of the following kind. We know that when water freezes it expands and erystallises in the hexagonal system; now we may suppose that, before the water solidifies, molecular aggregations are formed which possess the same property as is possessed by the ice crystals, viz. that when the molecules are arranged in this way they occupy a greater volume than when arranged uniformly-the formation of these aggregations would tend to increase the volume and might be sufficient to more than counterbalanco the diminution due to the nearer approach of those particles which do not form these aggregations.

CHANGE OF STATE FROM SOLID TO LIQUID.

When the temperature of a solide is raised sufficiently high it begins to melt. There are two kinds of melting; in the one, as in the case of ice, it the heat is applied slowly the tempera-

ture remains constant until all the substance has passed from the solid into the liquid state In this case there is a definite melting-point. In the other case, of which an example is the melting of sealing-wax, the substance first begins to soften, then as more heat is applied it gets softer and softer, its temperature, however, increasing until when a certain temperature is reached the substance is liquid; in this case there is no definite melting-point, as the process is sprsad over a considerable range of temperaturs. This would seem to imply that the substancss which melt in the second way are not perfectly homogeneous in structure, but contain molscular aggregations of various degrees of complexity, which get gradually split up as the temperature rises; while those substances which melt liks ice have a more uniform constitution, so that any change of state takes place simul-taneously through the molecules. This would obviously tend to make the transition more definite. This view is in accordance with the fact that crystalline bodies, which are generally regarded as being more uniform in structuro than non-crystalline, all melt in the same way as ice. Scaling-wax in the state of transi-tion is what is called a viscous body; so that on this view a viscous substance is regarded as a mixture of molecules some of which are in the same state as they are when the substanco is liquid, and some are in the same state as they are when the substanco is solid. A dynamical illustration will enable us to see how such a body might behave like a rigid body under the action of rapidly changing forces, and like a fluid under constant or slowly varying forces. Suppose we have a series of heavy spheres connected by strong springs placed upon a horizontal table, and that one end of this row of spheres is fastened to a peg which cannot sustain a tension greater than T without breaking; then if a steady pull, r, be exerted at the other end of the row of spheres the string will break, but if the tension at the end, instead of being steady, be periodic, and if the period of vibration be greater than the natural period of vibration of the spheres and springs, then if the number of spheres be great enough the string will not break, though a tension enormously greater than r is acting at the other end. This is quite a parallel caso to that of the viscous fluid; the springs and spheres correspond to those molecules which are in the same condition as when the substance is solid, the string to those in the fluid condition.

The change of state from solid to liquid seems to be always accompanied by a change in volumo, and when this is so the melting-point -as J. Thomson proved-must be altered by the application of pressure. Thomson showed that this followed from thermodynamical considerations; and that when the substance xpanded on solidification, like icc, the melting-point was lowered by pressure, but when the substance coutracted on solidification the melting-point was raised by pressure. The most important substances which expand on solidification are water, bismuth, antimony, and cast-iron; nons of these crystallisss in the regular system, so that we may suppose that the molecules are arranged unsymmetrically, that while some are nearer together | alcohol (Baumann, B. 6, 1371).

than when in the liquid state, others are further apart, producing on the whole an increase of volume.

The specific heat of a body in the solid condition is in general less than when it is in the liquid, except for those substances whose melting and boiling-points are very far apart, when it seems to be about the same in the twostates; if we know the specific heat of a snbstance in both the fluid and the solid state at all temperatures, we can find the amount of heat necessary to convert unit mass of the substance from the solid to the liquid stats. For Clausius has proved that if \(\lambda \) be the latent heat at the temperature t, s_1 and s_2 the specific heats in the solid and liquid states respectively at ths same temperature, then

$$\frac{d\lambda}{dt} + s_2 - s_1 = \frac{\lambda}{t}.$$

A similar equation will apply to the change from the liquid to the gaseous state. Some bodies, such as camplior and iodine, sublims, that is pass directly from the solid to the gaseons state, and as these bodies exhibit a definite vapour-pressure - they must also pass directly from the gaseous into the solid states.

In the solid and liquid states the moleculs is probably a very much more complex thing than the gaseous molecule, it is probably also not nearly so defluite. Maxwell, in the articlenearly so definite. 'Atom' in the Encyclopædia Britannica, shows how the hypothesis of groups of molecules of different degrees of stability would explain the residual effects of clasticity, and states that in his view a solid consists of groups of molecules, some of which are in different circumstances from others.

AGONIADIN C10 II 140 o. A crystalline bitter substance occurring in Agoniada bark (from I'lumeria longifolia), which is used in Brazil as a remedy for intermittent fever. Needles, very bitter, v. sl. sol. ether, v. sol. hot alcohol, and CS. Nearly insoluble in cold, easily soluble in boiling water. When boiled with sulphuric soid it yields a sugar (Peekolt, Ar. Ph. [2] 144, 34).

ALACREATINE C, H, N, O, i. e. NII. C(NII).NH.CIIMe.CO.Ha-guanido-profionio acid. S. 8.3 at 15°. Formed by mixing conc. solutions of alanine and eyanamide, adding a little NH₃, and allowing the mixture te stand (Baumann, A. 167, 83). Small prisms, v. sl. sol. cold alcohol. At 180° it changes into its anhydride, alacreatinine. Boiling baryta-water forms alanine and urea, or its decompositionproducts, CO2 and NH3. HgO oxidises it, forming guanidine.

Methyl-alacreatine . NH.C(NH).NMc.CUMe.CO.II. From a msthylamido-propionic act cyanamide, and a little NII, (Lindenberg, J. v. [2] 12, 253). Monoclinic prisms, sl. sol. cod water or alcohol.

ALACREATININE C.H.N.O aq. Formed by dehydration of alacreatine by the action of heat or dilute acids, crystallises from water in long prisms, which give off aq in dry air or at 100°. M. sol. alcohol, more soluble in water than alacreatine. With zine chloride it forms crystalline scales (C.H.N.O)₂ZnCl₂, S. 4.35 at 20°, v. el ad ALANIME C.H.NO., i.s. OH.OH(NH.)CO.H. a-amido-propionic acia. Mol. w. 89. S. 22 at

17°; S. (cold alcohol of 80 p.c.) 2.

Formation.-1. From ethylic a-chloropropionate and ammonia (Kolbe, A. 113, 220; Strecker, A. 75, 29).—2. From α-bromopropionic acid and alooholic ammonia (Kekulé, A. 130, 18).

Preparation.—An equoous solution of 2 pts. aldehyde-ammonia is mixed with aqueous hydroevanic acid containing 2 pts. HCy; hydrochloric acid is added in execss; tho mixture is evaporated to dryness over a water-bath; the residuo is digested with a mixture of alcohol and other, which leaves N11,Cl undissolved (Strecker).

Properties. - Tufts of colourless needles or oblique rhombic prisms, having a nacreous lustre. Sublimes at 200°. V. sl. sol. cold alcohol, insol. other. The aqueous solution has a sweet taste, docs not affect vegetable colours and gives no precipitates with any of the ordinary reagents. Alanine is isomeric with urethane, lactamido, and sarcosine; distinguished from the two former by not melting below 100°, and from the last by its solubility in water and its behaviour to metallic oxides.

Reactions .- 1. Not altered by boiling with dilute acids or with alkalis .- 2. Fused with KOH, it gives off hydrogen and ammonia and forms cyanido and acetato of potassium .- 3. Resolved by boiling its aqueous solution with PbO, into aldeliyde, carbon dioxide, and ammonia; $C_3H_7NO_2 + O = C_2H_4O + CO_2 + NH_3 - 4$. Decomposed in aqueous solution by nitrous acid, with evolution of nitrogen and formation of lactic acid.

ALANT CAMPHOR C10 1116O. Occurs in elecampane root, and is obtained together with solid alantic anhydride by distilling with water. Liquid smelling like peppermint, boiling at 200°. Heated with P2O3, it yields a hydrocarbon C10H11 which boils at 175°, and is converted by oxidisation with chromic acid into tercphthalic acid (Kallen).

ALANTIC ACID C_{1.}H_{...}O₄ [91°].—Obtained from its anhydride (v. sup.). Slender needles (from alcohol). Dissolves sparingly in cold, more readily in boiling water, very casily in alcohol. The barium salt forms warty masses moderately soluble in water. The silver salt AgC, 11, 0, forms small scales having a silvery lustre (Kallen, B. 9, 155).

Alantic Anhydride C, 11200, [66°], (275°). crystallises from dilute alcohol in prismatic Easily sublimable. Dissolves very sparingly in water, very casily in alcohol, ether, &c.—The chloride C1,1121O2Cl [140°] formed by passing HCl-gas into a solution of alantic acid in absolute scolol, crystallises in large rhombic tablets, melting, with evolution of HCl, at 140°. It unites with Asses, forming salts which readily decompose, with separation of metallic chlorides. By seess of caustic alkali it is converted into the second of the se is converted into dialantic acid, Cooling (?)

The a mide Ci, H2, O2, N112 [210], obtained by passing ammonia-gas into an alcoholic solution of the anhydride, forms small crystals, melting, with decomposition at 210°, slightly soluble in alcohol, resolved by potash into ammonia and plantic acid. H. W.

ALBUMEN v. PROTEÏDS.

ALCAMINES v. ALEANINES. ALCOGEL. A gelatinous compound of silicia acid with alcohol (q, v).

ALCOHOL C.H.O or EtOH ethyl alcohol, Anound Calley of Action (78:2°) at 762-7 mm. (R. Sohiff); (78-3°) (Regnault); (78-4°) at 760 mm. (Kopp, 4. 92, 9); (78-5°) (Perkin); (12-8°) at 20-9 mm.; (21°) at 41-3 mm. (Kahlbaum, B. 16, 2480). S.G. ½ 79367 (S.); ½ 79503 (P.); ½ 78820 (P.); ½ 8000 (Brühl). S.V. 62-18 (S.); 62-7 (Ramsay). (S.) (15.15 (Kopp)) (559 (20°) to 789) Lussac). S.H. 615 (Kopp); 659 (20° to 78° (Rcis); '6019 (16°-20°); '6007 (16°-35°); '6120 (16°-40'5°) (J. II. Schüller, P. Ergbd. 5, 116, 192). H. F. p. 58,470. H. F. v. 57,020 (Th. iv. 229). μ_B 1·3667 R_{∞} 20·31 (B.). M.M. 2·78 (P.).

Name. - The term alcohol was used in the timo of Libavius (1595) to denoto a powder. Spirit dried over powdered potassic carbonate was called spiritus alcolisatus. Kopp (Geschichte, iv. 281) suggests that this term does not mean spirit that has been treated with the powder, but that it is a corruption of spiritus alcalisatus, or spirit that has been treated with alkali. Alcolised or alcoholised spirit was then shortened to alcohol.

Occurrence.-1. In fermented saccharine juices .- 2. In putrid, and even in healthy, tissues, such as ox-brain (Béchamp, C. R. 89, 573) -3. 1n crude coal-tar benzene (about 2 parts per million) (O. N. Witt, C. C. 1878, 416) .--4. In the fruits and juices of some living plants (Gutzeit, A. 177, 314).—5. In bread (Bolas, C. N. 27, 271).-6. In crude wood spirit (V. 11emilian, B. 8, 661).-7. Together with acctone, in the urine of diabetic patients (Markownikoff, B. 9, 1441, 1603),

Formation.-1. By the decomposition of glucose under the influence of ferments (v. Fermentation): $C_6 11_{12}O_6 = 2C_2H_6O + 2CO_2$. Levulose, maltose, and melitose also give alcohol on fermentation .- 2. From olefiant gas by dissolving it in conc. H₂SO₂, diluting and distilling (Hennel, P. M. 1826, 240; Berthelot, A. Ch. [3] 43, 385): $C_2\Pi_4 + H_2SO_4 - C_2\Pi_5SO_4H$

 $C_2H_3SO_4H + H_2O = C_2H_3OH + HSO_4H$ The absorption of cthyleno is greatly facilitated by heating the H,SO, to 100° or, better still, 170° (Goriainow a. Butlerow, A. 169, 147).-3. By reduction of acetic anhydrido (Linnemann, A. 148, 249), acetyl chloride (Saytzeff, J. pr. [2] 3, 76), or aldchyde, by means of sodiumamalgam.—4. By heating ether at 170° with water slightly acidulated with H.SO. (Erlenmeyer a. Tscheppe, Z. [2] 4, 343).

Preparation. - When aqueous solutions of grape-sugar are fermented by yeast, 95 p.c. of the sngar splits up into alcohol and carbonio acid, but 4 p.c. goes to form succinic acid and glycerin, while 1 p.c. is used by the yeast as food. Small quantities of n-propyl, iso-butyl, and the two isp-amyl, alcohols, Me, C11.CH, CH, OH and MeEtCH Cll OH, are also formed. The mixture of these four alcohols is known as fusel oil. According to Rabuteau (C. R. 87, 500), potato spirit contains also iso-propyl, n-butyl and secondary amyl alcohols.

The liquid to be fermented must contain nitrogenous matter and some inorganic salts to scrve as food for the yeast; grape-juice, or a mixture of water with germinating barley (malt), to which a mash of potatoes may be added, are unorganised ferment in malt, converts the starch of the potatoes into a sugar, maltose, which then undergoes alcoholic fermentation.

When any of these elcoholio liquids are distilled the first portions of the distillate are rich in alcohol. By repeated rectification 'rectified spirit' containing 91 p.o. of alcohol may be got. Fusel oil may be removed by adding to the spirit about 7 of its weight of ccarsely powdered charcoal, leaving the mixture to stand for several days, and stirring repeatedly, then decanting and distilling. Animal or blood charcoal may also be used.

Absolute Alcohol.—The last traces of water may be removed by repeated rectification over freshly heated K.CO., CaO, BaO, CuSO., or CaCl. The best way is to digest strong spirit with quick lime at 40° for two hours, and then, on distilling, to reject the first and last portions (Mendelécff, Z. 1865, 260). If the spirit contain more than 5 p.o. of water a second treatment with lime will be necessary (Erlenmeyer, A. 160, 249). If dry baryta be used to complete the drying, as soon as the alcohol is absolute it will become yellow, dissolving a little BaO (Borthelot, J. 1862, 392).

References.-C. Bullock, Ph. [3] 4, 891; J. L. Smith, Am. Ch., 5 120; Dittmar a. Stewart, C. N. 33, 53; Friedel a. Crafts, A. Ch. [4] 9, 5. Properties .- A transparent, colourless, mobile, liquid. It has a characteristic odour and burning taste. When undiluted it acts as an inflammatory poison. It solidifies at -130.5° (Wroblewsky a. Olszewsky, M. 4, 338). Very hygroscopic. Miscible with water. Burns with a pale flame. Snow (1 pt.) mixed with alcohol (2 pts.) produces a freezing mixture $(-20^{\circ}, Bn.)$

1, 237). Alcohol dissolves fats, oils, resins, alkaloids and most organic substances. It dissolves CaCl, and SrCl₂ but not BaCl₂; Ca(NO₃), but not Sr(NO₃), and Ba(NO₃)₂; LiCl but not KCl and NaCl. It does not dissolve carbonates or suiplates. It dissolves I, Br, P, and S.

The critical point of alcohol is 234.6° at 48,900 mm. At this point 1 g. occupies 3.5 e.c. (Ramsay a. Young, Pr. 38, 329). Alcohol vapour in contact with liquid acquires its normal density, 23, at 50° (R. a. Y.).

When alcohol is mixed with water contraetion takes place and heat is evolved. Tho maximum contraction occurs when 49.8 vols. water and 53.9 vols. alcohol at 0° produce 100 vols. of mixture instead of 103.7 vols. This corresponds to a possible compound, EtOll 3aq (Mendeléeff, Z. 1865, 262).

The greatest difference between the observed specific heats of solutions of alcohol and the values calculated on the assumption of merc mixture occurs in a solution containing about 80 p.o. of alcohol by weight, corresponding to the formula Et011 6aq. The greatest difference between the observed and calculated boilingpoints and between observed and calculated capillarity also occurs in the same mixture, but the maximum deviation from calculated (or mean) compressibility is exhibited by a solution containing 40 p.c. of alcohol by weight (Dupré a. Pago, Pr. 17, 833; P. M. [4] 38, 158). The maximum rate of transpiration through capillary

the liquids usually employed. Diastase, an tubes is exhibited by the solution EtOH 3aq (Graham, A. 123, 102).

Detection of Water in Alcohol .- 1. CuSO ought not to turn blue (Cassoria) .- 2. Benzene ought not to form a cloudiness, due to waterdrops (Gorgen, O. R. 30, 691) .- 3. Wet alcohol produces a pp. of BaO when added to a solution of BaO in absolute alcohol (Berthelot, A. Ch. [3] 46, 180) .-- 4. If alcohol be added to a mixture of anthraquinono ('001 g.) with a little sodium amalgam, a green coloration indicates absence of water, otherwise a red colour is produced (Claus, B. 10, 927).

The following table gives the percentages of absolute alcohol, determined by Tralles:

Volumes	Weights	Specific	. Volumes	Weights	Specific
per	per	gravity	per	per '	gravity
cent.	cent.	at 15.56°	cent.	cent.	nt 15.56°
		·			
0	0	1.0000	51	43.47	·9315
1	0.80	9976	52	44.42	.0295
2	1.60	9961	53	45.36	.9275
3	2.40	.9947	54	46.32	9254
4	3.20	•9933	55	47.29	.9234
ŝ	4.00	9919	56	48.26	9213
6	4.81	-9906	57	49.23	9192
7	5.62	•9893	58	50 21	
	6.43	9881		51.20	9170
8			59		9148
9	7.24	.9869	60	52.20	9126
10	8.05	9857	61	53 20	.9104
11	8.87	9845	62	54.21	·9082
12	9.69	.9834	63	55.21	•9059
13	10.51	9823	64	56.22	.9036
14	11.33	.9812	65	57.24	·9013
15	12.15	9802	66	58.27	·8989
16	12.98	9791	67	59.32	·896 5
17	13.80	9781	68	60.38	*8941
18	14.63	9771	69	61.42	·8917
19	15.46	9761	70	62.50	8892
20	16.28	.9751	71	63.58	8867
21	17.11	9741	72	64.66	8842
22	17.95	9731	73	65.74	8817
23	18.78	.9720	74	66.83	·8791
24	19.62	9710	75	67.93	8765
25	20:16	.9700	76	69.05	.8739
26	21.30	•9689	77	70.18	8712
27	22.14	9679	78	71.31	8685
28	22.99	9668	79	72.45	8658
29	23.84	9657	80	73.59	8631
30	21.69	9646	81	71.74	8603
31	25.55	9634	82	75.91	·8575
32	26.41	9622	83	77.09	8547
33	27.27	9609	84	78.29	8518
34	28.13	•9596	85	79.50	-8488
35	28.99	9583	86	80.71	
36	29.86	9570	87	81.94	8458
		9556			8428
37 38	30.74		88	83 19	·8397
	31.62	9541	89	81.46	·8365
39	32.50	9526	90	85.75	.8332
10	33.39	\$10	91	87.09	8299
5.ª	34.28	9244	92	83.37	8265
49	35.18	.9178	93	89.71	.8230
43	36.08	9161	94	91.07	.8194
41	36.99	9114	95	92.46	8157
45	37.90	9127	96	93.89	·8118
46	38.82	.9109	97	95.34	·8 077
47	39.75	·9391	98	96.81	·8034
48	40.66	.9373	99	98.39	7988
49	41.59	.9354	100	100.00	·7939
50	12.52	•9835			
3					-

The specific gravity of squeous alcohol is given by Mendelsess (P. 188, 108, 280) as follows:

Weight p.c.	Specific	Gravity, re	ferred to Wa	ster at 40
Alcohol	at 0°	at 1000	at 20°	at 30°
0	•99988	•99975	•99831	.99579
- 5	•99135	99113	•98915	98680
10	·98493	•98409	•98195	97892
15	97995	.97816	•97527	97142
20	97566	97263	.96877	96413
25	97115	.96672	•96185	95628
80	.96540	.95998	.95403	.94751
35	95784	95174	.94514	.93813
40	·94939	.94255	.93511	.92787
45	.93977	93254	.92493	91710
50	92940	.92182	•91400	90577
55	.91818	.91074	·90275	89456
60	90742	89944	*89129	*88304
65	89595	88790	·87961	87125
70	*88420	·87613	86781	85925
75	87245	86427	85580	84719
80	86035	*85215	*84366	*83483
85	84789	-83967	83115	*82232
90	*83482	82665	·81801	.80918
95	*82119	·81291	80433	•79553
100	·80625	·79788	·78945	.78098

The following table is given by Fownes (Manual, 3rd ed. 591), the specific gravities being taken at 15.6° C.:

Percent- age by Weight	Specific Gravity	Percent- age by Weight	Specific Gravity	Percent- age by Weight	Specific Gravity
0.5	0.9991	34	0.9511	68	0.8769
1	0.9981	85	0.9490	69	0.8745
2	0.9965	36	0.9470	70	0.8721
8	0.9947	37	0.9452	71	0.8696
4	0.9930	38	0.9434	72	0.8672
5	0.9914	39	0.9416	73	0.8649
6	0.9898	40	0.9396	74	0.8625
. ř	0.9884	41	0.9376	75	0.8603
· 8	0.9869	42	0.9356	76	0.8581
9	0.9855	43	0.9335	77	0.8557
10	0.9841	44	0.9314	78	0.8533
11	0.9828	45	0.9292	79	0.8508
12	0.9815	46	0.0270	80	0.8483
18	0.9802	47	0.9249	81	0.8159
14	0.9789	48	0.9228	82	0.8134
15	0.9778	49	0.9206	83	0.8408
16	0.9766	50	0.9184	84	0.8382
17	0.9758	51	0.9160	85	0.8357
18	0.9741	52	0.9135	86	0.8331
19	0.9728	53	0.9113	87	0.8305
20	0.9716	54	0.9090	88	0.8279
21	0.9704	55	0.9069	89	0.8254
22	0.9691	5 6	0.9047	90	0.8228
23	0.9678	57	0.9025	91	0.8199
24	0.9665	58	0.9651	92	0.8172
25	0.9652	59	8979	93	0.8145
26	9.9638	60	0.8956	94	0.8118
27	0.9623	61	0.8932	95	0.8089
28	0.9609	62	*0.8908	96	0.8061
29	0.9593	63	0.8886	97	0.8031
80	0.9578	64	0.8863	98	0.8001
81	0.9560	65	0.8840	99	0.7969
82	0.9544	66	0.8816	100	0.7938
88	0:9528	67	0.8793		

Proof spirit was a tarm originally intended to denote spirit that was just strong enough to ignite gunpowder when burnt upon it, but it was defined by law in the reign of George III. to be spirit 'such as shall at the temperature of 51° K weigh exactly twelve-thirteenth parts of an equal amount of distilled water.' It has, therefore, S.G. 920 at 156°C, and contains 49-24 pts. alcohol to 50.76 pts. water by weight, or 100 vols, alcohol to 81.82 vols. water.

Alcoholic Drinks.—Beer contains from 2 to 6 p.c. of alcohol; hock and claret from 8 to 10 p.c.; port and sherry from 15 to 20 p.c.; gin,

rum, and whisky from 51 to 54 p.c.

Detection .- 1. The liquid supposed to centain alcohol is repeatedly rectified, after drying with K2CO3. The alcohol is recognised by its boiling-point, and by converting it into ethyl iodide, and noting the boiling-point of the iodide (72°) .- 2. The suspected liquid is distilled and some of the distillate (8 o.c.) mixed with water (10 c.c.) and H₂SO₄ (5 o.e.); some KMnO₄Aq, and after five minutos a solution of magenta, bleached by SO2, are added. A red colour indicates that aldehyde had been formed by the oxidation of the alcohol. Acetone, formic acid and methyl alcohol do not show this reaction, so that it may be used to detect ethyl alcohol in wood spirit. Other primary alcohols behave more or less like ethyl alcohol (Riche a. Bardy, C. R. 82, 768).-3. An aqueous solution of alcohol warmed with KOH and iodine deposits This 'iodoform reaction' is given also by aldehyde, acctone, n-propyl, n-butyl, sec. butyl, and octyl alcohols, by propionic and butyric aldehydes, by lactic, quinic, and meconic acids, by acctophenone, mothylic butyrate, acetic other, and oil of turpentine.

The 'iodoform reaction' is not given by methyl and amyl alcohols, by formic, acetic, butyrie, valeric, oxalic, succinic, malic, tartaric, racemic, citrie, pyrotartaric, suberic, sebacic, uric, mucic, isethionic, benzoic, salicylic, anisic, cinnamic, and picric acids, phenol, valeric aldohyde, benzoic aldehyde, glycol, glycevin, mannite, glycocoll, leucine, chloral, ethyl chloride, ethylene chloride and bromide, chloroform, tetrachloride of carbon, sulphide of carbon, toluene, and ether (Lioben, A. Suppl. 7, 226). Sugar and dextrin give a small amount of iodoform.

The formation of ethyl acetate and beuzcate is also recommended as a test for alcohol.

Estimation.—The liquid is distilled and the S.G. of the distillate taken.

Detection of Fusel Oil.—1. The liquid is diluted with water until it contains about 12 p.c. alcohol; it is then shaken with chloroform. This extracts the amyl alcohol, which it leaves behind on evaporation; by warming with KOAc and H₂SO₄ this is converted into amyl acctate, smoking like poar-drops.—2. The alcohol is diluted until it forms a 50 p.c. solution. 100 c.c. are then shaken with 20 c.o. chloroform at 15° in a graduated cylinder. If the chloroform layer is 37·1 c.c. the alcohol is free from higher

layer is 37.1 c.c. the alcohol is free from higher homologues, but if it occupy a larger volume, fusel oil is present. Thus 39.1 c.c. indicates 1 p.o. amyl alcohol (Röse, B. 19, R. 154).—3. The height to which the alcohol will rise in capillary tubes of known diameter is observed.

Pure alcohol rises higher than alcohol adulterated with fusel oil (J. Traube, B. 19, 892).

Reactions. -1. Potassium and sodium not upon alcohol, evolving hydrogen and forming EtOK and EtONa respectively. -2. Phosphorus trichloride forms EtCl, HCl, ethyl phosphite, and phosphorous acid (Béchamp, C. R. 40, 944); 6EtÓH + 2PCl, = 3EtCl + 3HCl + Et, PO, + H, PO, A smaller quantity of PCl, acts in the cold thus: PCl, +HOEt = PCL, (OEt) + HCl (Menschutkin, A. 139, 343). -3. PCl, reacts thus: EtOH + PCl, = EtCl + ClH + PCl₃O.

4. P2S, produces mercaptan:

 $5EtOH + P_2S_3 = 5EtSH + P_2O_3.$

5. Alcohol coagulates albumen, and, on this account and partly by arresting the development of low organisms, it prevents the putrefaction of dead animal matter.—
6. Vapour of alcohol passed through a red hot tube produces CO., water, hydrogen, CH., C2H. naphthalene, and charcoal. If the tube be filled with pumice, benzene, phenol, and perhaps also aldehyde and acctic acid, are also formed (Berthelot, A. Ch. [3] 33, 295; A. 81, 108).—7. Zinc dust at 300°-850° forms ethylene and hydrogen: $C_2H_4O + Zn = ZnO + C_2H_4 + H_3$. Alcoholvapour passed over zino dust at a dul! red heat forms CO, CH, and H2 (Jahn, M. 1, 378).-8. Alcohol scarcely conducts an electric current, but when acidulated with 5 p.c. H2SO, tho current passes, hydrogen comes off at one pole and, at the other, aldehydo, ethyl formate sulphate and acetate, together with small quantities of acetal, and CH, CH(OH)(OEt) are formed (Renard, A. Ch. [5] 17, 295). Alcohol containing a little potash produces hydrogen at the negative pole and aldeliyde-resin at the positive pole (Connell).—9. Alcohol burns with a pale flamo forming CO, and H₂O. Alcohol vapour undergoes rapid, but incomplete, combustion when mixed with air and exposed to finely divided platinum; acetic acid, aldeliyde, formio acid, acetal, and acctie ether are formed. Hence a coil of red hot platinum wire will keep red hot if placed round the wick of a spirit lamp that is not burning (glow-lamp of Sir 11. Davy) .- 10. Finely divided rhodium, iridium, and ruthenium, in presence of an alkali, decompose alcohol, with elimination of H and formation of an acciate (Deville a. Debray, C. R. 78, 1782),-11. Oxygen does not attack cold pure alcohol, but ozone forme acetic and formic acids (Boillot, C. R. 76, 1132).-12. Chromic acid mixture oxidises alcohol to aldehyde and acetic acid.- 13, An ammoniacat solution of CuO at 180° attacks alcohol, forming acctio acid and Cu2O (A. Letellior, C. R. 89, 1105) .- 14. KMnO, in acid, but not in alkaline, colution forms aldehyde and acetio acid (Chapman a. Smith, C. J. 20, 301). 15. Strong nitric acid acts violently, giving off copious red fumes containing nitrous other, nitric oxide, CO, aldehyde, acetic and formio acids. If the action be moderated by making three layers of funing HNO, water, and alcohol, and allowing them to mix slowly by diffusion, the following bodies are formed: aldehyde, acetic acid, glyoxal, glyoxylic acid, glyoollic acid, and oxalic acid (Debue) .- 16. In presence of Vol. L

tion of fulminate of mercury (q. v.); in a similar way fulminate of silver may be made.

If mercury (1 pt.) be dissolved in HNO. (12 pts.) (S.G. 1.3) and the liquid left for some days till no more nimous fumes appear and the liquid is colourless, and then alcohol (12 pts. of S.G. 8) be added and the mixture be warmed, a pp. is produced which is not mercurie fulminate. th may be crystalised from diluted (4 vols.) HNO₃ (1 vol.). It is C₂H₂Hg₃O₄(NO₃). At 120°-130° it explodes. It is insoluble in water, alcohol, and ether. Potash converts it into C₂H₂Hg₃O₄(OH)₃; while cold K₂C₄O₄(NO₄O₄(NO₄)). elowly converts it into the oxalate, C. 11, Hg, O, C,O,, a body which is browned by sunlight. A mixture of HNO₃ and alcohol converts it into mercuric fulminate (Cowper, c. J. 39, 242; v. Gerhardt, A. 80, 101).—18. Chlorine is rapidly absorbed by alcohol, and in sunlight the liquid may even take fire. The nltimate product is chloral alcoholate, CCl3.Cl1(OEt)(OH), but this is probably the result of a long scrice of reactions (v. Chloral). Besides chloral, there are formed HCl, aldebyde, acetal, acetic acid, EtCl and other chlorinated bodies .- 19. Bromine forms HBr, water, EtBr, bromal, and bromal alcoholate (E. Hardy, C. R. 79, 806).-20. Dry chlorins passed into alcohol mixed with K.Cr.O, gives aldehyde, EtCl, acetyl chloride, and EtOAc (Godefroy, Bt. [2] 40, 168).—21. When alcohol is distilled with much water and bleaching powder, chloroform (q. v.) is formed. When bleaching powder (300 grms.) is mixed with absolute alcohol (67 grms.) in 10 minutes the mixture gets hot and alcohol distils over together with a green oil, which explodes when exposed to sunlight or heated, and among the products of the explosion are mono- and dichloro-acetal (Schmitt a. Goldberg, J. pr. [2] 19, 393), aldehyde, and small quantities of chloroform (Goldberg, J. pr. [2] 21, 97).-22. Hydric chloride produces ethyl chloride: EtOH + 11Cl = EtCl + 112O.

Bit when excess of alcohol is used and the solution heated in a scaled tube at 240°, ether is also formed: EtOH+ClEt=Et_.O+HCl (Reynoso, A. Ch. [3] 48, 385),—23. Sulphura acid mixes with alcohol with evolution of heat and formation of hydrogen ethyl sulphate:

EtoH + H,SO, = EtHSO, + H,O.

About half the alcohol and H,SO, take part in
the reaction; when more dilute acid ie used
hydrogen ethyl sulphate is not formed until
heat is applied. If a mixture of alcohol with
an equal volume (or less) of H,SO, be heated,
a further reaction sets in at 1202-1502, ether
and water distilling over; this is due to action
of alcohol upon hydrogen ethyl sulphate
(Williamson, C. J. 4, 106, 2227; v. Ernea);

16. Strong nitric acid acts violently, giving off copious red fumes containing nitrous other, nitric oxide, CO,, aldelyde, acetic and formio acids. If the action be moderated by making three layers of fuming HNO,, water, and alcohol, and allowing them to mix slowly by diffusion, acetic acid, glyoxalic acid, glyoxylic acid, glyoxlic ac

C.H. SO O, or 'carbyl sulphate,' together with ethionic, isethionic, and sulphuric acids, and HEISO,—25. Heated with sulphurous acid at 200° it forms HEISO, ether, H.SO, mercaptan, and S (Pagliani, J. 1878, 518).—26. CISO,H forms EHSO, and other bodies (Baumstark, A. 140, 75).—27. When alcohol is drawned upon but give Vibraits the greater party page. dropped upon hot zinc Chlorice the greater part is decomposed in accordance with the equation :

 $2C_2H_0O = C_2H_1O + H_2 + C_3H_1 + H_2O$. Hydrogen, ethane, HCl, and polymerides of aldehyde are also formed (W. H. Greene, C. R. 86, 1140). When wet alcohol is heated with ZnCl, at 155°, ether is formed, as well as EtCl, basic zinc chloride being left .- 28. Phosphoric acid mixed with alcohol forms some di-hydrogen ethyl phosphate, EtH.PO₄. Alcohol heated with P₂O₅ forms HEt.PO₄ and Et.₂PO₄ (Carius, 4. 137, 121).—29. Alcohol heated with B₂O₅ forms EtBO₅ and Et.₂BO₅.—30. Phosphorus sulpho-chloride, PSCl., forms di-hydrogen ethyl this phosphate. thio phosphate (Chevrier, Z. [2] 5, 413):

 $PSCl_a + 3HOEt = PS(OH) \cdot (OEt) + 2EtCl + HCl.$ 31. Chloride of sulphur, S.Cl., acts upon alcohol forming ethyl chloride, ethyl sulphite. and a small quality of mercaptan (Carius, A. 106, 316).—32. Chloride of antimony dissolves in alcohol; if the solution be heated to 150' the following reaction ensues (H. Schiff, A.

Suppl. 5, 218):

 $SbCl_1 + 4EtOH = SbOCl + 2EtCl + Et_0 + 2H_0$. 33. Heated with carbon tetrabromide at 100° for 12 hours, bromoform is produced (Bolas a. Groves, C. J. [2] 9, 781): $CBr_1 + C.H_2O = CHBr_2 + C_2H_2O + HBr$. Alcohol here acts as a reducing agent, as it does also in the next reaction. - 34. Heated with a di-azo salt, nitrogen is evolved and the entire di-azo group displaced

by hydrogen:

 $\mathbf{C}_{e}\mathbf{H}_{s}^{T}\mathbf{N}_{2}\mathbf{C}\mathbf{I} + \mathbf{C}_{2}\mathbf{H}_{s}\mathbf{O} = \mathbf{C}_{e}\mathbf{H}_{3}\mathbf{H} + \mathbf{N}_{2} + \mathbf{H}\mathbf{C}\mathbf{I} + \mathbf{C}_{2}\mathbf{H}_{1}\mathbf{O}.$ In some cases the di-azo group is displaced by ethoxyl-35. Heated with ammoniacal zinc chloride at 260°, alcohol is converted into a mixture of mono-, di-, and tri-ethylamine; the yield of mixed bases amounts to 45 p.c. of the alcohol used (Merz a. Gasiorowski, B. 17, 637). -36. Zinc acctate heated with excess of alcohol at 100° is converted, in about 30 hours, into zine ethyl acetate and zinc oxide (Kraut, A. 156, 323). 87. When stannic chloride is distilled with alcohol, ether and EtCl pass over at 140°-170°; afterwards a compound of EtCl with SnCl, (Kuhlmann, A. 33, 106, 192). - 38. Crystallised stannous chloride distilled with alcohol yields ether, but no EtCl (Marchand); the same decomposition takes place in a scaled tube at 240°. Crystallised ehloride of manganese, and ferrous chloride also etherise alcohol completely at 240°; the chlorides of cadmium, nickel, and cobatt partially (Reynoso, A. Ch. § 2] 48, 385).—39. Platinic chloride (1 pt.) dissolved in alcohol (40 pts.) of S.G. 82 and distilled to 1, yields aldehyde, ethylchloride, HCl, and the solution contains the st-called inflammable chloride of platinum C2H4PtCl2, which is left as a sticky mass when the liquid is evaporated (Zcise, P. 9, 632; 21, 498; 40, 249) .- 40. Platinous chloride boiled with alcohol forms a black explosive powder called detonating platinum deposit, C,11,PtO(?) (Zeise, loc. cit.) .- 41. Mercuric chloride is slowly reduced to calomel by alcohol. 43. Alcohol hested with soda lime, air being excluded, is converted into sodic scetate, with evolution of hydrogen; at a higher temperature the sodic acctate breaks up into sodic carbonate and marsh gas.—43. Chloride of cyanogen is readily absorbed by alcohol but does not decompose it immediately. After a few days, or more quickly at 60°, NH,Cl separates, and the liquid then contains ethyl chloride, ethyl carbamats (or urethane), and ethyl carbonato (Wurtz, A. 79, 280).

Combinations.-Alcohol combines with many salts, acting like water of crystallisation .-SbCl, EtOH [67°] needles (from alcohol); resolved by heat into HCl, EtCl, Sb.O., and SbCl, Soluble in ether and chloroform, but decomposed by water (W. C. Williams, C. J. 30, 463).— AsCl₃EtOII (148°): liquid; fumes in the air; decomposed by water (Luynes, A. 116, 368).-CaCl, 4EtOII: got by cooling an alcoholic solution of CaCl, with ico.—CaCl, 3EtOH: got by evaporation of such solution over H, SO, (Heindl, M. 2, 207).—LiCl4EtOH (Simon, J. pr. [2] 20, 376).—MgCl,6EtOH (S.).—Mg(NO,),6EtOH (Chodnew, A. 71, 256): a crystalline mass deposited from boiling solution.—PtCl,2EtOH (Schützenberger, J. 1870, 388).—SnCl,2EtOH: crystals formed by evaporation over H,SO. (Lewy, C. R. 21, 371; Robiquet, J. Ph. [3] 26, 161); heated with acids, this compound readily forms ethyl salts.—TiCl,EtOH [105°-110°] erystals, decomposed by water (Demarcay, B. 8, 75).

Alcoholates or Ethylates are formed by displacing the typical hydrogen by motals. They are decomposed by water:

 $MOEt + H_vO = HOEt + MOH.$

Aluminium ethylate Al(OEt)3 [130°] S.G. 1.147. Aluminium does not attack alcohol, but if iodino be present and the liquid be warmed, hydrogen is evolved and aluminium ethylate is formed (Gladstone a. Tribe, Pr. 30, 546): $2AI + 6HOEt = Al_2(OEt)_s + 3II_2$. The reaction probably takes place in three stages:

SHOEt + All Is = Al Is (OEt) + SHI

 $Al_2(OEt)_3I_3 + 3HOEt = Al_2(OEt)_6 + 3HI$ $\tilde{A}l_2 + 6H1 = Al_2I_4 + 3H_2$

Aluminium (4g.), iodine (2g.), and alcohol (40 c.c.), are heated in a flask with inverted condenser; when no more H comes off, the contents are distilled in vacuo at 300°. (Good yield (12 g.). G. a. T., C. J. 39, 2). When aluminium ethylate has been fused it remains liquid for a long time even at 70°. It is decomposed by water thus:

 $Al_2(OEt)_6 + 6II_2O = Al_2(OII)_6 + 6HOEt.$ When distilled under atmospheric pressure it decomposes: $Al_2(OEt)_4 = Al_2O_3 + 3C_2II_4 + 3IIOEt$, (G. a. T., C. J. 41, 5).

Barium othylate Ba (QEt)2Aq (Berthelot, A. Ch. [3] 46, 180), Ba(OEt), Ba(OH), (Dostrom, A. Ch. [5] 27, 8, 22; C. R. 90, 1213). A granular pp. formed by boiling an alcoholic solution of BaO, or by heating alcohol with BaO in a digester at 150°. A white powder, turned yellow by oxidation. Convorted by CO₂ into baric ethyl-carbonate. Destructive distillation gives C2H4, methane, H, and BaCO2

Calcium othylate Ca(OEt), resembles the

barium compound.

Ferricethylate *Fe2(OEt), (?).-When the proper quantity of sodic ethylate is added to an pleisoile solution of Fe,Cl, all the chlorine is ppd. as NaCl, and the filtrate leaves, after evaporation, a black pasty mass, sol. alcohol, McOH, etherabenzene, chloroform, or benzoline (Grimaux, C. R. 98, 105). A solution of ferric thylate poured into water produces a solution of colloidal ferric hydroxide.

Potassium ethylate KOEt.—Similar in sharacter to sodie ethylate.

Sodic othylate NaOEt.—When sodium is dissolved in dry alcohol, II is evolved, and ultimately crystalline laminæ of NaOEt 2HOEt separate. If the solution be evaporated in vacuo at 20° needles of NaOEt 3HOEt are got (Fororand, Bl. 40, 177). The alcohol of crystallisation may be driven off at 180°.

tion may be driven off at 180°. Reactions .- 1. When mixed with water and dietilled, alcohol passes over and NaOII is left .-2. Converted by Etl into other (Williamson) .-3. Forms ether when it acts on EtNO3: but it acts like Na upon ethers of organic acids; thus it converts formic ether into CO and alcohol, oxalic ether into CO and carbonic ether, carbonic ether into NaCO, Et and Et.O, benzoic ether into NaOBz and Et.O, acctic other into sodium aceto-acetic ether (teuther).-4. CO combines with NaOEt at 100° forming sodie propionate. Carbonic oxide passed over a mixture of NaOEt and NaOAc at 205° produce n-butyric acid, diethyl-acetic acid, mesitylenic acid, an acid C₁₉H₁O₂ (250°-260°), and two kotones C₂(I₁₉O and C₁₅H₂O (Geuther, A. 202, 305).—5. PCI₂ gives NaCl, PO(OE1)₂, and EtCl.—6. With chloroform it forme ortho-formic ether, CH(Olit), (Williamson a. Kay, C. J. 7, 224).—7. Chlorine forms aldehyde and acetic acid (Maly, Z. [2] 5, 345) .- 8. Bromine forms bromal, EtBr, and acctic ether (Barth, B. 9, 1456).-9. Chloroacctic acid forms sodium cthyl-glycolate (Heintz, P. 109, 301). - 10. Iodine forms Nal, sodie formate, and iodoform.-11. Iodoform is reduced by NaOEt to methylene iodide.—12. Nitrobenzene

aniline (Béchamp a. Saint-Pierre, C. R. 47, 24).
Thallium ethylate *TIOEL S.G. 3-5 to
3-685. Formed by heating EtoH with thallium
at 100° (Church), or by exposing thallium to the
vapour of alcohol in a bell-jar full of oxygen
(Lamy, A. Ch. (4) 3, 378). It may be solidified
by great cold. It dissolves in dry alcohol or
ether, but addition of a trace of water causes
separation of thallous hydrate. TIOEL is slowly
decomposed by CHCl, with separation of TICI.

is reduced to azoxybonzene, azobenzene, and

ALCOHOLS.—The term alcohol, originally limited to one substance, viz. spirit of wine, is now applied to a large number of compounds, many of which, in their external characters, exhibit but little resemblance to common alcohol. All alcohols are compounds of carbon, hydrogen, and oxygen, and are derived from hydrocarbons containing even numbere of hydrogen-atoms by sabstitution of one or more hydrogen-atoms: thus from propane C₃H₂ or CH₃.OH₂.CH₃, are derived the three following alcohols:—

 $\begin{array}{cccc} \text{Propyl alcohol} & \textbf{C}_1\textbf{H}_1\textbf{O} & = \textbf{C}_2\textbf{H}_1(\textbf{OH}) \\ \text{Propylene alcohol} & \textbf{C}_2\textbf{H}_2\textbf{O}_2 & = \textbf{C}_3\textbf{H}_2(\textbf{OH})_2 \\ \text{Proponyl alcohol} & \textbf{C}_2\textbf{H}_2\textbf{O}_3 & = \textbf{C}_2\textbf{H}_2(\textbf{OH})_3 \\ \text{of glycerin} & \textbf{C}_3\textbf{H}_3\textbf{O}_3 & = \textbf{C}_3\textbf{H}_2(\textbf{OH})_3 \\ \end{array}$

Alcohols are classed as monohydric, dihydric, trihydric, &c., or generally as mono-

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alcoholic solution of Fa,Cl., all the eblorine is and poly-hydric, according to the number of ppd. as NaCl, and the filtrate leaves, after hydroxyl-groups which they contain.

An alcohol is saturated or unsaturated according to the nature of the hydrocarbon from which it is derived. Thus, all the three alcohols derived from propane C₂H₂, which is a saturated hydrocarbon, are themeelves saturated moleculee not capable of forming addition-compounds; but *from** the nasaturated hydrocarbon C₂H₄ is derived tho unsaturated compound allyl alcohol, C₃H₄O or C₃H₃(OH), which is capable of taking up 2 at bromine and forming the compound C₃H₄Br₂O.

The replacement, partial or total, of the hydroxyl in an alcohol by Cl, Br, I, or F, gives rise to haloid ethers; thus:

From C₃II₁(OH) are derived C₃H₁Cl, C₃H₂Br,

From C₃H₆(OII)₂ are derived C₂H₂Cl(OH),

C₃H₂Cl₂, &c.

From C₃H₃C(OII)₃ are derived C₄H₃Cl₄(OIH)₄,
C₃H₃Cl₄(OH)₄, Cl₃, &c. These substitutions are effected by treating the alcohols with the chlorides, brounides, and iodides of hydrogen and phosphorus, as in the formation of ethyl chloride from cthyl alcohol:

 $\begin{array}{c} C.II.OH + HCl = H(OH) + C_2H_3CI\\ 3C.H_3(OII) + PCl_2 = P(OII)_3 + 3C.H_3CI\\ 3C.II_3(OH) + POCl_2 = PO(OII)_3 + 3C_2H_3CI. \end{array}$

Instead of the bromides and iodides of phosphorus, a mixture of phosphorus and bromine or iodine, in the proportions required to form them, are often used in these processes.—The haloid ethers are also formed in many instances by direct substitution of chlorine &c. for hydrogen in hydrogerbons.

The treatment of the alkyl chlorides, bromides, or iodides with aqueous caustic alkalis gives rise to a substitution opposite to that shown in the above equations, reconverting the ethers into alcohols; e.g. C₂H₃Cl+KOH=KCl+C₂H₄(OH). A considerable portion of the alcohol thus formed is, however, converted by delaydration into the corresponding olefin; e.g. C₄H₄O+L₄O=C₅H₆. A better yield of alcohol is obtained by heating the haloid ether with moist silver oxide, which acts like a hydroxide AgOII; and a still better method ie to convert the alcoholic chloride, &c. into an acetate by heating it with silver acetate or potassinm acetate, and to boil the resulting alkyl acetate with caustic potash or soda;

C.H.Cl+KC.H.O. = KCl+C.H.C.H.O. C.H.C.H.O. + KOH = KC.H.O. + C.H.OH. This reaction is of great importance in the preparation of some of the higher alcohole.

The replacement of the hydroxyl in an alcohol by the corresponding radiclee, methoxyl OCH₃, ethoxyl OC₂H₃, &c. or of the hydrogen in the OH by Me, Et, &c., gives rice to simple or mixed alkyl oxfles or ethers: thus EtOH yields EtOK, EtOMo, and EtOEt; and ethylene alcohol C₂H₄(OH)₂ yielde C₂H₄(OH)(©Et) and C₂H₄(OEt)₄. These substitutions may be effected in various waye, the simplest being to replace a H-atom in the alcohol by K or Na, and act on the resulting compound with a haloid ether; e.g.

2C₂H₁(OH)₂+Na₂-2C₂H₁(OH)(CNa) + H₂; C₂H₂(OH)(ONa) + EtI = NaI + C₂H₂(OH)(OEt). These oxides may be looked upon as anhydrides formed by elimination of one molecule of water from two molecules of the same or different aloohols.

In the polyhydrio alcohols, where the two hydroxyls occur in the same molecule, the slimination of water gives rise to another class of oxides; thus from ethylene alcohol C,H,(OH), is

derived ethylene oxide C2H4O.

The replacement of the hydrogen in an alcohol by acid radicles produces alkyl salts (also ealled compound ethers or esters); thus from methyl alcohol, MeOH, are derived a nitrate MeONO, an acctate MeOAc, an acid sulphate Me.O.SO, H and a normal sulphate (MeO), SO,. These alkyl salts may also be derived from the corresponding acids by substitution of alkylradioles for hydrogon, being indeed related to the alcohols in the same manner as nictallic salts to metallie hydroxides. They may also be looked upon as anhydrides formed by climination of a molecule of water between one molecule of an alcohol and one molecule of an acid. By distillation with alkalis they are resolved into acid and alcohol; e.g.

EtOAc + KOH = KOAc + EtOH.

MONOHYDRIO ALCOHOLS. 1. Series $C_nH_{2n+2}O$ or $C_0H_{2n+1}OH$. Of this

series the following members are at present known, each being derived from the corresponding paraffin CaH20+2 by substitution of OH for H:

Methyl Alcohol	· CH,OH
Traberi	C,H,OH
Danamal	C.HOH
Butyl ,	C,H,.OH
Amyl ,,	$\mathbf{C}_{\mathbf{x}}\mathbf{\Pi}_{\mathbf{H}}$.OH
Hexyl ,	· C ₆ H ₁₃ .OH
Heptyl "	· C,H,sOH
Octyl "	$O_8H_{17}OH$
Ennyl or Nonyl Alcohol .	· CallinOH
Deoyl Alcohol	$\cdot C_{10}H_{21}.OH$
Hendecyl "	• C ₁₁ H ₂₃ .OH
Dodseyl "	• C ₁₂ H ₂₅ .OH
Tetradecyl ,,	· CHIL.OII
Hexadecyl or Cetyl Alcohol	. C ₁₆ II ₃₃ .OII
Octadecyl Alcohol	. C ₁₈ H ₃₇ .OH
Ceryl Alcohol	. C ₂₇ H ₅₅ .OH
Melissyl or Myricyl Alcohol	. CaoHen.OH

The first and second of these alcohols do not admit of isomerie modifications: for supposing, as is most prohable, that all the hydrogenatoms in the paraffins methane CH, and ethane CH. CH, have the same value and are attached to their respective carbon-atoms in the same way, the result of the substitution of OH fer II in them must be the same, whichever of the hydrogen-atoms is thus replaced. But in all the higher terms of the series the cass is different. Thus in propane, CH₃.CH₂.CH₃, the substitution may take place either in one of the exterior groups CH3, or in the middle group CH2, giving rise to two alcohols of different structure, distinguished as primary and secondary, viz.

CH, OH, CH, OH . CH, CHOH, CH, Primary Secondary

In the primary alcohols the carbon atom id; 40, 20 hydroxyl is connected immodiately with alcohol nother carbon atom, that namely called delonating; but in the secondary alcohol (Zeise, loc. cit.).—41:0 other carbon atoms, and

these are the only forms of a 8-carbon alcohol of the series.

The 4-carbon alcohol of the series admits of. a greater number of modifications. For in the first place, the hydrocarbon, butane, O.H., from which, it is derived, is itself susceptible of two forms, viz., Normal butane OH3.CH2.CH2.CH3. and Isobutane CH2. CH(CH3)2; and further the first of these hydrocarbons is capable of yielding one primary and one secondary alcohol-these terms having the meanings above explainedwhile the second yields another primary alcohol, and likewise a tortiary alcohol, in which the C-atom joined to the hydroxyl is linked also to three other atoms of earbon. These four derivatives are represented by the following formulæ :-

Normal Primary CH3.OH2.CH2.CH2OH. Isoprimary (CH₃), CH.CH₂OH. Secondary CH₃.CH(OH).CH₂(CH₃). Tertiary (CH₃)₂C(OH).CH₃.

The higher alcohols of the series admit of a still larger number of isomerio modifications; but all these alcohols must be either primary, secondary, or tertiary; for the C-atom joined to the OH cannot be joined to a number of other carbon-atoms greater than three. In other words the replacement of an H-atom by the group OH must take place, either in a methylresidue CH3, a methylene-residue CH2, or a methenyl residue CH, producing respectively a primary, secondary, or tertiary, alcohol.

A very convenient nomenclature for these isomeric alcohols has been introduced by Kolbe (A. 132, 102). Mcthyl alcohol is called carbinol and the higher alcohols named as its substitution.

products, thus: Carbinol or Methyl Alcohol CH₁OH. Methyl-carbinol or Ethyl Alcohol MeCH2OH. Ethyl-carbinol or Propyl Alcohol EtCH,OH. Dimethyl-carbinol or Isopropyl Alcehol

Me, CHOH. Propyl-carbinol or Butyl Alcohol PrCH2OH. Isopropyl-carbinol or Isobutyl Alcohol PrCH2OH. Methyl-cthyl-carbinol or Secondary Butyl Alcohol MeEtCHOH

Trimethyl-carbinol or Tertiary Butyl Alcohol

Primary, secondary, and tertiary alcohols are distinguished from one another by their products of oxidation. The primary alcohols of the series $C_nH_{2n+2}O$, containing the group CH,OH, are converted by exidation with chromio acid mixture, first into the corresponding aldehydes, by removal of H₂, or conversion of CH₂OH into CHO, and then by further oxidation into fatty acids C, H, O2; thus:

 CH_3 . CH_2 . CH_2 OII + O = H_2O + CH_3 . CII_2 .CHOPropyl Alcohol Propionic Aldehyde CII_3 . CH_2 . $CII_2OH + O_2 = H_2O + CII_3$. CH_2 .COOHPropionic sold

A secondary aloohol on the other hand which coutains two alcohol-radioles united by Clioh, is converted, by removal of H, from this group, into a ketone, i.e. a compound consisting of two alcohol-radieles united by the group CO.

 CH_3 .CHOH.CH₂ + O = H_2O + CH_3 .CO.QH₃ Secondary Propyi Alcohoi Conversely the aldehydes treated with nascent

hydrogen (action of sodium-amalgam) are converted into primary alcohols, and the ketones

into secondary alcohols.

Tertiary alcohels do not yield by oxida-tion either aldohydos or ketones, or acids containing the same number of carbon-atoms as themselves, but are split up into bodics containing smaller numbers of carbon-atoms-tertiury butyl alcohol for example into formic and propionic acids:

 $(CH_3)_3COII + O_4 = CH_2O_2 + C_3H_6O_2 + H_2O.$ The three classes of alcohols may also be distinguished by the following test: -A quantity of dry silver nitrite, mixed with an equal weight of dry sand, is introduced into a small distillation-flask fitted with a side-tube; the iodide of the alcohol under examination is then added; the mixture, after the reaction has begun, is distilled; and the distillate, received in a testtubo, is shaken up with potassium nitrite and potash-ley, and then aciduluted with dilute sulphuric ucid. If no coloration of the mass ensues, the alcohol-radicle present is a tertiary, whereas a red coloration indicates the presence of a primary, and a blue coloration that of a secondary, radiolc. The reaction may be re-cognised with great distinctness with the use of not more than 0.3 to 0.5 grm. of the alcoholic iodide (Meyer a. Locher, B. 7, 1510). Secondary hexyl iodidc does not give this test.

2. Series Callano. The most important member of this series is ally! alcohol C3H8O, which is a primary alcohol, convertible by exidation into acrylic aldchyde C₂H₄O, and acrylic acid C₂H₄O₂. They are unsaturated compounds capable of taking up 2 at. bromine, and forming

the compounds C.H., Br.O.

3. Series C_nH_{2,-2}O. This series includes propargyl alcohol, di-allyl-carbinol, and the higher

homologues of the latter.

4. Series C_nH_{2n-8}O. These alcohols are derived from the aromatic hydrocarbons, C_nH_{2n-6}, in the same manner as the fatty alcohols C_nH_{2n+2}O from the parafins. The lowest member, viz., phenol C₀H₀O or C₀H₀(OH), which may be formed from benzene, C₀H₀, by oxidation with H₂O₂ or with nascent ozone (Leeds, B. 14, 96), is the only alcohol of the series containing 6 at. carbon. The higher terms admit of isomeric modifications: for all the homologues of benzeno may be regarded as derived from benzene by substitution of one or more of its hydrogen atoms by alcohol radicles CaH_{2n+1}, and the formation of an alcohol from such a hydrocarbon by substitution of OH for H may take place either in the benzene nucleus or in one of the substituting alcohol-radicles: thus from toluene C.H. CII, may be obtained the two alcohols, C.H. CH.OH (benzyl alcohol) and C.H. (OH). CH. (cresol), und the higher hydrocarbons of the series are capable of yielding a still greater number of metameric alcohols. The properties of the compounds thus formed differ considerably, according as the hydroxyl is introduced into the benzene nucleus, or into one of the associated alkyls. The compounds formed in the latter case—benzyl alcohol for example are true alcohols analogous in all their reactions to those of the fatty series; But those in which the OH replaces a hydrogen-atom in the benzene nucleus (phenols) exhibit very different

properties, the hydroxyl being much less easily displaced by other radioles (Cl. Br. &c.), v. PHENOLS).

5. Series C.H. O. To this series belong oinnamy I alcohol C.H. O, cholesterin C.z.H. O, and allyl-phenol C.H. O.

DIHYDRIC ALCOHOLS.

These alcohols are derived from hydrocarbons by substitution of two IIO-groups for two H-atoms, and may therefore be regarded as compounds of divalent alkyls with hydroxyl. Two series of them are known, viz., glycols derived from the fatty hydrocarbons, and dihydric phenols from the aromatic hydrocarbons.

H. W The lower glycols are described as GLYCOL. PROPYLENE OLYCOL, and tri-METHYLENE OLYCOL, but the higher members as di-Oxy-BUTANE,-PEN-TANE, &c. Unsaturated glycols are described as di-Oxy-Butinene, -HEXINENE, and -HEPTINENE. The chief di-hydric phenols are Pyro-catronin, Resoacin, and Hydaoquinone. Di-Oxy-naptha-Lene and di-Oxy-anthaacene belong to this class.

Tainydrio Alconols.

This class is represented by five fatty alcohols: GLYCEAIN, and tri-OXY-BUTANE, -PENTANE, -HEX-ANE, and -HEXINENE. There are also several aromatic representatives, e.g. Pyrooallol, Philo-BOOLUCIN, and tri-OXY-NAPTHALENE.

TETRALITOR C ALCOHOLS.

Erythrite is the only fatty tetra-hydric alcohol Tetra-oxy-benzene and tetra-oxytetra-phenyl-cthano, and tetra-oxy-tri. phenyl-methano are aromatic tetra-hydric alcohols.

PENTAUYDRIO ALCOHOLS.

Pinite and quercito are the only ones known.

HEXAUYDAIC ALCOHOLS.

Mannite, dulcite, sorbite, perseite and hexa-oxy-diphenyl make a complete list. Formation of Alcohols.—1. From haloid cthers as described above. - 2. From aldehydes or ketones by reducing with sodium-amalgam.-3. From acid anhydrides by reduction with sodium-amalganı (Linnemann).-4. From primary amines by the action of nitrous acid; this reaction is, however, accompanied by an intramolecular change in the case of all fatty amines except ethylumine and methylamine. As a result of this change n-propylamino gives rise to secondary as well as n-propyl alcohol.-5. Secondary Alcohols may be got by the action of zino alkyls upon aldeliydes:

RCHO + ZnEt, -- R.CHEt.OZnEt R.CHEt.OZnE+ H.O =

R.CHEt.OH + EtH + ZnO 6. Tertiary alcohols can be formed, similarly, from zinc alkyls and scid chlorides: CH₃.CO.Cl + ZnMe₂ = CH₂.CClMe.OZnMe

> CH, CClMe.OZnMe + ZnMe CH, CMe, OZnMe + ClZnMe

 CH_3 . CMo_2 . $OZnMe + H_2O =$

CH, CMe,OII + ZnO + CH, v. ZING-METHYL. -7. From olofines, by dissolving them in H,SO, and distilling the product with water: $Me_2C.O.SO_4H + H_2SO_4 = Me_3C.O.SO_4H$ $Me_3C.O.SO_4H + H_3O = Me_3C.OH + H_3SO_4$

Just as: O.H. + H.SO. = O.H.SO. H O.H. HSO. + H.O = O.H.OH + H.SO.

By this reaction primary alcohols can be turned into secondary. Thus cone. H2SO, converts propyl alcohol into propylene, which is converted by the above treatment into iso-propyl alcohol.

Reactions of Alcehols.—Besides the general reactions mentioned above, the following are important:—1. Any reaction that might be expected to produce an alcohol of the form R.CH:CH.OH, produces an aldehyde, R.CH., CHO, instead (Erlenmeyer, B. 13, 309; 14, 320). Similarly an alcohol of the form R.CH:CR'.OH bocomes a ketone, R.CH., CR'O .- 2. On heating methyl, ethyl, butyl, octyl, and capryl alcohols with ammoniacal ZnCl, at 240°-280° a mixture of the mono-, di- and tri-alkylamines is got, the yield of which amounts to 50-75 p.c. of the alcohol (Merz a. Gasiorowski, B. 17, 623).—3. Tertiary alcohols differ from primary and secondary alcohols in not combining with baryta (Menschutkin, J. R. 10, 368).-4. Conc. HNO, converts tertiary alcohols into nitro-alkylenes, thus (CH_n)₄COH becomes nitro-iso-butylene C₄H₁NO₂ (Bn. 1, 232).—5. The boiling-points of tertiary alcohols are lower than those of the isomerie secondary alcohols, and these again lower than those of the isomeric primary alcohols.—6. The alcohols C, H2, 2 O are decomposed by sine dust at 300°-350° into olefine, C.H., and water. Methyl alcohol gives, however, CO and hydrogen (Jahn, M. 1, 378) .- 7. Carbonic oxide above 100° acts upon sodium alcoholates (RONa) mixed with sodium salts $(C_nH_{a_{n-1}}O_nNa)$ as follows: $RONa + CO + C_nH_{a_{n-1}}O_nNa =$ $CHNaO_2 + C_nH_{a_{n-2}}RO_nNa$

the elements of NaOH being abstracted so that R displaces H. A secondary reaction is

RONa + CO = RCO.Na(Genther a. Froehlich; Looss; Poetsch, A. 218, 56). But CO does not act on a mixture of sodie phenylate and sodic acctate at 200° (Schroeder, A. 221, 35), or on one of sodic sthylate and sodic benzoate at 200°. On a mixture of sodio cthylate and sodic phenylacetate CO forms various acids including ono (810°-320°) which may be phenyl-vinyl-butenyl acetio acid, Ph.C(C,H₂)(C,H,Et)CO,H. On mixture of NaOEt and sodic cinnamate, carbonic oxide forms di-ethyl-cinnamic acid, C.H. CEt:CEt.CO.II and di-butyl-cinnamic acid, C.H., C(C,H.,):C(C,H.,).CO,II. Both are oils.—8. Primary alcohols heated with soda-lime form acids and give off hydrogen thus: RCH_OH + KOH = RCO, K + 2H, (Dunias a. Stas, A. 35, 129). But at a higher temperature a second reaction occurs: $RCO_2K + KOH = RH + CO_2K_2$. If the hydrogen evolved be measured, some conclusion may be drawn as to the molecular weight of the alcohol; but the lower alcohols cannot give good results, as the hydrocarbons RH are gases. Myricyl alcohol gives off 17 of the calculated hydrogen (C. Hell, A. 223, 269).— 9. When ah alkyl carbonate is heated with an alcohol, exchange of radicles occurs if the radiclo of the alcohol contains more carbon atoms than that of the ether (Röse, A. 205, *240). But when an alcohol is heated with an acetal, exchange takes place only if the alcohol has the smaller radicle (Geuther, A. 218, 45). When an alcohol is boiled with a simple

ether or with an ether of acetic or butyris acid with inverted condenser, no change occurs (G.) .- 10. If a small quantity of a secondary alcohol, other than isopropyl alcohol, ba moistened with HNO, and then mixed with water and shaken with ether, on adding alcoholio KOH to the residue left after evaporating the ether, yellow prisms of a potassium alkyl nitrite separate (Chancel, C. R. 100, 601). -11. Benzoin, isohydrobenzoin, and pyrocatechin givo, when their sodium compounds are treated with ClCO, Et, neutral carbonates of the form R"CO, while resorcin, hydroquinone, and orcin give di-carbonates, R"(CO₃Et)₂ (M. Wallach, A. 226, 87).—12. On the Rate of Etherification of alcohols v. CHEMICAL CHANGE .- 13. Fe,Cl, gives a colour-reaction with all oxy-compounds whether aromatic or fatty, though in the latter case the reaction is faint and a nearly colourless solution of the reagent is required. Such a solution can be prepared by diluting two drops of 10 p.c. solution of Fe₂Cl₈ with 60 o.c. of water. If an excess of the substance to be tested is added to this solution a sulphur-vellow colour will be produced if a fatty alcohol, oxyacid, or carbolydrate is present (Landwehr, B. 19, 2726)

ALDANE. A term proposed by Riban (C. R. 75, 98) to designate products formed by the union of two or more molecules of an aldehyde, with elimination of water-e.g. crotonic aldehyde CII3.CH:CH.CHO from aldeliyde.

Di-aldane C,H,O, i.e. CH,CH(OH).CH,CH:CH.CH(OH).CH,CHO [130°]. S. (ether) 87 at 22°. Formed by the condensation of aldol, CH, CH(OH).CH, CHO, under the influence of hydrochloric acid (Wurtz, Bl. [2] 24, 100; 28, 169). Crystalised from water. Sl. sol. cold water, v. e. sol. boiling alcohol. May be distilled in vacuo. It reduces silver solution. Aqueous NH, at 100° forms a crystalline base, $C_{16}H_{28}N_2O_3$, v. sol. water, alcohol or ether (Wirtz, C. R. 91, 1030). The aqueous solution of the baso deposits, after some time, an amorphous isomeride.

Iso-di-aldane C_nII₁O₂ [114°]. Formed by heating aldol at 125° (W.) or by slow action of aqueous HCN upon aldol (Lobry de Bruyn, Bl. [2] 42, 161).

Di-aldanic acid C,H,O, i.e. CH,CH(OH).CH,CH:CH.CH(OH).CH,CO,H [80°]. (198°) at 20 min. Formed by treating an aqueous solution of di-aldane with Ag.O or KMnO, (Wartz, C.R. 83, 255, 1259). Monoclinic crystals. V.e.sol. alcohol or water, m. sol. ether. Salts:-KA' deliquescent crystals (from 98 p.c. alcohol).-NaA': plates (from alcohol).-BaA' ppd. as powder by adding ether to an alcoholio solution.-CaA', xaq: v. e. sol. water, but not deliquescent.—AgA': small laminæ (from boiling water); insol. alcohol.

Di-aldanic alcohol CaH, O. i.e. CH, CH(OH).CH, CII:CH.CII(OII).CH, CH, OH [49°-53°]. (162°-165°) at 10 mm. Prepared by reducing di-aldane in aqueous solution with a large excess of (I p.o.) sodium amalgam, the liquid being kept slightly acid with HCl. The liquid is neutralised and evaporated, freed from NaCl by alcohol, and the alcoholic solution distilled (Wurtz, C. R. 92, 1871). White, crystalline, deliquescent mass. V. c. sol. water and alcohol; v. sol. ether. Ac.O forms an acetate $\Omega_1 H_{11} Ac_2 O_2$ (c. 159°) at 20 mm. Di-aldanic alcohol does not reduce ammoniacal AgNO₂.

Constitution.—The formation of a di-acetate seems inconsistent with the constitution assigned to di-aldanic alcohol. Di-aldane may be conaidered to be derived from di-aldol.

OH, OH(OH). OH, OH(OH). OH, CH(OH). CH, CHO, by removal of H,O. If this delaydration is to destroy two hydroxyls an anhydride must be formed:

CH,.CH.CH,.CH.CH,.CH(011).CH,.CHO

and the formulæ of dialdanc derivatives must be altered accordingly.

ALDEHYDE (AČĒTIC) C.H.O, i.e. CH. CHO. Mol. w. 44. (21°). S.G. 2 800 (Kopp, A. 64, 214); 3 7799 (Brühl); 1 7951; 1 67876 (Perkin, C. J. 45, 475). S.V. 56·6 (Ramsay). μ_{β} 1°3359. R. $_{\infty}$ 18·18 (B.). H.F.p. 48,740 (Thomsen); 46,000 (Berthelot). H.F.v. 47,870. M.M. 2°383 at 16·3°. V.D. 1·532 (for 1·520).

Occurrence.—In the first portions obtained by rectifying spirit that has been filtered through charcoal, where it is perhaps formed by oxidation in the charcoal (Krämer a. Pinner, B. 2, 403; 4, 787; Kekulé, B. 4, 718). The name aldehyde was invented by Liebig as a contrac-

tion of alcohol dehydrogenatum.

Formation. - 1. In the oxidation of alcohol, either by slow combustion in centact with platinum black, or by the action of CrO₃, chlorine water, HNO, or a mixture of H2SO, and MuO2 (Liehig, A. 14, 133). Also by exidation of acetic ether and other ethyl compounds (e.g. ethylamine, Carstanjen, J. pr. 89, 486), and by slow combustion of ether.—2. By action of ZuCl₂ on glycol: C₂H₄(OH)₂ = H₂O + C₂H₄O (Wurtz, A. 108, 86).—3. From ethylene bronide and water at 160° (Carius, A. 131, 172) C₂H₄Br₂ + H₂O = C₂H₄O + 2HBr. From ethylene bromide and mereurie acetate (Linncmann, A. 143, 347).-4. From ethylene and CO at 400° (Schützenberger, Bl. 31, 482) .- 5. From acetylene and aqueous HgBr, (Kutscheroff, B. 14, 1510) .-6. By electrolysis of potassic lactate (Kolbe, A. 113, 244), of sugar solutions (II. T. Brown, C. J. 25, 578), or of alcohol containing H,SO, or KOH (Jaillard, C. R. 58, 1203).—7. By dissolving acetylene in dilute H₂SO₄ (S.G. 1-35) and distilling the product with water (Lagermark a. Eltekoff, B. 10, 637).—8. By the dry distillation of a mixture of calcie acetate and calcie formate $Ca(CO_2CII_3)_2 + Ca(CO_2II)_2 = 2CaCO_3 + 21ICO.CH_3$ (Ritter, A. 97, 369).—9. By oxidation of ethylene with aqueous CrO, at 120° (Berthelot, C. R. 68, 334) .- 10. By reducing chloral with zinc and dilute H₂SO₄ (Personne, C. R. 71, 227).—11. Together with formic acid by heating lactic acid with dilute H2SO, at 130°

CH₂CH(OH).CO₂H = CH₂CHÔ + HCO₂H (Erlenmeyer, Z. [2] 4, 343). Also, together with lactide, by the dry distillation of lactic acid.—12. Among the products of the action of H₂SO₄ and MnO₂ or K₂Cr₂O₇ upon albumen, fibrin, casein, gelatin (Guekelberger, A. 64, 46, 86), on gluten (Keller, A. 72, 31).—13. By heating acetal with glacial acetic acid at 180° for two days (Beilstein, C. R. 48, 1121).—14. By passing alcohol through a red-hot tube. From from oil in the same way (Hess, P. 38, 380).

Aldehyde occurs among the products of the dry distillation of wood (Kane, A. 19, 288; Krämer, a. Grodzki, B. 9, 1921; Mabery, Am. 5, 256), and of sugar (Völckol, A. 87, 303).—15. By distilling a-di-alkylated-\$-oxy-propionio acids:

CH₂,CH(OII).CEt₂,CO₂H =

CH₃.CH(OH).CEt₂.CO₂H = CH₃.CHO + CHEt₂CO₂H.

Preparation.—1. From the 'first runnings' in the rectification of fermented liquors.—2. Alcohol (3 pts. of S.G. '842) and K.Cr.O. (3 pts.) are placed in a retort and conc. H.SO. (4 pts.) is slowly run in. The heat evolved causes the aldehyde to distil off (W. a. R. Rodgers, J. pr. 40, 240). It is collected in dry ether, which is afterwards saturated with dry NH₃; aldehyde animonia separates in cubes, and thie is distilled with H.SO. (3 pts.) mixed with water (4 pts.), the receiver being cooled with ice and salt. The product is dried over CaCl₂ and rectified.

Properties.—Characteristic odour, miscible with water, but separated by CaCl₂ from sclution. Neutral. Readily polymerised. Mixes with alcohol and ether. A mixture of aldehyde (1 pt.) and water (3 pts.) boils at 37°. Aldehyde dissolves S, P, and 1. Burns with blue flame. It dissolves 7 times as much SO₂ as water does.

Tests.-1. Heated with annuoniacal silver nitrate forms a mirror .- 2. Heated with aqueous potash forms a yellow body (aldehyde resin) and gives off a characteristic odour (Weidenbusch, A. 66, 153). The solution then contains formate and acetate. - 3. Restores the colour to a solution of a rosaniline salt that has been bleached by SO2-4. Reacts with hydroxylamine forming a liquid oxim (v. infra) .- 5. Reacts with phenylhydrazine forming a crystalline phenyl-hydrazide (v. infra).—6. Combines with NaHSO, -7. Combines with NH, -8. H,S passed into an aquecue sclution forms an oil, converted by acids into solid thio-aldehyde (q. v.). -9. Alkaline aqueous solutions produce a red coloration when treated with diazo-benzene eulphonic acid and a little sodium-amalgam.

Reactions. - 1. Oxidised to acetic acid slowly by air, more rapidly in presence of platinum black, most rapidly by oxidising agents.—2. Passed over red-hot soda lime, it forms sodie acetate and hydrogen. Passed over red-hot quieklime, it gives acetone and various ketones and gases (Schloemilch, Z. [2] 5, 336) .- 3. HI at high temperatures reduces it to ethane (Berthelot, Bl. [2] 7, 59) .- 4. Sodium amalgam reduces it to alcohol, some di-oxy-butane being also formed (Kekulé, A. 162, 309) .- 5. Converted into crotonic aldehyde (q. v.) by ZnCl, or by squeons solutions of sodie acctate or Rochelle salt at 100°. Zinc shavings at 100° produce an aldehyde C_oII₁₀O₂ (220°); it is an oil and combines with NaHSO₃ (Riban, Bl. [2] 18, 63).—6. With chlorine forms, in sunlight, acetyl chloride (Wurtz, A. 102, 93).—7. Chlorine passed into aqueoue aldehyde forms chloral, bntyfo-chloral, dichloraldehyde, and other bodies (Pinner, A. 179, 21; B. 8, 1321, 1561; Wurtz a. Vogt, Bl. 17, 402). Bromine converts aldehyde, dissolved in acetic ether, into bromal and disharm addehyde. di-bromo-aldehyde.-8. PCl, gives ethylidene chloride, CH₂,CHCl₂ (Beilstein, A. 118, 110). COCl₂ acts similarly (Eckenroth, B. 18, 518). PCl₃Br₂ gives ethylidene bromide, CH₃.CHBr (Paterno a. Pisati, B. 5, 289).—10. Dry HCT

passed into cold aldelyde forms ethylidone chlor-hydrin UH₃.CHOl.OH, (25°) at 40 mm. This changes epontaneously, or more quickly if heated or treated with HCl, into 'ethylidene oxy-chloride,' C4H6Cl2O, (0. 59°) at 40 mm. A small quantity of another body, CaH12Cl2O, (c. 100°) at 40 mm., is also got. Ethylidene oxy-chloride is probably di-ohloro-di-ethyl oxide (CK3.CHCl)2O. It is converted by boiling water into aldehyde and HCl, and by ammonia into efflorescent needles of (CH, CHNH₂)₂O 2HCl (Lieben, C. R. 46, 662; Kessel, A. 175, 46; Hanriot, C. R. 92, 302). Aldehyde saturated with IICl is converted into crotonic aldehyde (q. v.), chloro-butyrie aldehyde, and a compound $C_{10}H_{18}Cl_{2}O_{3}$ [98°] (Kekulé, A. 162, 102).—11. Aldehyde left for some days with aqueous HCl forms aldol (q. v.) .- 12. HCl passed into a mixture of aldehydo and alcohol forms chloro-ethylether (q, v_1) .—13. HCl passed into a mixture of aldeliyde and mercaptun forms di-thioacetalCH3.CH(SEt), a mobile liquid (Baumann, B. 18,884).—14. Aldehyde forms with zinc ethyl a compound which is decomposed by water with production of secondary butyl alcohol (q. v.).— 15. When paraldehyde (I g.) is added to cold $\mathbf{H_2SO_4}$ (100 g.) and the solution is shaken with benzene, di phenyl ethane is got :

CH₃-CHO + 2C_nH₃ = CH₃-CH(C_nH₃)₂ + H₂O (Baeyer, B. 7, 1190).—16. With cyanamide it forms a compound (C_nH₃)₃N₃Cy₃Aq (Knop, A. 131, 253).—17. With HCN it gives lacto-nitrile (q, v).

18. With HCN, HCl, and NH, in aqueous solution it gives having (q, v). solution it gives, on boiling, alanine (q. v.). A mixture of aldehyde-ammonia and HCN in 30 p.c. solution acidified by HCl gives in the cold amido-propionitrile, which changes first to imido-propionitrile, and then, in about a month,

to hydrocyanaldine.

Hydrocyanaldine C, II, N, [115°]. S. 18 at S. (alcohol) 1.27 at 18°. Prisms (from ether). May be sublimed. V. sol. acctone, m. sol. ether, v. el. sol. CS. Decomposed into its components by boiling AgNO, or boiling KOII.

Parahydrocyanaldine C,H,N, [232°]. 8. 01 at 20°; S. (alcohol) 01 at 18°. This is a similar body formed by allowing the liquid containing hydrocyanaldine to stand several months, and also by warming a mixture of amido and imido- propionitrile with HCl. Rhombic crystals (from acetone). Insol. ether, v. e. sol. acetone. Decomposed by AgNO, or KOII like hydrocyanaldine.

Combinations.-1. With bisulphites of the alkalis: C2H,ONaHSO3Aq: pearly plates by evaporation over H SO,; sating needles when ppd. by alcohol. - C. II OKIISO, : hard indistinct crystals composed of minute needles .-(C₂H₄O)₂Ba(HSO₃)₂e₃, soluble seales.—If a solution of (NH₄)HSO₃ be mixed with aldelyde and evaporated, it deposits crystals of C.H.OSO.NH., S. 16 at 16° (Bunte, A. 170, 305). But by assing GO, into alcoholio aldehyde-ammonia, Bedtenbacher (A. 65, 37) got unstable needles of an isomerit body, S. 70 at 16°. When strongly heated with potash lime, this decomposed with production of di-methyl-amine or ethylamine (Gössmann, A. 91, 122).-It may be C2H4ONH4HSO4 (Beilstein) .- The compound of aldehyde with acid sodium sulphite may perhaps be represented by the formula CH, CH(OH) SO, Na, as a oxyesthyl sodium sulphite.

2. With ammonia: CH, CH(OII).NH, Aldshyde-ammonia [70° 80°]. (100°). V.D. 80°36. By passing NH, inthe ethereal solution of aldehyde (Liebig, A. 14, 133). Rhombohedra, best got by mixing a cono. alcoholio solution with ether, very soluble in water, hardly soluble in ethor. Alkaline. Turns brown in air. Decomposed by diluto acids, even by CO, giving off aldehyde. Reactions.—(a) H₂S forms thialdine (q. v.).—(b) H₂Se forms selen-aldine C₂H₁₃NSe.—(c) Alcoholio CS₂ forms carbo-thialdine (q. v.).—(d) HCN and HCl form, in the cold, hydrocyanaldine, C_sII_sN_s, or on heating, alanino (q. v.).—(e) At 120° in a scaled tubo it forms tri-methyl pyridine, oxy-tetraldine C. H., NO, and oxypentaldine C. HI, NO. The two latter are monacid amorphous bases, sl. sol. water (Babo, J. pr. 72, 88; Heintz a. Wislieenus, J. 1858, 317; Schiff, A. Suppl. 6, 10).—(f) Aotion of SO is described above.—(g) CS, forms earbothialdine C,II, N.S. (q. v.).—(h) NaOEt and MeI in the cold form isocholino iodide, C.H., NOI (G. Meyer, B. 16, 207).

3. With ammonia and silver nitrate or sulphate: When AgNO, (100 c.e. of a 3 normal solution) is added to aqueons NII, (15 o.c. four times normal) and, after filtration, aldehyde is added as long as the pp. first formed redissolves, a liquid is got in which more NII, (15 c.c.) eauscs separation of the compound Cillin NaOa Ag laq. which must be washed with alcohol and ether and dried at a low temperature (Reychler, B. 17, 41). It forms unstable white six-sided plates. Sl. sol. water, v. sl. sol. alcohol, insol. ether. Its warm aqueous solution deposits a eilver mirror.-If the same solutions be mixed in the following proportions: 20 e.e. NII, Aq, 33 c.o. AgNO,Aq and 20 e.e. aldehyde, and 250 c.o. alcohol be added, a white microcrystalline pp. $C_iH_{in}N_iO_iAg$ is got. This body is represented by Liebermann a. Goldschmidt (B. 10, 2179; II, 1198) as AgNO,2C.H.NH;

Reychler writes AgO.N (NH CII.CH,)2. $\begin{array}{l} {\rm Ag.SO_4(C.H._1NH)._3aq.-Ag.SO_4(C.H._1NH)._6aq.-}\\ {\rm Ag.SO_4(C.H._1NH)._3NH._3aq.(W.~G.~Mixter,}\\ Am.~\ref{S_\bullet} \end{array}$ [3] 17, 127).

4. A solution of aldehyde in alcoholic ammonia in six months becomes brown. If it be then evaporated, tri-cthyllidene diamine, or hydracetamile (CH₂CII)₃N₂, is left as a yellow amore hous powder, soluble in water. Its hydrochloride is CaH12N2HCl. Boiling water or acids convert it into oxy-trialdine C. HuNO, an amorphous yellow powder, soluble in water; ealts.—C.H., NO.HCl.—(C.H., NO).H., SO. (H. Schiff, Bl. [2] 8, 443; A. Suppl. 6, 1).

5. With pressive acid: C.M.O.N.H. i. e.
C.H.(OH)CN, ethylidene cyanhydrin or
lacto-nitrile (q. v.).—6. With ethyl nilrate:
C.H.(O2ENO) (86°). S.G. 12 1045. Formed by distilling a mixture of KEtSO, with KNO, is an oil. Vapour explosive. Decomposed by potash into aldehyde (Nadlor, A. 116, 173).

7. With ethyl chloride v. Chloro- n-ETHYL

8. With alkoyl chlorides or bromides. The following compounds may be viewed as dorived from ethylidone glycolchlorhydrinCH, CH (OH) CI (v. reaction 10), by displacement of H by acid radioles.

(a) With acetylchloride: C.H.O. C.H.O.Clor | C.H. (OA) Cl, ethylidene chloracetin (121.5° cor.) S.G. 12 1.114. Combination takes place at 100° (M. Simpson, Pr. 27, 120; Franchimont, R. 1, 246; Rübencamp, A. 225, 274). The compound was discovered by Wurtz (Z. 1871, 362; A. Ch. [3], 49, 58; C. R. 73, 528). Decomposed by potash into KCl, acetic acid, and aldehydc. KOAe forms CH, CH(OAe), (Schiff, B. 9, 304). Chlorine at 120°, in presence of iodine, forms CHCl., CHCl.OAe, tri-ehloro-ethyl acetate (250°-280°). Bromine dropped into it at 100° forms bromethyl bromo-acetate CH2.CHBr.O.CO.CII2Br (v. Bromo-acetic acid).

(b) With acetyl bromide forms a corrosponding, but unstable, compound (0.140°) (Tawildaroff, A. 176, 21).

(e) With propional chloride: C₃H₉ClO₂ or CH₂ClICl.O(C₃H₃O), ehloro-ethyl-propionate (135° uncor.). S.G. ¹⁵ 1·071.

(d) With butyrylchloride: CH4.CHCl.O(C4H7O)

(149° uncor.). S.G. 15 1.038.

(e) With valerylehloride; CH2.CIICl.O(C5II2O) (c. 163°). S.G. 15 .997.

DERIVATIVES OF ORTHO-ALDEHYDE.

The following combinations between aldehydo and compounds of the form M2O may bo viewed as derivatives of ortho-aldehyde, CH3.CH(OH)2. Ortho-aldchyde itself is not known, but chloral hydrate is tri-chloro-orthoaldeliyde.

Alkyl derivatives, Acetals or Aldehydates. The term 'acetal,' originally applied to CII, CH(OEt),, is now often extended to the whole series of di-alkylated ortho-aldehydes.

These bodies are formed, together with other products, by the exidation of alcohols. Each of them may be formed from one of its higher homologues, by heating the latter at 120°, with an alcohol containing a lower radicle. Thus di-ethyl-acetal heated with methyl alcohol yields dimethylacetal, whereas the latter heated with ethyl, propyl, isobutyl, or amyl alcohol yields only traces of an acetal containing different alcohol-radicles. Similarly diethyl-acetal heated with methyl-alcohol is converted, for the most part, into dimethylacetal, but is practically unaltered by propyl and amyl alcohols (Bachmann, A. 218, 38). Aldehydates may also be formed by heating aldehydes with alcohols and HCl (Wurtz a. Frapolli, A. 108, 226; Claus a. Trainer, B. 19, 3004).

Ethyl-ortho-aldehyde CH3.CH(OH)(OEt) (80°-90°) (Renard, B. 8, 132) (c. 50°) (Jacobsen, B. 4, 215). Among the products of electrolysis of mixture of alcohol and diluto ILSO, (R.). By action of water on ohloro-ethyl-ether,

CH. CHCl.OEt (J.).

Di-methyl-acctalC,II,O2i.e.CII,CH(OMe)2 (62.7°-63.3°) at 751.6 nm. S.G. 22 .8655 (Bachmann); 8590 at 14° (Daneer, A. 132, 240). V.D. 8·10 (for 3·11). S.V. 110·81 (R. Schiff, A. 220, 104). Occurs in crudo wood spirit (D.). Formed by oxidising a mixture of MoOH and EtOH with MO, and H,SO, (Wnrtz). Prepared by heating aldshyde (4 vols.), methyl alcohol (8 vols.), and glacial HOAo (1 vol.) at 100° (Alsberg, J. 1864, 485). A colourless liquid burning with a white, blus-cdged finms (Wnrtz, A. Ch. [3] 48, 373). EtOH at 120° has hardly any action, traces of

methyl-sthyl-acetal being formed. Propyl, isobutyl, and iso amyl alcohols act similarly.

Methyl-ethyl-acetal CH, CH(OMe)(OEt).

Reactions that might be expected to produce this body yield only a mixture of di-methylacetal and di-ethyl-acetal (A. Genther, A. 225,

Di-ethyl acetal v. Acetal.

Methyl-propyl-acetal C_aH₁₁O₂ i.e. CH₃CH(OMo)(OPr) (103°-105°). Very little is formed from di-methyl-acetal and PrOH at 120°.

Ethyl-propylacetal C,H₁₀O₂ i.e. CH₂CH(OEt)(OPr)(124°-126°). Very little is formed from di-ethyl-acetal and PrOH at 120°.

Methyl-isobutyl-acetal C, II, O, (126°). Dimethyl-acetal (15 g.) heated with isobutyl alcohol at 120° forms a little (1 g.) of this body.

Methyl-iso-amyl-acetal C,H18O2 (141°·I44°). A little got from di-metliyl-acetal

and iso-amyl alcohol at 120°.

Di-propyl-acetal C₈H₁₈O₂ (146°-148°). S.G. 22'5 825. Got by passing pure PH₃ into a mixture of aldehyde and Proll at -21° (Girard, C. R. 91, 629).

Di-iso-butyl-acctal C10H22O2 (168°-170°). S.G. 22 816. Prepared like the preceding.

Di-iso-amyl-acetal C12H26O2 i. c. CH, CH(OC, H₁₁)₂ (e. 195°) (Bachmann); (210.8° eor.) (Alsberg, J. 1864, 485; Claus a. Trainer, B. 19, 3008). S.G. ²² ·801 (B.); ¹⁵ ·835 (A.). Aldebydo (1 vol.) and iso-amyl alcohol (5 vols.) are mixed, saturated with SO2 and warmed with glacial HOAc (1 vol.). A small quantity is got by heating amyl alcohol with acetal at 120°.

Ethylenc-acctal C,H,O, i. e.

CH₃CH<0>C₂H₄ (82·5°) at 766 mm. S.G. 21·002. 67. From aldeliyde and glycol at 100° (Wurtz, A. 120, 328). Separated by CaCl, from its aqueous solution. Not attacked by KOH. Acetic acid gives glycol di-acetin.

Propylene-acetal $C_5\Pi_{10}O_2$ i.e. CH₂CH<0>C₃H₆ (93°). From aldehyde and propylenc glycol at 160° (Gramout, Bl. 4I, 36I). Decomposed by water into aldeligde and propylene glycol.

Oxy-propylene-acetal C₅H₁₀O₂ i.c. CH₅CH_CO>C₅H₅OH (c. 186°) S.G. 2 1.081.

From aldehyde and glycerin at 180° (Harnitzky a. Menschntkin, A. 136, 126). Docom-

posed by water into its components. $Di \cdot (\beta) \cdot naphthyl \cdot acctal \quad C_{22}H_{18}O_2$ i.e. CH, CH(O.C, $_{13}I_{1})_{2}$. [201°]. Slowly formed when (β) -naphthol and aldehyde are dissolved in acctto acid and a few drops of HClarc added (Claisen, B. 19, 3318). Crystals; insol. aqueous alkalis. Changed by warming with HOAo and HCl into

ethylidene-di-naµhtliyl oxide, CH_s .CH $< C_{10}H_s$ > 0.

Alkoyl derivatives or ethylidenesalts. When both alkoyls (acid radicles) are the same these bodies may be viewed as compounds of acid anhydrides with aldehyde. They are slowly decomposed by water, more rapidly by potash, into aldehyde and acids. Mono-alkoyl derivatives, CH, CH(OH)(OR) are not known; they appear to split up into water and the an-

hydrides {CH, CH(OR)},O. These anhydrides may be formed from di-chloro-di-ethyl oxide, (CH,CHCl) O and sodium salts. They are volatile liquids, decomposed by water into aldehyde

and acid (Geuther, A. 226, 223).

Di-acetyl derivative CH₂.C11(OAc)₂ (168-4° cor.). S.G. ½ 1.073. μ = 1.10 at 28°. 1. From CH₂.CHCl(OAc) and AgOAc (Rübenoamp, A. 225, 274). -2. From aldehyde and Ac.O at 180° (Genther, A. 106, 249).—3. From ablehyde and AeCl at 100° (Franchiment, R. 1, 248).

Di-propionyl derivative CH₃.CH(OC₃H₅O), (192.2° cor.). S.G. 15 1.020 From CH, CHCl(OC, H,O) and $\mu = 1407.$

 $AgO(C_3H_5O)$.

Di-butyryl derivative CII, CH(OC, II,O)2 (215.5° cor.). S.G. $\frac{15}{2}$ 9855. $\mu = 1.411$ Di-valeryl derivative CH, CH(OC, H,O)2

(225° cor.). S.G. 25 ·947. μ=1·414.

Acetyl-propionyl derivative CH, CH(OAc)(OC, H,O) (178.6° cor.). S.G. 15.1.044 μ=1.402. From AgOC, H,O and CH, Cl1C!.OAe or from AgOAc and CH3.CHCl.OC3H3O (Geuther a. Rübencamp, A. 225, 281).

Acetyl-butyryl derivative CH, CH(OAe)(OC, H,O) (192.6° cor.). S.G. 15 1.015 μ=1.047. From AgOC, H,O and CH, CHCl.OAo, or from AgOAc and CH, CHCl.OC, H,O.

Acetyl-valeryl derivative CH, CH(OAc)(OC, H,O) (194°-199° cor.). S.G. 14 •991 $\mu = 1.408$. Similarly prepared.

POLYMERIDES OF ALDEHYDE.

Aldehyde reauty polymerises, forming aldol C.H.O. (q. v.), parallelnyde C.H.p.O., or metal-dehyde C_{2n}H_mO_n. Pure aldehyde may be kept without change, but when impure it spontaneously changes to paraldehyde or metaldehyde (Weidenbusch, A. 66, 155; Fehling, A. 27, 319; Geuther a. Cartmell, A. 112, 16; Lieben, A. Suppl. 1, 114; Kekulé a. Zincke, A. 142, 141; B. 3, 468). Meta dehyde is formed from aldehyde at a low temperature by the same reagents that eause the formation of paraldeliyde at high temperatures (K.a. Z.). Neither of these bodies is affected by hot potash, but both of them are converted by PCl, into ethylideue chloride, CH3.C11Cl2, and by HCl into 'ethylidene oxy-chloride' (v. supra). A little alcoholie KOH converts aldehyde into a mixture of metaldehyde, paraldehyde, and a little crotonic aldehyde (Perkin, C. J. 43, 88).

nyae (rerkin, C. J. 45, 85).

Paraldehyde C.11, O., Mol. w. 132. [10°-12°].

124°i. V.) (K. a. Z.); (124·4′) at 752mm.(R. Schiff, 4. 220, 104). S.G. 2° ·9943 (Brühl); ½5 ·9993;

4 ·9900 (Perkin, C. J. 45, 470). S.V. 150·7 (S.).

V.D. 4·35 (for 4·55, S.). μβ 1·4095. R∞ 52·48 (B.)

M.M. 6·662 at 17℃. S. 12 at 13°; 6 at 100°.

Preparation .- In presence of a small quantity of HCl, COCl2 or SO2, aldeliyde gradually becomes hot, often reaching 40°. It is then thanged to paraldehyde. H₂SO, and ZnCl₂ effect this change even more vigorously. product is cooled to "0", when paraldehyde erystallises.

Properties .- Colourless liquid, smelling like scetal and aldchyde. It is partially converted into all chyde by distillation. Distillation with H.SO., HCl, ZnCl₂, HgBr₂, or COCl₂ completely effects this change. The reactions of parallehyde, in presence of any of these bodies, are

therefore the same as those of aldehyde. It also forms CH, CH(OAo), with Ac.O. But it does not react with ammonia. HNO, oxidises it to glyoxal (Liubawin, J. R. 18, 496).

Constitution. — The S.V. agrees with that

required by Kekulé's formula $C_2H_4 < 0.C_2H_4 > 0$ (Brühl, A. 203, 44).

Metaldehyde $C_8H_{12}O_9$. S.(chloroform) 1034 at 26°; 4·235 at 60°; S. (benzene) 12 at 23°, ·181 at 80°. Formed by passing a few bubbles of SO2 or HCl into aldehyde in a freezing mixture; metaldehyde crystallises out, and the mother liquors are distilled and treated as before (K. a. Z.). CaCl2 effects the same transformation at the ordinary temperature.

Properties.-Long striated prisms, sublimes about 115° without melting. Insoluble in water, slightly soluble in cold alcohol or ether. It may be converted into aldeliyde: (a) by heating for a day in vacuo at 180°, (b) by repeatedly distilling under atmospheric pressure, (c) by heating its solution in ehloroform. The vapour density may be found in the usual way, due allowance being made for its partial dissociation, the amount of undecomposed metaldehyde being estimated after cooling. The V.D. is thus found to lie between 72.2 and 59.1, tho mean value being 62.5. Metaldehyde is not attacked in the cold by KMnO4, chromic mixture, or NH3. Chlorine forms ehloral; PCl, gives othylidene chloride (Hanriot a. Oeconomides, C. R. 93, 463; A. Ch. [5] 25, 226).

Di-aldehyde v. ALDOL

ALDEHÝDE-ACETAMIDE C.H.2N2O2 i.s. CH, CH(NHAe), di-acetyl-ethylidene diamine [169°]. Got by heating aldehydo with acetamide (Tawildaroff, B. 5, 477),

ALDEHYDE ACETATE v. p. 106, l. 6. ALDEHYDE ACETYL CHLORIDE v.p.105,1.1. ALDEHYDE ALCOHOLATE v. p. 105, l. 53. ALDEHYDE GREEN v. Rosaniline.

ALDEHYDE GUM C10H18O. The barium salt is formed by allowing a solution of aldehyde in baryta-water to stand for some time. From this salt H2SO4 liberates the 'gum' as a syrup, soluble in water and alcohol. It reddens rosaniline decolorised by SO2, and it gives iodoform with I and Na₂CO₃. Reduces hot Fehling's solution—Ca(C₁₀H₁₇O₄)₂: amorphous (Tollens, B. 17, 660).

ALDEHYDE PHENYL-HYDRAZIDE C,H,ON2 i.e. CH,CH:N2HPh.

Frem aldehyde and phenyl-hydrazine in ether v. Aldenydes, reaction 4. Crystallised from benzoline. Deliquescent. V. sol. alcohol or ether. Resolved by boiling water or dilute acids into its constituents.

ALDEHYDE RESIN. Formed by the action of aqueous or alcoholie potash, hot or cold, upon aldehyde, or by heating aldehyde with NaOAc in sealed tubes at 100°. It is accompanied by a strongly smelling yellow oil which may be removed by distillation. Aldehyde resin resembles colophony. It produces, when fused with potash, oxy-iso-phthalic acid [2832], o-oxy-m-toluic ccid [173°], and m-xylenol, C. II, Me, OH When strongly heated with zine dust it gives ethyl-benzene, m. and p. methyl-ethyl-benzene and methyl-naphthalene. Conc. HNO, given iso phinalis asid (Weidenbusch, A. 66, 158; Ciamician, M. 1, 199).

ALDEHYDES:—An aldehyde is a body de-

ALDENTDES.—An aldehyde is a body derived from a primary alcohol by removal of two atoms of hydrogen from each molecule, and having the general formula R.CO.H. It may therefore be locked upon as a ketone in which one alkyl is represented by H. Aldehydes may also be viewed as hydrides of acid radicles, hence CH₂.CO.H is called acetic aldehyde and not ethyl aldehyde, although the latter name is, etymologically, the more correct (p. 103).

Enumeration.—In the following list the numbers denote values of n. C_nH_{2n}O. 1. Formic; 2. Acetic; 3. Propionic; 4. Butyric; 5. Valeric; 6. Hexcic; 7. Heptoic; 10. Decoic; 12. Lanric; 14. Myristic; 16. Palmitic; 17. Steario.—O_nH_{2n-2}O. 3. Acrylic (acrolein); 4. Crotonic; 5. Tiglic; 6. Hexcenic; 8. Octenic; 14. Tetradecencic; 15. Cimicic; 21. Tri-ænanthio.—O_nH_{2n-2}O. Enuzoic; Phenyl-propionic.—O_nH_{2n-2}O. Enuzoic; Phenyl-propionic.—O_nH_{2n-1}O. Di-phenyl-acetic.—C_nH_{2n-1}O. Glycollic (?). Oxypropionic. Oxyputyric (aldel).—O_nH_{2n-2}O_x Elyoxul.—C_nH_{2n-1}O_x Oxyp-tenzoic; Furfur-acrylic, Furfurerotonic.—C_nH_{2n-1}O_x Snberio. Azelaic and Brassylio.—C_nH_{2n-1}O_x Di-oxy-benzoic; Piperonal.—C_nH_{2n-1}O_x Di-aldehydo-reserein; Di-aldehydo-orein.

Formation.—1. By oxidation of primary alochels by air and platinum-black, by aqueous chromic acid or by H₂SO₂ and MnO₂:

 $2R.CH_{x}.OH + O_{z} = 2R.CO.H + 2H_{z}O.$ 2. By distilling a mixture of barium or calcium formate with some barium or calcium salt: Ca(O.CO.R), + Ca(O.CO.H), = 2CaCO, + 2H.CO.R [Limpricht, A. 97, 368; Piria, A. Ch. [3] 48, 113; Krafft, B. 16, 1717). This process is a particular case of Williamson's method of producing mixed ketones. Instead of calcic formate, a mixture of calcic oxalate and lime may be used (Begusch, J. R. 7, 47).—3. From chlorides of the type R.CH.Cl. by heating with dry exalic acid (Anschütz, A. 226, 19).—4. Chromyl dichlorido, CrO.Cl., unites with toluene and its homologues when added to their solution in CS₂, forming brown powders, possibly of the form RCH(O.CrCl_OII)₂, which are decomposed by water with production of aldehydes (A. Étard, O. R. 90, 534; 97, 909; Bornemann, B. 17, 1462).—5. Aromatic aldehydes may be prepared by heating dichlorides R.CHCl, with NaOHAq, or the monochlorides, R.CH,Cl with aqueous lead or copper nitrate.—6. Alcohols of the form R.CH:CH.OH appear to change, at the moment of their formation into aldehydes, R.CH. CHO. The formation of aoroloin from glycerin, and of aidehyde from brome-ethylene are instances .-7. Some aldehydes, as benzeic, acetic, propionic, and butyrio, are produced by distilling albumen, fibrin, casein, or gelatin, with MnO, and H2SO. 8. Many aldehydes can be obtained from essential oils derived from plants; e.g., benzoic, cin-namic, cuminio, and salicylic aldehydes.

Properties.—Almost all are volatile liquids.

Reactions.—1. Are readily oxidised to acids,
and consequently are powerful reducing agents.

Retenic alcohels, R.OO.OH.OH, resemble aldehydes in reducing power (Zincke, A. 216, 817).—

2. Many are converted by alcoholic rotach or by potash-fusion into an alcohol and an acid: 20,H,.CHO+ KOH = 0,H,.CO,K+O,H,.CH,OH. Glycols with double the number of carbon atoms in the molecule are often formed.—3. Sodiumamalgam, or zine and glacial HOAc, reduce them to alcohols (Krafft, B. 16, 1714; Tiemann, B. 19, 355).-4. They combine with NaHSO, These compounds are usually soluble in water and in alcohol, but insoluble in saturated solutions of the bisulphites. Hence by shaking a liquid containing an aldehyde with excess of such a saturated solution, the aldehyde may be completely separated in the form of a crystalline compound. From these compounds the aldehyde may be set free by dilute H.SO, or Na, CO, and may then be distilled with steam (Bertagnini, A. 85, 179, 268) .- 5. They combine with phenylhydrazine (q. v.). A solution of phenyl-hydrazinc hydrochleride (1 pt.) and sodic acetate (1 pts.) in water (8 pts.) when added to an aqueous solution of an aldehyde or ketone, produces an inscluble compound, usually an oil appearing in drops producing a milkiness, but sometimes a crystalline pp. These compounds are not volatile with steam, but on boiling with dilute HCl they are resolved into phenyl hydrazine hydrochloride and the aldeliyde or ketone (E. Fischer, A. 190, 131; B. 15, 2252).—6. They form a silver mirror when heated with conc. annuoniaeal silver nitrate. The reduction is promoted by adding NaOH (Tollens, B. 15, 1635). 7. A solution of a rosaniline salt, bleached by SO₂, is reddened by aldehydes, in the cold (Schiff, Z. 1867, 175; Caro; V. Meyer, B. 13, 2342). This test is not infallible (Tiemann, B. 14, 791); it is given by aldehyde, paraldchyde, propiouic, iso-valeric, and cenanthic aldehydes. chloral, butyre-chloral, aerolein, furfurol, benzeic, cinnamic, and furfurcrotonic aldehydes, furfuracrolein, salicylic aldehyde, cimicic aldehyde; it is not given by chloral hydrate, formic acid, carbo-hydrates, prepyl alcohol and higher alcohols, pinacene, glycol, the phenels, or quinone; a faint colour is produced after some time by acetone, and methyl and ctliyl alcohols (G. Schmidt, B. 14, 1818).-8. Alkaline aqueons solutions produce a coloration like magenta when treated with diazo-benzene sulphonic acid and a little sodium-amalgam. Acctone and acetoacetic ether give a dark red coloration without the viclet shade (Penzoldt a. E. Fischer, B. 16, 657) .- 9. Aldehydes are converted by hydroxylamine into aldoxims

R.CHO + H₂NOH = R.CH: NOH + H.O (V. Meyer, B. 15, 1164, 1324, 1525, 2784; 16, 822, 2992).—10. u-di-methyl-p-phenylene diamina acts vigerously on aldehydes—in alcoholic selntion forming crystalline compounds (A. Calm, B. 17, 2938): Ph.CHO + NH₂.C₃H₁.NMe₂ = Ph.CH: N.C.H₂.NMe₂ + H₂O.

Ph.CH:N.C_sH.NMe₂+H₂O.

11. Hemologues of acctic aldehyde form crystalline compounds with anmenia, R.CH(NH₂)OH.

These are converted by H₂S into sulpifur bases (v. p. 104, l. 9). The aromatic aldehydes are converted by ammonia into hydramides:

3Ph.CHO + 2NH, = (Ph.CH), N, + 3H.O. Some fatty aldehydes, e.g. iso-bntyrio aldehyde (Lipp, A. 211, 344) behave similarly. Acroleia loses only half its oxygen:

 $2C_3H_4O + NH_3 = C_4H_4NO + H_2O_4$

Primary and secondary bases act upon aldehydes with elimination of water. The neutral products are split up by HCl into their components. 12. Chlorine forms derivatives by substitution.—18. PCl, displaces O by HCl,.—14. H.S displaces O by S, forming thio-aldehydes, or their polymerides.—15. PCl, combines with aldehydes; the compounds are converted by water into phosphinic acids (q. v.).—16. PH,I forms crystalline compounds (v. Phosphines).—17. Aldehydes react with alcohols forming acetale (p.):

 $R.CHO + 2HOR' = R.CH(OR')_2 + H_2O.$ Similarly, mercaptans form mercaptals (Baumann, B. 18, 884).—18. Alkyl chlorides form chlorinated ethers (e.g. CH₃CHCl.OEt). Alkoyl chlorides act similarly (p. 105, l. 1).—19. Hydric cyanide combines with aldehydes, forming a oxynitriles. These nitriles give (a) on saponification, oxy acids, (b) on treatment with ammonia, amido-nitriles, whence amido acids may be got (Tiemann, B. 14, 1965) .- 20. Aldehydes may be converted into amido-acids by allowing them to stand for 30 minutes with a 3 p.c. solution of NH,CN, and then boiling with HCl (Liubawin, J. R. 13, 506).-21. Benzoic aldehyde reacts with nitro-parassins thus: Ph.CHO + H.C(NO2).CH3 = Ph.CH:C(NO₂).CH₂ + H₂O (Priebs, A. 225, 319). 22. Aldehydee condense with aromatic compounds with elimination of II,O and formation of tri-substituted methanes. Thus aldehydo and benzene give di-phenyl-methyl-methane; benzoie aldehyde and phenol givo di-oxy-triphenyl-methane; benzoie aldeliyde and aniline give di-amido-tri-phenyl-methane. -23. In presence of small quantities of acids, aldehydes form red resins when warmed with phenols. Many of these are converted by excess of acide into crystalline isomerides. Thus benzoic aldehyde forms with pyrogallic acid prisms (from ether) of $C_2H_2O_6$; this forms an acetyl derivative $C_{20}H_{24}Ac_2O_6$. Benzoic aldehyde and resorcin form $C_{24}H_{24}O_4$. If a few drops of a liquid containing an aldehyde be boiled with an alcoholic solution of resorcin and a little HCl, and be then poured into water, a pp. is formed. This may be used as a test for presence of aldehydes (Baeyer, B. 5, 25; Michael a. Ryder, B. 19, 1388) .-- 21. In dilute alkaline solution aldehydes condense with ketones or other aldehyles with elimination of H.O, and production of complicated aldehydes or ketones.-25. For the reaction between aldehydes and o-di-amines v. ALDEHYDINES.

Perkin's Synthesis of Unsaturated Acids.—Benzoic aldelyde, acetic anliydride and sodic acetate, heated togother form sodic cinnamate.

In this reaction the sodic acetate may be exchanged for sodie butyrate or valerate, but the product will still be sodic cinnamate: hence Perkin concludes that the reaction takes place between the aldehyde and the anhydride. Fittig-came to the opposite conclusion, viz., that the aldehyde acted on the sodium ealt and that the nature of the anhydride was immaterial, thus if sodic succinate and acetic anhydride were used, the condensation took place with the eodic succinate. To this Tiemann (B. 15, 2061) objected that possibly the acetic anhydride acting on the sodic succinate formed sodic acetate and succinic anhydride, and that the latter reacted upon

the aldehyde. Stuart (B. 16, 1486) then showed that when sodic malonate was used condensation took place between it and the bengoic aldehyde, although no malonic anhydride is known. He also showed that in this case glacial acttic acid might be substituted for acctic anhydride. According to Fittig, aldol-like condensation-products are first formed, and these, when they split off water, give the unsaturated acids (A. 227, 49). This is shown by the action of eodic iso-butyrate on benzoic aldehyde in presence of isobutyric anhydride, whon the anhydride of the isobutyric derivative of β-oxy-β-phenylvalerie acid Ph.CII(OH).CMe, CO, H (q. v.) is formed; in which there is no H for the OH to split off with. If NaOAo be used inetead of sodium isobutyrate oxy-phenyl-valerie acid is still formed, a result that supports Perkin's vicw (Perkin, C. J. 49, 317).

Enanthol and valeric aldchyds may be substituted for benzoic aldchyde in these syntheses, while the sodium salt and anhydrido of propionic or n-butyric acid may be used instead of the corresponding derivatives of acetic acid. Condensation then takes place in the α position:

Ph.CHO + CH_3 - CH_4 - CO_2 Na = Ph.CH: $CMe.CO_2$ Na + H_2O .

A dibasic acid can unite with one equivalent of an aldehyde for each CH₂, CO₂H contained in its formula; the product may then lose H₂O₂, becoming a lactonic acid or an unsaturated acid.

ALDĒHYDINES.—This name was formerly applied to the base C_sH_1N obtained by heating aldehyde-annonia, since shown to be tri-methylpyridine (q, v). The same name has since been used by Ladenburg (B. 10, 1126; 11, 590, 1650) to denote bases obtained by mixing dilute aqueous solutions of aromatic c (but not m or p) di-amine hydrochlorides with aldehydes. Condensation occurs with evolution of heat; the yield of aldehydine after crystallisation ie 60 to 70 p.c. of the theoretical. $C_xH_y(NH_2)_+ 2H.CO.R = C_xH_yN_zC_H_zR_+ 2H_zO$. Thus o-tolylene-di-amine hydrochloride and benzoic aldehyding ive rise to $C_zH_xN_zC_zH_zP_b_z$ tolylene benzaldehydine.

The same body is formed by the action of benzyl chloride on anhydro-benzoyl-tolylene di-

amino at 160°, C_1H_0 $\stackrel{N=C.Ph}{\sim}$. Hinsberg con-

cludes from this that its formula is

N = C.Ph

C,H₆ (B. 19, 2025). Under the con-

ditions of this experiment molecular change is more likely to occur than in the usual preparation of aldehydines in aqueous colution. All other considerations point to a symmetrical formula; and since in stahility and other properties these bodies resemble the quinoxalines it is probable that, together with the latter, thay belong to the class of azines: tolylene benzalde-N-CHPh

hydine would then be C.H. N-CHPh
formula given by Ladenburg is
N.—CHPh

C,H, N CHPh

Phenylene-anis-aldehydine C₁₂H₂₈N₂O₂ [129°]. Needles. Soluble in alcohol. Prepared

by shaking anisic aldehyde with an aqueous solution of o-phenylene-diamine hydrochloride.

B'HCl: needles; difficultly coluble in water, Phenylene-benzaldehydine C₂₀H₁₆N₂ i.e.

N_CH.C.H, [133°-134°]. CH.C.H.

Six-sided prisms. Insoluble in water, easily soluble in alcohol and benzene.

Preparation .- (1) By heating o-phenylenediamine with benzaldohyde. (2) By chaking benzaldehydo with an aqueoue colution of ophenylenediamine hydrochloride.

Salts .- B'HCl: eolourless prisms.

(B'HCl), PtCl : yellow precipitate. B'HNO. Slightly soluble prisme. B',H,SO4: colourless

Ethylo-iodide C20H16Nz(C2H5)I [2110-2130].

Colourless prisme. Methylo-iodide C20H16N2(CH3)I. Prisms.

Phenylene-furfur-aldehydine C18H12N2O2 i.e. (1)N__C1I.C,H₃O [95°-96°].

 $\searrow_{(2)N} \stackrel{\triangle}{\hookrightarrow}_{CH.C,II_{a}O}$ Colourlese crystals. Soluble in alcohol and CoH4, with difficulty in ligroin, insoluble in water. Prepared by shaking furfurol with an aqueous

solution of o-phenylenediamine hydrochloride. Salts. - (B'HCl), PtCl4: yellow leaflets. B'HNO,: slightly coluble needles.

Melhylo-iodide $C_{16}H_{12}N_2O_2(C\Pi_3)I$. [192°-195°]. Prisms.

 $C_{23}H_{22}N_2O_2$ Tolylene-anisaldshydine

(1)N _CH.C₆H₄(OMe) [152°-156°] (2)NCH.C₈H₄(OMe)

Needles. Propared by the action of anisio aldohyde on an aqueous solution of o-tolylenediamine hydrochloride.

Tolylene-benzaldehydine $C_{21}H_{18}N_2$ (1)N_CH.C₆H₅ [195·5°]. `(2)N´

Colourless prisms. Forms unstable salts with acids. Prepared by heating benzaldehyde with o-tolylene diamino to 110°. Yield 45 p.c. of tho diamine.

B'HCl + H2O: long needles: difficultly soluble in etrong HCl.

Elhylo-iodide [180°-181°]. Needles or prisms (+ ½H₂O). Soluble in water. With iodine it forms a periodide C₂₁II₁₂N₂(C₂H₅)I₁ [123].

Givee a strongly alkaline solution with Ag.O which on neutralising with HCl and adding PtCl, gives a crystalline platino-chloride (C₂₃H₂₃N₂Cl), PtCl₁.

Methylo-todide [209°]. Thin white needles. Phenylene-benzalde Hydine carboxylic acid

(1)N _CH.C,H, C₂₁H₁₆N₂O₂ i.e. CO₂H.C₆H₃ (2)N CH.C.H.

Prepared by the oxidation of tolylenc-benz-aldehydine with KMnO. Is not altered by heating with HCl to 200°.

Salts: A'Ag: white precipitate. A'2Ca: dillicultly soluble needles.

Telylene furfuraldehydine

[128]°]. Thin white prisms. Easily soluble in alcohol, ether, &c., with difficulty in cold ligroin.

Preparation.—(1) By heating furfured with o-tolylenediamine. (2) By adding furfured (20 pts.) to a solution of o-tolylenediamine bydrochloride (20 pts.) in 80 pts. of water; on standing the hydrochloride separates out and is purified by crystallising the base repeatedly from ligroin; yield 55 p.c. of the theoretical.

Salts: B'HNO,: needles. B'₂H₂SO₄: prisms

(B'HCl) PtCl,: yellow crystals.

Methylo iodide [195.5°]. Leaflets. Difficultly soluble in water; bitter taste; powerful poison.

Methylo chloride: leastets; easily soluble in water. Powerful poison.

Methylo-triiodide C, H, N, O, (CII,)I, [126c_ 128°]. Light brown needles.

Methylo-pentiodide C1, H11N2O. (CH3)Is [109°]. Steel-blue pillars.

p-ALDEHYDO-BENZOIC ACID C,H,O, i.e. C, H, (CHO)(CO, H) [c. 216°]. Formed by careful oxidation of terephthalic aldehyde with chromio mixture (Löw, A. 231, 365; B. 18, 917). Needles (from water). Small needles when sublimed. M. sol. ether or chloroform, sl. sol. hot water.

Salt.-AgA'. Ether.-EtA'. Shows oharac-

ters of benzoic aldehyde. Reactions .- 1. Doce not reduce ammoniacal AgNO₃. Its ethyl ether, however, reduces ammomacal AgNO2.-2. Doce not give Perkin's reaction with NaOAo and Ac.O .- 3. With ZnCl. and alcoholic NPhMe2 it forms the zinc carboxy-

late of lenco-malachite green, [147°]. Phenyl hydrazide [226°].

p-Aldehydo-nitro-benzoic aeid

C_sII₃(NO₂)(CHO)CO₂H. Nitro-terephthalie-aldehyde-acid [160°]. Large four-sided prisms. Easily soluble in alcohol and other, sparingly in chloroform. Formed by nitration of p-uldehydo-bonzoie acid.

With acetone it gives the indigo-reaction (Löw, B. 18, 948).

ALDEHYDO-ĆINNAMIC ACID v. CINNAMIO

ALDEHYDO-NAPHTHOL v. OXY-NAPHTHOIG ALPEHYDE

ALDEHYDO-OXY ACIDS. Got by heating aromatic oxy acids with chloroform and aqueous NaOH (Tiemann a. Reimer, B. 9, 1268). C_oH₄(ONa)CO,Na + 3NaOH + CHCl₂ =

 $\hat{C}_{a}H_{a}(COH)(ONa)CO$, $Na + 3NaCl + 2H_{a}O$. The COH takes either o or p position towards the hydroxyl.

ALDEHYDO-OXY-BENZOIC ACIDS C.H.O. m-Aldehydo-salicylic acid

C, H, (Oll)(CllO)(CO,H)[1:4:2]. [249°]. S.7 at

100°; .038 at 25°. Preparation. - Salicylie acid 714 pts.), NaOH (25 pts.), water (50 pts.) and chloroform (15 pts.), are boiled for some hours, the product dissolved in water, and acidified with HCl. A yellow pp. is formed and is extracted with ether. The ethercal solution is shaken with agueous NaIISOa; this solution, when boiled with dilute H2SO, deposite a crystalline pp. consisting of the (1, 2, 6) acid; the (1, 4, 2) acid remaining in the solution, from which it may be extracted by ether (Tiemann a. Reimer, B. 9, 1268; 10, 1563).

Properlies .- Long delicate yellowish needles. V. sol. ether or not alcohol, v. sl. sol. chloroform.

action on the skin, affected the colour of many plants, and reacted with acids with effervescence and the production of new substances wherein neither the properties of the acids nor those of the alkalis were prominent. Van Helmont and his successors recognised two kinds of alkali, fixed and volatile; Duhamel, in 1736, divided fixed alkali into two classes, vegetable (potash), and mineral alkali (seda). Little or nothing was known regarding the composition of alkali until the year 1755, when Black (on the occasion of graduating as M.D. at Edinburgh) published his dissertation on 'Magnesia Alba, Quicklime, and other Alkaline Substances.' Magnesia alba dissolved in acids with effervescence; but after being strongly heated no effervescence attended the solution of this alkali. The notion of Basil Valentine (end of 15th and beginning of 16th century), that lime when burnt combined with 'matter of fire,' had been accepted by many as an explanation of the difference in the behaviour towards acids of burnt and unburnt lime. If this explanation applied to magnesia it should be possible perhaps to get hold of this 'matter of fire,' which combined with the magnesia alba when that body was heated. But Black found that a given mass of magnesia alba weighed more than the calcined magnesia obtained from it. Hence something was lost instead of gained during the process of heating. This something proved on further quantitative examination to be a gas different from common air; to it Black gave the name of fixed air. The effervescence or non-effervescence of alkalis with acids was proved by Black to accompany the presence or absence of fixed air (carbonic acid). From this time a distinction was clearly drawn between alkalis, which dissolved in acids without effervescence, and carbonated alkalis, the solution of which in acids was accompanied by the escape of oarbonic acid gas. It was recognised that whether a caustic or a carbonated alkali were dissolved in an acid, the body which remained in solution, and which had no close resemblance either to the acid or the alkali, was one and the same.

The properties of the alkalis were supposed by the older chemists to be due to a 'principle of alkalinity,' or sometimes to a 'principle of saltness,' which latter principlo was common to acids, alkalis, and the products of their mutual action, i.e. salts. Closely allied to, and sometimes regarded as identical with, the alkalis, was the group of earths. These bodies were known to neutralise acids and affect colouring matters like alkalis, but they were much less soluble in water than the alkalis. It was taught by some chemists that an alkali is hidden in every earth, and by others that an alkali is an earth refined by the presence of acid and combustible matter. Black's exact quantitative investigations tended to disparage all such explanations as these; but it yet remained to find the precise composition of the alkalis and the earths. Lavoisier thought that these bodies must be compounds; but, as he had no means of proving this, he classed them with the elements, while suggesting that the earths were probably compounds of oxygen with ınknown metals. In 1807 Davy decomposed wo alkalis, potash and soda, by passing an dectric current through these substances when molten; and a year later he succeeded, by the same agonoy, in separating the earthy bodies lime, baryta, and strontia, into oxygen and, in each case, a metal.

The name alkali is now generally applied to the compounds of hydrogen and oxygen with one or other of the five metals, lithium, sodium, potassium, rubidium, cæsium (v. Alkalis, Metals or the); an aqueous solution of ammonia is also regarded as containing an alkali, viz. a compound of hydrogen and oxygen with the radiele ammonium (v. Ammonium compounds). The alkalis are classed with the hydroxides, i.e. compounds of hydrogen and oxygen with a third element, rather than with the hydrates, i.e. compounds of water with an oxide or a salt (v. Hydrates). The general formula of the alkalis. is written MOH rather than M2OH2O; M=Li, Na, K, Cs, Rb, or NH. The alkalis are very soluble in water; these solutions neutralise acids forming salts, and also precipitate most of the heavy metals from their solutions in the form of oxides or hydrated oxides; aqueous solutions of the alkalis act corrosively on animal and vegetable substances, and also alter the tint of many colouring matters. When moist, the alkalis, with the exception of ammonia, readily combine with carbonic acid to form carbonates. Lithia is much less solublo in water than the other alkalis. The solid alkalis are not decomposed by the action of heat alone.

M. M. P. M.

ALKALI-BLUE v. PHENYL-ROSANILINE SUL-PHONATE OF SODA.

ALKALIMETRY.—The estimation of alkaliaby volumetric methods, v. ANALYSIS.

ALKALINE EARTHS, METALS OF THE. -CALCIUM, STRONTIUM, BARIUM .- Certain substances, more or less alkaline in their properties, but differing from alkali chiefly in being insoluble in water, were known from early times; these substances were called earths. After a time some of the earths were found to dissolve in water, although to a less extent than alkalis: these comparatively soluble carths were separated from the others and classed together as the alkaline earths. The best known alkaline earth is lime; this substance was long considered identical with baryta and strontia, but in 1774 Scheelo proved that baryta was different from lime, and in 1792 Hope distinguished strontia from the two other alkalino earths. After decomposing the alkalis potash and soda, Davy applied the agency of electricity to the three substances just named, and in 1808 he succeeded in separating each into oxygen and a metal. Davy made his experiments quantitative; he also synthesised the three alkaline earths from oxygen and the metals he had himself discovered; thus he proved the alkaline earths to be metallic oxides. The metals calcium, barium, and strontium were not obtained in a state of approximate purity until 1855. The metal magnesium is sometimes classed with calcium, barium, and strontium; but, on the whole, it seems better to place magnesium with zinc and cadmium (v. p. 114, also MAONE. SIUM GROUP OF METALS).

Some of the principal data regarding the metals of the alkaline earths are presented in the following tables.

ALKALIS, METALS OF THE.

		CALCIUM	BTBONTIUM	BARIUM
Atomic weights .		39-9	87.3	136.86
No compounds oxides and oblorio weights unknown.	gasified les dedu	Combining weigh	ts determined; and m	ost probable formulæ or oxides, &c. Molecula
Melting points (data uncerta	'I	high red-heat above strontium	moderate red-heat	below red-heat
Specific gravitie	28	1.58	2.5	8.75
Specific heats Atomic weight		0.17	not determined	not determined
Spec. grav.	}	25.3	34.9	86 ·5
	. Hee	ats of formation in a	queous solutions (Thor	nsen).
[M,Ol ² ,Aq]		187,600	195,700	196,800
[M,Br2,Aq]	:	165,800	173,800	174,900
[M,I2,Aq]	, 1	135,000	143,400	144,500
,	•		oxides:	
. (pA,O,M)	• 1	149,260	157,780	158,760
[M,O',H',Aq] .		217,620	droxides:	227,120
	• •	•	tion (Thomsen).	,
			loid salts:	
Morenioj .	•	21,750	18,640	[BaOl ¹ ,2H ² O] 7,000
[MBr*,6H*O] •	•	25, 600	23,330	[BaBr ² ,2H ² O] 9,110
4	•	(2) Of	rides.	. 0,110
мо,н,ој	. 1	15,540	17,700	22,260
	Harla of		ides in solution (Thom	
MOAq,H'SO'Aq] .	. i	81,150	31,150	81,150
MOAq,H2Gl2Aq] MOAq,H2N2O2Aq]	1.	27,640	27,640	27,640
	1			
		Calcium	STRONTIUM	Barium
Malleability, co- lour, &c.	ham tle ;	ductile, but when mered becomes brit- whitish - yellow; lness about same as	Ductilo and malle- able; colour re- sembles calcium but olearer; harder than lead.	gold-yellow colour.
Wave-lengths of .mostcharacteris- bic lines in spectra.	H (\	cllow) 5588. riolet) 3969. do.) 3933·8.	Sr. (blue) 4604	Ba, (yellow) 5538.
Chemical proper- ties.	water to rewith coming the coming	y exidises in moist decomposes cold r rapidly; heated dness in air, burns out smoke; readily bines with Cl, Br, I, d S, at high tem- tures.	Closely resembles calcium; decomposes water more rapidly.	Resembles calcium burns when heated i O-H flame.
Decurrence and preparation.	rocke anim sulpl silica elect	widely diffused in s, waters, plants, and hals, as carbonate, hate, phosphate, and hate: prepared by rolysis of mixture 22 SrOl ₂₁ and NH ₂ Cl.	Not very widely dif- fused; occurs as carbonate and sul- phate in rocks, and wators; prepared by electrolysis of fused SrCl _s .	Not very widely diffuse occurs as carbonate sulphate, and sil- cate, in rocks, water and certain plants prepared by electro "s of BaCl, mixe

General Formula and Character of Salts.

MO, MO, MO, MO, MS, MS, MS, MS, MX, (X=Cl, Br, I, F, CN), MSO, M2NO, MCO, &c., where M = Ca, Sr, or Ba. MO, decomposed by heat. Salts for the most part white; no great tendency to form double salts; polysulphides known, SrS,6H,O and BaS,H,O, in definite crystals. Oxides and hydroxides markedly basic; latter, except that of Ba, decomposed by heat alone into oxides and water; almost all similar salts isomorphous; many salts isomorphous with corresponding compounds of Mg, e.g. carbonates; most, with corresponding compounds of lead; MO and MO2H2 not very soluble in water, solubility increases as atomic weight of metal increases; MC12 and MBr2 easily soluble, colubility decreases as a tomic weight of metal increases; CaSO, very slightly soluble (S. 272 at 38°), SrSO, nearly insoluble (S. 01 at 100°), BaSO, insoluble. CaCO, slightly soluble (S. 1.13 at 100°), SrCO, and BaCO, nearly insoluble. Nitrates all soluble, solubility decreases as atomic weight of metal increases; Ca2NO, S. 93 I at 0°. Sr2NO, S. 54 9 at 10°. Ba2NO, S. 7 at 10°.

These data show that the metals of the alkaline earths differ from the alkali metals (compare data for latter on p. 115); the former are not so readily oxidised as the latter; the heats of formation of the oxides of the alkaline cartly metals are smaller than those of the alkali metals; the hydroxides of the alkali metals cannot, but the hydroxides of the alkaline earth metals except that of Ba can, be separated into oxides and water by the action of heat alone. The alkali metals are epecifically lighter than those of tho alkaline earths; the composition of the oxides and chlorides of the former is represented by formulæ containing two atoms of metal to one of oxygen or two of chlorine, while that of the corresponding salts of the latter is represented by formulæ containing one atom of metal to one a of oxygen or two of chlorine. The salts of the alkali metals, as a class, are much more soluble in water than those of the alkaline earth metals; the hydroxide, carbonate, and phosphate of lithium are, however, considerably less soluble than the corresponding salts of the other alkali metals (v. Alkalis, Metals of the, p. 115). though magnesium forms the oxide MgO, the ohloride MgCi, and the sulphate MgSO, salts analogous in composition to the oxides, chlorides, and sulphates of the metals of the alkaline earths. nevertheless this metal is clearly cut off from these by the following, among other, characteristics. The heats of formation, in aqueous solutions, (1) of the haloid salts of Ca, Sr, and Ba, (2) of Mg, Zn, and Cd, indicate the existence of two groups, in the first of which (Ca, Sr, Ba) the value of the reaction increases, and in the eccond of which (Mg, Zn, Cd) the value of the reaction

ц,	[M, Ol*, Aq]	[M, Br*, Aq]	[M, 1°, Aq]
Ca	187,600	165,800	135,300
Sr	195,700	173,800	143,400
Ba	196,300	174,400	144,000
Mg [•] Zn Cd	186,900	165,000	134,600
	112,800	90,900	60,500
	96,300	74,400	44,000

tin affected the colour of many | molten; and a year later he superaction of the metals formula and Character of Salts. | usoresses; as the atomic weights of the metals increase. The data are from Thomsen:

Magnesium is scarcely oxidised in ordinary air; it does not decompose cold water; nor does it combine so readily with the halogens as the metals of the alkaline earths do. The spectrum of magnesium, as produced in the electric arc, is marked by a series of triplets alternately sharply marked and diffuse, and diminishing in brightness towards the more refrangible side; the spectra of barium and strontium show no triplcts, but a series of lines only; the spectrum of calcium is marked both by lines, perhaps homologous with those of barium and strontium, and also by well-marked triplets (Liveing and Dewar). Magnesium eulphate is very soluble in water: this salt, and also the carbonate and chloride, readily combinee with salts of the alkali metals to form double compounds. Magnesium oxide is scarcely if at all soluble in water; the formation of the hydroxide by the action of water on the oxide is attended with the production of very little heat: [MgO, H'O] = (approx.) 3,000 (Thomsen).

The mutual relations of the two groups of elements-the alkaline earth metals and the magnesium metals-are suggested by the position they occupy in the classification based on the periodic law (q. v.; v. also CLASSIFICATION). Both belong to Group II.; but Ca, Sr, and Ba occur, along with Be, in even series, and Mg, Zn, and Cd, along with Hg, in odd series, of that group. The metal beryllium exhibits analogies both with the alkaline earth, and with the magnesium, metals; it is one of those elements called 'typical' by Mendeléeff (v. BERYLLIUM). For accounts of the metals of the alkaline earths and their. binary compounds v. the articles Bankum, Calcium, and Strontium; and for the other salts of the metals v. Carbonates, Nitrates, Sulphates, &c. M. M. P. M.

ALKALIS, METALS OF THE. (LITHIUM, Sodium, Potassiem, Rubidium, Casium.)-The history of the namo alkali has been briefly traced in the article under that heading. The alkalis potash and eoda were decomposed by Davy in 1807; lithia (discovered by Arfvedson in 1817) was decomposed by the same chemist about 1818; casia and rubidia were discovered by Bunsen and Kirchoff in 1860-61, rubidium being obtained in the same year by Bunsen, by electrolysing the chloride; approximately pure casium was not prepared until 1882, in which year Setterberg obtained the metal by cloctrolysing the double eyanide of easium and barium. The more important properties of these metals and of their principal salts are presented in the tables on the next page and page 116.

Thermal values of reaction with water.—When an alkali metal reacts with water an alkaline hydroxide is formed and dissolved, and hydrogen is evolved; thus:-

 $M_2 + xH_2O = 2MOHAq + (x-2)H_2O + H_2$

This reaction would be expressed in the notation of thermal chemistry thus:--.

 $[M^2,2H^2O] = -2[H^2,O] + [M^2,O^2,H^2,Aq].$

The value of 2[H2,O] is 136,720 gram-units when H.Q represents 18 grams liquid water; when the value of [M2,O2,H2,Aq] considerably

· <u></u>	Leihiùm	SODIUM	POTABSIUM	Busingon	CASIUM
Atomic weights .	7.01	23	39-04	86-2	182-7
No compounds a oxides and chlorides salts which take par	deduced by ch	emical metho	ls from consid	dering emallest	ble formula of
Melting points	180°	95°-5°	58°_62°	1 38°	26°-27°
Specific gravities .	0.59	0.98	0.87	1.52	1.88
Specific heats	0.94	0.29	0.17	not determined	not determine
Atomic weight Spec. grav.	11.9	23.5	44.9	56.1	70.6
	Heats of form	ation in aqueo	us solutions ('.	Thomsen).	•
		(1) of haloi	d salts :		
[M ² ,Cl ² ,Aq]	204,500	193,000	202,300	_	_
$[M^2,Br^2,Aq]$	182,600	171,200	180,500	_	
[M ² ,I ² ,Aq]	152,200	140,600	150,000	_	
	(2)	Of oxides and	hydroxides:		
[M²,O,Aq]	166,500	155,300	164,600	ı –	
$[M^2,O^2,H^2,Aq]$	234,900	223,600	232,900		
· .	Teats of neutral	isation of oxid	les in solution	(Thomsen):	
M2OAq,H2SOAq] .	31,150	31,150	31,150	i –	
[M°OAq,H°Cl²Aq]	27,640	27,640	27,640		
[M ² OAq,H ² N ² O ⁶ Aq] }	,020	,,010	,,010	1 -	

exceeds 136,720, we should expect the metal M to decompose liquid water. Thomsen has determined these values:—

 $\begin{array}{c} M & [M^2, O^2, H^2, Aq]. \\ M = Li_2 & 234,900 \\ Na_2 & 223,600 \\ \bullet & K_2 & 232,900 \end{array}$

General formulæ and characters of salts .- M_2O_1 (M_2O_2 , M_2O_4), MOH_1 , M_2S_1 , (M_2S_2 , M_2S_3), MSH_1 , $MX(X=Cl_1,Br,I,F,CN)$, M_2SO_4 , $MHSO_4$, MNO₃, M₂CO₃, MHCO₃, &c., where M = Li, Na. K, Rb, or Cs. No oxides or sulphides of Rb and Cs have been prepared in a state of purity. Li.O is the only oxide, and Li.S the only sulphide, of Li known with certainty. Na O and K2O4 are very stable towards heat, but quickly decompose in moist air, giving off oxygen and forming NaOH and KOH. Salts for the most part white, and very soluble in water; but LiOH is much less soluble than the other hydroxides, and Li2CO3 and Li2PO4 than the other carbonates and phosphates—(Li,CO₃, S. 769 at 18°, S. 778 at 100°; Li,PO₄, S. 04 at 18° [approx.].) Chlorides, except LiCl, form many double salts with chlorides of heavy metals, e.g. M.PtCl., SbCl.6MCl, &c. Sulphates, except Li.SO., form alums, also double salts with sulphates of magnesium group. Most salts are isomorphoue, but some of the lithium salte are not strictly isomorphous with corresponding ealts of the other metals; some compounds of silver and thallium are isomorphous with those of the alkali metals. All the metals of this group are electropositive towards any other elements; thoir oxides and hydroxides are strongly basic. The latter cannot be decom-posed by heat alone into oxides and water. Lithium differs from the other members of the group in the comparative insolubility in water of its hydroxide, carbonate, and phosphate, in

the non-formation of an alum, and in some other respects (compare heate of formation of analogous salts); this element serves to connect the group of the alkali metals with that of the metals of the alkalino earthe in somewhat the same way as the latter group is connected with zinc and cadmium by the olement magnesium (v. Alkaline earths, Metals of the). metals copper and silver are to some extont connected with the alkali metals. Copper forme two series of salts represented by Cu2O and CuO respectively; the former, so far as composition goes, are analogous to the alkali salts. They are, however, much more insoluble in water than these, and, with the exception of the iodide and cyanide and some double salts, are mach less stable than the salts formed from the oxide CuO. The salts of silver, as a class, are much less soluble in water than those of tha alkali metale; their composition is similar-Ag₂O, AgNO₃, Ag₂SO₄, &c.; some of them are isomorphous with corresponding sodium ealts, e.g. Ag, SO. Silver forms an alum, and its oxide is markedly basic.

The alkali metals are placed in Group I., according to the classification of elements based on the periodic law, and this group also contains the metale Cu, Ag, and An. Li, K. Cs, and Rb belong to even sories, and Na. Ou, Ag, and Au, to odd series, of Group I. There can be no doubt, however, that eodinm is closely connected with the other alkali metals, and that the three heavy metals (Cu, Ag, An) present only feebly marked analogies to each other, and to the metals of the alkalis. In considering the classification of elements which the periodic law presents, attention must be paid, net only to the group in which any given family of elements occurs, but also to the character of the elements which precede and those which

position of the family in the complete scheme must also be considered (v. Periodio Law).

In some respects thallium exhibits a markod chemical resemblance to the alkali metals; it forms an oxide Tl₂O and a hydroxide TlOH, both of which dissolve in water, producing a strongly alkaline and basic liquid, marked by most of the properties which characterise aqueous solutions of soda and potash; it also

follow the given family in the same series; the | forms salts-Tl₂CO₂, Tl₂SO₄, TlNO₂, &c.which, as a class, are easily soluble in water, and many of which are isomorphons with the oorresponding alkali salts. Some of the thallous salts, however, resemble those of lithium in being comparatively insoluble, e.g. TICl and Tl.PO. Thallium also forms an alum, and a double platinum chloride Tl.PtCl. On the other hand, the metal itself differs much from the alkali metals; it is heavy, is not very easily

	LITHIUM	Sodium	POTASSIUM	RUBIDIUM	Cæsium
Malleability, colour, &c.	Silver-white; easily drawn into wire, butless tenacious than lead; very soft, may be welded at ordinary temperature; not volatile at red heat.	soft as wax at ordinary	tie at 0°, malleable at 5° or so, pasty at 15°: can be		Silvor-white; soft at or- dinary tem- perature.
Wave-lengths of most characteristic lines in spectra.	Li. (red) 6705 (blue) 4602	D ₁ (orange) 5895 D ₂ (orange) 5889	(yellow) 5800 K _s (violet) 4044	Rb, (red) 7800 Rb, (orange) 6297	Cs, (blue) 4597 Cs. (blue) 4560
Chemical properties.	Oxidises in ordinary air but not so rapidly or completely as other metals of the group; decomposes cold water rapidly but without itself melting; ignites at temperaturo much above its melting - point; readily combines with halogens and sulphur.	pidly in air; decomposes water ra- pidly; com- bines very energetic- ally with ha- logens and sulphur, de- composes	air ; decom-	air so rapidly that usually takos fire; decomposes water most rapidly;	essily oxi- dised. Pro- perties not yet exactly studied.
Occurrence and preparation.	Widely diffused in rocks, waters, plants, and some animal secrotions; occurs as silicate and phosphate with other alkali metals; prepared by electrolysis of mixture of LiCl and NH ₄ Cl.	Inlarge quantities as chloride, silicate, fluoride, nitrate, &c., prepared by deoxidising Na ₂ CO ₃ by hot carbon.	Inlargequan- tities as ni- trate, sili- cate, sul- phate &c. propared as Na.	diffused, but in very small quan- tities; in	a rare min- cral. Inmi- nute quan- tities in many rocks and waters; prepared by electrolysis of double oy-

oxidised, does not decompose water except at a red heat, and is much more electro-negative than the alkali metals. Thallium forms an oxide, Tl.O., from which a series of salts—Tl.3SO., TlOl., &c.—is obtained; these salts exhibit analogies with those of the earth metals. The heats of formation of thallous oxide, hydroxide, and chloride, are much smaller than those of the alkali salts; Thomsen gives these numbers: [Tl2,O,Aq] = 39,200; [Tl2,O2,H2,Aq] = 107,500; [Tl2,Cl2,Aq] = 76,900 (v. EARTHS, METALS OF THE, and THALLIUM).

An aqueous solution of ammonia is strongly alkaline; when neutralised by acids salts are obtained which, as a class, closely resemble those of the alkali metals, with which they are, for the most part, isomorphons. These salts are con-sidered to be compounds of the radiole ammonium (NH,) with acid radicles; the general formnlæ given for salts of the alkali metals apply to the ammonium salts if M be taken to represent NH. This radiole ammonium replaces the elements Li, Na, K, Rb, or Cs, in most compounds without altering the crystalline

form, and without changing the chemical type, of these compounds. The salts of ammonium are, therefore, classed with those of the alkali metals. (For more details regarding the constitution of those salts, and for an account of their properties, see Ammonium Compounds.) For accounts of the individual alkali metals and their binary compounds, see the articles Cæsium, Lithium, Potassium, Rushium, and Sodium; and for the other salts of these metals see Carbonates, Nitrates, Sulphates, &c.

M. M. P. M.

ALKALIS, Action on Organic Compounds. The tendency of alkalis is to form salts. Thus they react with acids and other hydroxylio compounds by displacing the hydrogen by potassinm or sodium (p. 53). Neutral substances are frequently saponified by alkalis, i.e. turned into salts. Saponification means soap-making; in the narrowest senso it means boiling a fat with potash or soda: $C_3H_3(OC_{18}H_{35}O)_3 + 3KOH = C_2H_3(OH)_3 + 3KOC_{18}H_{35}O$. In a broader sense it means the splitting up of any compound ether into its alcohol and its acid, whether by means of an alkali, an acid, or by water alone. In the broadest sense it means the conversion of a neutral substance into an acid or the salt of an acid. Alkalis saponify compound ethers, nitriles, amides, and amic acids. In the case of nitriles the reaction takes place as follows: $R.CN + KOH + H_2O = R.CO_2K + NH_3$. act upon chlorinated or brominated substances with production of haloid salts: the reaction is either one of substitution; CH₃Cl + KOH = CH, OII + KCl: or else HCl or HBr is abstracted; $CH_{..}Br.CH_{.}Br + KOH = CH_{..}CHBr + KBr + H_{..}O.$ The latter equation represents the action of alcoholic KQH on chlorinated or brominated hydrocarbous. Hydrogen and halogen are always taken from contiguous carbon atoms. Alcoholie potash sometimes displaces halogen atoms by cthoxyl: CH_Cl.CO_K + KOH + EtOH = CH₂(OEt).CO₂K + KCl + H₂O. γ-Chloro-acids are converted by neutralisation with potash into lactones (q. v.). When the halogen is in place of hydrogen in the benzene nucleus, it cannot be turned out by aqueous potash unless a nitrogroup is also present. Thus chloro-benzene is not affected by potash, while o- and p- chloronitro-benzenes are converted into nitro-phenols.

When phonol is boiled with chloroform and NaOHAq, oxy-benzoic aldehyde results (Tiemann a. Reimer 'sreaction): C₆H₂ONa + 3NaOH + CHCl₂ = C₆H₄(ONa)COH + 3NaOH + 2H₂O (B. 9, 824).

By the same method the group CHO can be introduced into many derivatives of phenol

(p. 109).

If tetrachloride of carbon be used instead of ohloroform, carboxyl enters the phenol, forming a carboxylic acid: C₄H₄ONa + 5NaOII + CCl₄ = C₅H₄(ONa).CO.Na + 4NaCl + 3H₂O. Alcoholic potash sometimes acts as a reducing agent (p. 99, 1. 42).

Potash-fusion (or soda fusion).

Converts aromatic sulphonates into phenols:
 C₆H₃.SO₃K + KOH = C₆H₃OH + K₂SO₃.

Displaces halogens by hydroxyl:

C₄H₄Cl.CO₂K + KOH = C₆H₄(OH).CO²K + KCl. However, owing to the high temperature required, a subsequent migration of the hydroxyl

sometimes takes place. Thus when any halogen benzene sulphonate or halogen phenol is fined with potash at 235°-270° resorcin is produced. 3. Converts carboxylates into hydrocarbons: CH₄.CO₂K + HOK = CH₄ + CO₂K₂. Soda-lime, lime, or baryta may also be used for this purpose. 4. Converts the higher fatty aldehydes and aromatic aldehydes into alcohol and salt of the acid: 2Ph.CHO + KÖH = Ph.CO₂K + Ph.CH₂OH. In other cases also, potash acts by oxidising one portion of the substance and reducing another. Thus glycerin distilled with potash gives (a) by reduction, propylene glycol, (b) by oxidation, potassic acetate and formate. Similarly anthraquinone sulphonate gives (a) by reduction, anthracene, (b) by oxidation, alizarin. 5. Splits up unsaturated acids at the point of non-saturation into two salts:

CH₂.CH:CH.CO₂H + 2KOH = 2CH₂.CO₂K + H₂. 6. Resins usually give p-oxy-benzoate protocatechuate, and phloroglucin.

ALKALOIDS.—The term alkaloid was first applied to any organic base. It is now usually restricted to organic bases that are of vegetable origin and produce marked toxicological effects. Thus such bodies as ethylamine, asparagine, and leucine, are not usually classed as alkaloids. All the alkaloids contain nitrogen, and all except conline, nicotine, and sparteino contain oxygen. These three alkaloids are volatile, the others are fixed. The vegetable alkaloids are ammonia-, not ammonium-, bases, that is, they combine with HCl without elimination of H₂O. The following alkaloids have been described:

From Achillea Mosehata: achilloine, moschatine.

From Aconitum Napellus, ferox, &c.: aconitine, piero-aconitine, pseudo-aconitine, japa-conitine, lycaconitine, myoctonine.

From Æthusa Cynapium: cynapine.

From Agaricus: agarythrine.
From Alstonia constricta: alstonine, porphyrine, alstonidine, alstonicine.

From Arariba rubra: aribine.

From Artemisia abrotanum: arbrotine.
From Aspidosperma: aspidospermine, aspidospermatine, aspidosamine, hypoquebrachine, quebrachino, quebrachamine, paytine, paytamine.

From Angustura bark: cusparine, gasipeine. From Atherosperma: atherospermine. From Atropa: atropine, hyosoyamine, by-

oscine, belladonine.
From Baccharis: baceharine.

From Baptisia tinctoria: unnamed.

From Bebeeru: beberine.

From Berberis: berberine, oxyacanthine, hydrastine.

From Buxus: buxine, buxidine.

From Calabat beans: physostigmine or eserine.

From Capsicum: eapsicine.

From Cannabis indica: an unnamed alkaloid (M. Hay, Ph. [3] 13, 998).

From Chelidonium: ohelerythrine, oheli-donine.

From Cinchona: quinine, cinehonine, conquinine, quinicine, homoquinine, hydroquinidine, cinehonidine, aricine, cusconine, enschnidine, cineholine, cuscamine, onscamidine, quinamine, cinehamidine, oinchotine, hydrocinchonine, con-

quinamine, hydroquinine, dicinchonine, diconquinine, javanine, paricine.

From Coca leaves; cocaine, ecgonine, hygrine.

From Cocoa beans: theobromine. From Coffee berries : caffeine. From Colchicum: colchieine.

From Conessi bark: conessine.

From Conium: oofline. From Corydalis: eorydaline.

From Crossoptera: crossopterine.

From Curare: curarine. From Cytisus: cytisine.

From Delphinium: delphinine, delphinoidine, delphisine, staphisagrine.

From Dita bark : ditamino or ditaine, echitamine, echitenine.

From Duboisia: duboisine or hydrocyamine.

From Ergot: ergotine.

From Erythrophleum: erythrophleine.

From Esenbeckia: csenbeckine. From Frazinus americana; an unnamed

alkaloid (F. B. Power, Ph. [3] 12, 812). From Fumaria: fumarine.

From Gelsenium: gelsenine.

From Geselmium: geselmine. From Glaucium: glaucine, glaucopicrine.

From Harmala: harmaline, harmine. From Humulus lupulus (Hops): lupuline

(hopeine), neurine. From Hymenodictyon: an unnamed alkaloid.

From Ipecacuanha: emetine. From Isopyrum: isopyrine, pseudo-isopyrine.

From Lobelia: lobeline.

From Lotur bark: loturine, colloturine, loturidine.

From Loxopterygium: loxopterygine. From Lupinus: lupinine, lupinidine.

From Lycopodium: lycopodine. From Macleya: mackleyine, sanguinarine.

From Menispermum: menispermine.

From Mustard: sinapine.

From Nicotiana tabacum: nicotine.

From Nymphaa alba: an unnamed alkaloid (Gräning, B. 16, 969).

From Olcander: oleandrine.

From Opium: morphine, codeine, thebaine, papaverine, narcotine, narceine, hydrocotarnine, pseudomorphine, codamine, laudamine, laudanosine, meconidine, lanthopine, protopine, eryptopine, crytopine, oxynarcotine.

From Papaver rheas: rheadinc.

From Papaver somniferum: v. OPIUM.

From Pennius: boldine.

From Piper nigrum (Pepper): piperine. From Perciro bark: geissospermine, percirine.

From Pilocarpus leaves: pilocarpine, jaborine, pilocarpidine.

From Pomegranate bark : pelletierine.

From Poppy: rheadine. - Opium Poppy v.

From Ratany root: ratanhine.

From Ricinus (castor-oil plant): ricinine.

From Salamanulra: samandrine.

From Saphora: saphorine. From Sinapis: sinapine.

From Spartium : sparteine. From Strophantus: strophantine.

From Strychnos: strychnine, brucine.

From Thalictrum: thalictrine.

From Taxus: taxine.

From Tea leaves ; caffeine. From Tobacco: nicotine.

From Trigomella: trigomelline, neurine.

From Veratrum: veratrine, veratridine, cevs. dine, cevadilline, jervine, rubijervine, pseudojervine, veratralbine.

From Vetch: vicine.
Formation of alkaloids in plants. Most of the above alkaloids are pyridine derivatives. They are probably produced by the action of ammonia or amido compounds upon non-nitrogenous bodies. Pechmann a. Welsh (B. 17, 2384) consider that the non-nitrogenous bodies are such acids as meconie, chelidonic, and cumalic, which are probably furfurane derivatives. These three acids are converted by ammonia into exypyridine carboxylic acids. Cumalic acid is formed artificially from malic acid by action of cone. H2SO4; and it is probable that the two other acids are also formed by condensation of simpler acids.

V. Meyer has suggested that hydroxylamine hy acting upon aldehydes may also play some part in the production of the nitrogenous con-

stituents of plauts.

Extraction: The tissue is extracted with dilute acid and the extract ppd. by ammonia, potash, soda, lime, or magnesia. Volatile alkaloids are then distilled, fixed alkaloids are crystallised from a suitable solvent. The extraction of alkaloids from animal matter, as in cases of poisoning, is described in the next article.

Reactions .- 1. Sodie phosphomolybdate added to solutious acidified with nitric acid gives, in the cold, a yellowish-white flocculent pp. Aniline, the alkylamines, and quinoline, as well as silver, mercurous, and lead, salts are also ppd. by this reagent (Somenschein, A. 104, 45). recover the alkaloid, the pp. is boiled with baryta, when volatile alkaloids distil over. The residue is saturated with CO, evaporated to dryness and extracted with alcohol. Somenschein's reagent is prepared by dissolving yellow ammonie nitro-molybdate in Na₂CO₃Aq, drying and strougly heating; if reduction of molybdic acid take place, the product is moistcued with HNO. and again heated. It is then heated with water, nitric acid is added, and the liquid diluted until 10 parts of the solution contain 1 part of solid residue.

2. Phosphotungstic acid may be used instead of phospho-molybdic acid (Scheibler, Fr. 12, 315; J. 1860, 157). The reagent, which is a mixture of sodic tungstate and phosphoric acid, is added to solutions acidified with H.SO. Phosphoantimonic acid got by dropping antimonic chloride into aqueous phosphoric acid, precipitates morphine, narcotine, and nicotine, but not

atropine (F. Schulze, A. 109, 177).

3. Potassio-mercurie iodide produces floceu-lent yellowish-white pps., insoluble in acids and in dilute alkalis, slightly soluble in excess of the reagent, easily soluble in alcohol, and generally also in other (F. Mayer, J. 1863, 703; A. 133, 236; De Vrij, J. 1867, 602). Theobromine, caffeine, glucosides, carbohydrates, and organie acids give no pp. with Mayer's solution. Albuminous and gelatinous substauces, in presence of free acid (but not in alkaline solutions) give sticky pps. (Valser, Fr. 2, 79). To separate the alkaloid from the pp., the latter is triturated

with SnCl, and excess of aqueous KOH; this reduces the mercury to the metallic state, and the base is then extracted by its proper solvent.

Mayer's solution contains 18.5 g. mercuric

chloride and 49 8 g. potassio iodide per litre.
4. Potassio-bismuthous iodide is prepared by dissolving Bi(OH)2NO, (80 g.) in HNO, (200 c.c. of S.G. 1'18) and adding a conc. solution of KI (272 g.). The solution is cooled until KNO, crystallises, and the mother liquor is then diluted to a litre (Dragendorff, Fr. 5, 406; Kraut, A. 210, 310). The solution is added to the alkaloid dissolved in dilute H.SO, or HI. Double iodides of the alkaloid and of bismuth are ppd. The alkaloid can be recovered by decomposing these double iodides with aqueous NaOlI, and extracting with a proper solvent.

5. Potassio-cadmic iodide forms flocculent pps. when added to solutions of alkaloids acidulated with H2SO4. The pps. soon become crystalline; they are soluble in alcohol, insoluble in other. They dissolve in excess of the reagent. The alkaloid can be recovered by treatment with NaOHAq and a solvent (Marmé,

Bl. [2] 9, 203).

6. Potassio-platinic iodide and potassio-auric iodide also pp. alkaloids (Selmi, G. 5, 255). These solutions are prepared by adding K1 to solutions of PtCl, or AuCl, until the pp. first formed is redissolved. The platinum salt gives, in acetic acid solution, a black pp. with nicotine, but nono with conessine; it also gives a winered pp. with solanidine but none with solanine. The gold salt gives, on evaporation, arborescent crystallisation with nicotine, but only oily drops with conline.

7. A solution of iodine (1 pt.) in Kl (1 pt.) dissolved in water (100 pts.) gives brown, often crystalline, pps. of the periodides. These polarise light like tourmaline. The alkaloids can be recovered by treating the pp. with SO2Aq.

8. Animal charcoal removes most of the alkaloids from aqueous solution. The alkaloid can thon be extracted from the charcoal by a suitable solvent (Graham a. Hofman, C. J. 5,

9. Picric acid pps. many alkaloids, even in presence of a large excess of H2SO4. Morphine, caffeine, and glucosides are not so ppd. reagent pps. English but not German preparations of atropine (Hager, Fr. 9, 110).

10. Tannin gives a white or yellowish-white The salts of morphine, with the exception of the acetate in strong solution, are not ppd. by tannin. The alkaloids can be recovered by treating the pp. with lime.

11. Platinic chloride gives, in conc. solutions, a yellowish-wifte or yellow pp. Chloride of gold does the same (cf. Coninck, Bl. [2] 45, 131).

12. Sodic nitroprusside usually forms oily drops of the nitroprusside, which crystallises on standing (Horsley, C. N. 5, 355; E. W. Davy, Ph. [3] 11, 756).

13. The electrolysis of solutions of salts of alkaloids has been studied by Bourgoin (Bl. [2]

12, 438).
14. The alkaloids are ppd. by sodium salts of glycocholic, hyoglycocholic, and taurocholic acids. The pps. appear to be acid salts of the alkaloids (W. F. de l'Abre, C. C. 1872, 231).

15. Hydric sulphide passed into alcoholic

solutions of alkaloids forms compounds containing sulphur (Palm, J. 1863, 483; E. Schmidt, B. 8, 1267).

16. The absorption-spectra of various alkaloids have been mapped by A. Meyer (P. [8] 18,

17. A solution of iodine monochloride in HCl added to a solution of an alkaloid in HCl gives a pp., usually yellow that sparingly soluble (Tildeu, C. J. 21, 145; Dittmar, B. 18, 1612; Ostermayer, B. 18, 2298). According to Dittmar, if the alkaloid contain one pyridins ring, the pp. is of the form XICI; if it contain two pyridine groups it is of the form YI Cl. However, NEt, Cl. NEt, HCl, caffeiue hydrochloride, and pyrrol, all give pps. though they contain no pyridine ring. while B-oxy-quinolines, and tetra-hydroquinolins give no such pps. The pps. may also be got by using a mixture of Kl, KNO,, and HCl, instead The chlore-iodides are converted by of ICl. excess of chlorine into unstable compounds XICl. Aminonia converts the chloro-iodides into iodoamides, X1NH,; these are dark-green or darkred unstable bodies, insoluble in water, but converted back into the chloro-iodides by HCl, and decomposed by boiling with alcohol according to the equation:

 $6XINII_{2} - 3X + 3XI_{2} + 4NH_{3} + N_{2}$ These iodo-amides are also produced by the combination of the alkaloids with iodide of

nitrogen.

Tests for alkaloids.—The above reactions are general. The following tests may be used in distinguishing the alkaloids from one another:-1. The alkaloid is sublimed. Sublimation begins below 100° in the case of eaffeine, theobromine, and cantharidine; between 150°-200° in the case of strychnine, morphine, and pilocarpine (Wynter Blyth, C. J. 33, 318). The following give no sublimate, but melt: (a) bolow 100° hyoseyamine, atropine; (b) between 100°-150° papaverine; (c) above 200° solanine. The sublimate is then examined microscopically (Helvig, Fr. 3, 43; Deaue a. Brady, C. J. 18, 34). 2. Conc. H.SO, produces colours with certain alkaloids, e.g. a blood-red colour with thebaine and a crimson with veratrine. -3. Nitrio acid usually produces a yellow solution, but morphine and brucine give a red colour.-1. Sulpburicacid containing a little molybdic acid, so-called sulphomolybdic acid, gives a violet colour with morphine, and characteristic colours with other alkaloids.—5. Erdmann's solution is prepared by adding HNO₃ (6 drops of S.G. 1.25) to water (100 c.c.) and adding ten drops of the diluted acid to 20 c.c. of conc. H_2SO_4 . This solution gives a blue colour when warmed with solutions of codeine, and characteristic colours with other alkaloids .- 6. Chlorine water, followed by ammonia, gives a green colour with quinins, a rad colour with narceine, and an orange colour with narcotine .- 7. A mixture of an alkaloid (1 pt.) with sugar (7 pts.) often gives colours with conc. H.SO. Morphino and codeino give a purple, aconitine a rose-red (R. Sehneider, P. 147, 128). -8. H.SO, and a little Cc.O, give with strychnine a fine blue colour; with brucins, orange; narcotine, brown, cherry-red, finally wine-red; morphine, olivo-brown, finally brown; codeine, olive green, finally brown; quinine, pale yellow; einchouine and calleine, no colour; veratrine. atropine, solanine, emetine, brown; colchicine, green becoming brown; papaverine, almost black (Sennenschein, B. 3, 652).

Optical Properties.—When the solution of

an alkaloid affects a ray of polarised light the specific rotatory power of solutions of its normal salts is independent of the nature of the acid if the alkaloid is mono-acidic and the salt is not decomposed by water, but if the alkaloid is diacidic the basic salts usually rotate much less than the normal salts (Oudemans, R. 1, 18).

ALKALOIDS, CADAVERIC v. Promaines.

ALKALOIDS, POISONOUS, Detection and Estimation of .- This article will be directed to the simple detection of the chief alkaloids; but incidentally their quantitative estimation will be touched upon. The detection and complete recognition of an alkaloid by chemical tests is often a matter of great difficulty, even in the case of some well-known and potent alkaloids. Indeed, in some cases these difficulties are at present insurmountable by chemical means alone, and physiological experiments have to be called in aid. The obstacles to the recognition of these bases, when several are present, is still greater. Indeed, it may be stated that the completo separation of a mixture of commonly occurring alkaloids is a problem still awaiting solution. Tho toxicologist has too often to content himself with the identification of one or two alkaloids in organic mixtures, and the determination of their aggregate quantity, being unable to ascertain their individual amounts.

History .- The earliest methods devised for the detection of alkaloidal poisons in forensic research were those found effective for the separation of the vegetable alkaloids from the other matters with which they are found associated in nature; and modifications of these methods are even now employed for this pur-The material to be operated on was extracted with diluted acids, aided by gentle heat, gummy and other substances removed by lead acetate, the excess of lead ppd. with hydrogen sulphide, and the alkaloid obtained as a salt-generally an acetate-in a greater or less degree of purity by evaporation of the solution. This and other crude processes were mostly abandoned when Stas (A. 84, 379) published his classical paper on the separation of alkaloids from organic mixtures, and devised a new and refined process, which was subsequently modified and improved by Otto (A. 100, 39), Dragendorff (Gerichtl. chem. Ermit. v. Gift., 1876), and others. In one form or another, this process, generally known as that of Otto-Stas, is still the one most generally employed in a, generally known as that of Otto-Stas, forensic analyses, though Sonnonschein (A. 105, 45), Selmi (C. J. 1877, 93), and more recently L. Brieger (Die Ptomaine, Pt. I., 1885; Pt. II., 1885; Pt. III., 1886) have each employed different but less refined methods for the general separation of the organic bases from the matters with which they are commonly associated.

Methods of Procedure. - If an apparently fairly pure solid body has to be examined, e.g. a crystalline medicinal powder, its alkaloidal nature inay be demonstrated by ascertaining in the ordinary way that it contains both organic carbon and nitrogen; by its sparing solubility in aqueous alkaline, and its ready solubility in

aqueous acid, solutions; and by adding to the acid solution reagents that react with the alkaloids as a class. The alkaloid will usually be ppd. from its acid or aqueous solution, if this he not too dilute, by caustic and by carbonated alkalis; and will appear either in the form of oily droplets (liquid and volatile alkaloids), or as an amorphous pp., becoming crystalline on stand. ing. If the pp. redissolves in excess of the caustic alkali, as in the case of morphine, it will again scparate when the alkali becomes carbonated, as by exposure to the air. Since none of the alkaloids are altogether insoluble in water, no pp. may form in very dilute solutions, and yet an alkaloid bo present. There are, however, certain group reagents that pp. the alkaloids from their barcly acid solutions, even when these are highly dilute; and these reagents are generally employed where the presence of an alkaloid is suspected. Such goneral reagents are the following; but it must be borne in mind that as any one of them may fail to give a precipitate with a given organic base, two or more of them must be used, under appropriate conditions, in order to prove or disprove the presence of an alkaloid in the solution to be tested .- 1. A weak solution of iodine in potassium iodide. This reagent gives a more or less coloured pp. with extremely dilute solutions of most of the vegetable alkaloids.—2. Bromins water yields similar pps., but has the disadvantage of yielding pps. with the phonols also.—3. Tannin pps. most of the vegetable alkaloids from their not too dilute solutions .-4. Mercuric chloride in aqueous, and also in alcoholic solution, is a valuable pptant., and is especially useful in the separation of the organic bases resulting from putrefaction (ptomaines), for the separation of which the Otto-Stas method to be presently described is inadequate (L. Brieger, op. cit.) .- 5. Potassio-mercuric iodide solution is perhaps the most generally useful pptant, of the alkaloids; and from the ppts. thus produced the alkaloids may be recovered in a high state of purity by trituration with stannous chloride and solution of NaOH, and extraction of the alkaloid thus liberated, with ether. - 6. Phosphomolybdic acid in nitric acid solution pps. the alkaloids from acid solutions. The alkaloids may be recovered from these pps. by decomposing them with barium hydrate, and either distilling off the alkaloid (volatile alkaloids), or after removing the baryta by means of a stream of carbon dioxide, subsequently extracting the alkaloid with absolute alcohol.-7. Picric acid is also a useful pptant., and from the pps. thus produced the alkaloids may be separated by acidification with HCl and agitation with other. - Many other general pptants, of the alkaloids have been proposed, but the above fulfil almost every useful purpose; and on the ppn. of organic natural bases by alkalis, and their resolution and extraction by other and other special solvents - or on their removal from organic solutions by one or other of the above pptants.—are based the most approved general methods for the separation of the

poisonous alkaloids in forensic analysis.

The method most generally employed for the extraction of the vegetable alkaloids from admixture with animal matters is that originally devised by Stas for the separation of nicotine in

he course of a forensie analysis, and now known | rith modifications as the Otto-Stas method. The writer of this article has introduced further nodifications which are embodied in the followng description, and have been found by him necessary in those cases where unstable and easily hydrolysed alkaloids are to be sought for, such as morphine, which is readily decomposed by heating its acidified solutions, and aconitine, which is unstable in alkaline and especially in ammoniacal solutions. In all cases the method, which is a quantitative one, is greatly dependent for success upon the care with which the preliminary operations are conducted.

The organic material to be operated upon, if solid, is brought into as minute a state of division as its nature permits, and is then digested with twice its weight of rectified spirit of wine at a temperature of about 35°. Liquids are also treated with twice their volume of rectified spirit. Redistilled methylated spirit of wine may be used for these operations. After eeveral hours' digestion the liquid is poured off from the deposited solids, and the digestion is repeated with a fresh quantity of spirit. This is again poured off, and mixed with the previous alcoholic infusion. If solid matter, e.g. liver, is operated on, the liquid is squeezed from the solid portion at each digestion in a piece of fine cambric which acts as a crude filter; and the liquids thus obtained are added to the other alcoholie liquids. After two or mere digestions, according to the nature of the organic matter, the andissolved portions are subjected to a new digestion, also at 35°, with spirit faintly acidified with acetic acid. Some recommend tartaric in preference to acetic acid, but this is objectionable, when, as is usually the ease, morphine has to be sought for: others use oxalic acid, but this reid may have to be sought for as well as the alkaloids. Enough acid must be added to keep the liquid just perceptibly acid, excess being avoided. After a prolonged digestion with the acidified alcohol, this is poured off, the solids squeezed, and the digestion repeated, but this time with macidified spirit. A final digestion may be required, the rule being to continue the exhaustion with spirit of wine so long as any colour is imparted to this. The alcoholic liquids obtained before acidification after mixing are momentarily and rapidly raised to a temperature of 70°, cooled, and the insoluble residue filtered and washed with spirit; and those obtained with and after the use of acctic acid are similarly treated. But the two liquids, the unacidified and the acidified are not mixed till a later stage is reached. In this way, by keeping the liquids separato, danger of hydrolysation of unstable alkaloids is as far as possible avoided. The alcoholic infusions are now evaporated at a temperature never exceeding 35° to the consistency of a syrup. It is advisable during the course of the evaporations to neutralise a portion of the free acid with caustie soda from time to time, so as to keep the liquids just per-ceptibly acid. The evaporations are easily effected with tolerable rapidity by placing the liquids in challow basins supported on large beakers some inches above the floor of an ordinary

advantages of this arrangement of the author's are, that overheating is avoided, evaporation is more rapid than by any other method, and all creeping of the liquids up the sides of the basins is obviated. This course of procedure is greatly preferable to distilling off and recovering the alcohol, as usually recommended. The syrupy liquid is now drenchod with about 30 c.c. of absolute alcohol, with constant stirring or grinding in a mortar; the alcohol is poured off from the pasty mass which usually separates, and replaced by successive quantities of 15 c.o. alcohol so long as a colour is imparted to this. The alcoholic liquids are mixed, filtered, the filter washed with alcohol, and the filtrate evaporated in the oven, as before, at a temperature not exceeding 35°. The syrupy residues—that from the plain and that from the acidified spirit of wine-are diluted with a small quantity of water, filtered, the filters washed with water, and They should, together, the filtrates mixed. measure 15-20 e.c. The liquid is introduced into an accurately stoppered tube, partially neutralised, if necessary, with caustic soda, taking care, however, to leave it slightly acid. If the method laid down has been scrupulously followed, we have now a liquid containing the whole of the alkaloids, and free from albuminoids. aqueous and faintly acid liquid in the tube is now covered with twice its volume of washed ether. and the whole is mixed by gently and repeatedly inverting the tube, care being taken not to emulsify the mixture by any violent agitation. The ether is allowed to separate, and this is favoured by giving an occasional sharp rotatory shake to the tube. The supernatant ether is then pipetted off, and replaced by a new and smaller quantity of ether. The tube is again shaken; and the operation of extraction with ether is continued till a few drops on evaporation leave no residue. Four or five extractions will generally suffice. The othereal solutions as they are pipetted off are successively washed by vigorous shaking in a second stoppered tube with 5 e.c. water to which a few drops of dilute sulphuric acid have been added. The ether on evaporation may yield an oily residue which may be reserved for further examination and for physiological tests. But the acid liquid subjected to ethereal extraction will still contain nearly all the alkaloids, as the acid salts of these are mostly practically insoluble in ether; but the salts of some of the alkaloids being perceptibly soluble in ether may be met with in the 'acidother 'extract. The acid aqueous solution, and the acidulated water with which the ether has been washed, are mixed, alkalised with sodium carbonate, and again exhausted four or five times as before with washed ether; only in this case the first exhanstion is made with a mixture of 1 vol. chloroform and 3 vol. ether, and the final extractions with other alone. These successive ethereal extracts are washed in a tube by shaking anew with 5 c.e. water. They are then transferred to a third and finally to a fourth tube, the first containing 10 e.c. water acidulated with a few drops of sulphurie acid, and the last containing 5 c.c. water alone, and agitated. By these operations the alkaloids are first liberated from oven, which is heated by a gas flame playing on their salts by the alkali, then transferred to the the top. The door is kept a little ajar. The ether-chloroform in which they are coluble; then

again converted into sulphates, which, being insoluble in other and chloroform, again pass into the acid solutions, impurities being left behind in the ether. We have now again the alkaloids in acid solution, but in a much purer state than before. The acid liquid and the final washwater are mixed, washed with a little ether once or twice, then re-alkalised with sodium carbonate, and well re-extracted with chloroform-ether and ether. These ethereal solutions are washed with water barely alkalised with sodium carbonate, then filtered through a dry filter, and evaporated to dryness in the oven below 35° in tared glass basins about eight centim. in diameter. Once dry, the residue may be transferred for a few minutes to the water-oven, dried at 100°, and weighed after cooling over sulphuric acid. This weight fairly represents that of the alkaloids. It is well before evaporating the bulk of the solution to evaporate a few e.c. only; if an oily odorous residue be left the presence of a volatile alkaloid is indicated; the evaporation is then modified by mixing the ether-ehloroform with so much ether previously acidulated by agitation with a strong solution of hydrochloric acid as is necessary to render it acid. In this case it is not the free alkaloids, but their nonvolatilo hydrochlorides which are left and weighed. The residue may therefore be dissolved in water and subjected to appropriate testsfirst for the alkaloids generally (vide aute), and secondly specifically for the volatile alkaloids. But if the solid free alkaloid has been obtained it must first be converted into a hydrochloride by moistening it with a very slight excess of very dilute hydrochloric acid, and evaporating to dryness in vacuo over sulphuric acid. The residue may then be dissolved in water and subjected to tests, which may be obtained from the ethercal extract either before or after conversion into hydrochloride and solution of this in water.

Morphine is practically insoluble in ether except immediately after ppn. from its solutions, hence if this alkaloid were present in the matters submitted to examination, but little of it would be removed by the chloroform and other, more especially if, as is advisable, the agitation with these solvents was not executed inomediately after the addition of sodium earbonate. In order to obtain the morphine, the first alkalised solution from which the other alkaloids have been removed must be re-extracted a few times with a well-washed mixture of equal volumes of acctio ether and ethyl ether, which is preferable to amyl-alcohol, and in which mixture morphine is soluble. The mixed ethers are washed with a little water, filtered through a dry filter, and evaporated just as the chloroform-ether was evaporated for the other alkaloids. The residue is usually not pure morphine, as acetic ether takes up other non-alkaloidal bodies, but these . do not usually interfere with the morphine

G. Bragendorff (Gerichtl. chem. Ermit. v. Gift., 1876, p. 141) has devised a process which proceeds upon the same general lines as that of Stas, but is much more elaborate. It requires, moreover, a higher temperature for the preliminary evaporation. The finely divided substance, if solid, is digested for several hours at a temperature of 40°-50° with dilute sulphurie

soid-about 2 p.c. by volume of the soid. Ideaids are acidulated with the same proportion of acid. The digestion is continued for several hours. and the mixture is then pressed, and filtered. The operation of extraction with dilute sulphuric acid is repeated two or three times, 100 o.o. of liquid being a convenient quantity for each extraction. The mixed filtrates, containing the alkaloids, are partially neutralised with magnesia. and carefully evaporated to a syrnp at a temperature much below 100°; but never to dryness. It is certain that in this operation some alkaloids may be destroyed, and it is also asserted that basic bodies are fermed by the decomposition of the albuminoids. A useful modification (L. Brieger, op. cit.) is to partially neutralise the liquid from time to time during the course of the evaporation, so that it is never more than very slightly acid in its reaction to litmus. The syrupy residue from the evaporation is mixed with three or four times its volume of rectified methylated spirit of wine, and a few drops of sulphurie acid, and allowed to digest at about 30° for twenty-four hours. The insoluble matter is separated by filtration and washed with spirit. and the filtrate and washings are distilled in a retort so as to recover the alcohol. The aqueous residue in the retort is diluted with water to 50 o.c., filtered, and introduced into a stoppered tube and exhausted successively with petroleum ether, benzene, and chloroform, 20-30 e.e. of each at a time, in the manner in which exhaustions are made with ether in Stas's process. The aquoous solution is then made alkaline with ammonia, and again exhausted successively with petroleum-spirit, benzeue, chloroform, and amylalcohol. On evaporation of the respective solvents (consult what has been said under Stas's process as to volatile alkaloids) alkaloidal residues are obtained, which when taken up with water, either with or without previous conversion into hydrochlorides as necessity demands, may be submitted to appropriate tests for the alkaloids, and specially for the various suspected alkaloids.

Among the commoner alkaloids, and poisonous neutral substances:—

Petroleum ether removes from the acid aqueous solution:—some piperine, resins such as capsicin, camplior, and plienol.

Benzene further removes from the acid solution:—more piperine, easterne, celchicine, santonin, digitalin, cubebin, colocynthin, cloctin, pieric acid, elaterin, and cantharidin.

Chloroform, again, removes from the acid aqueous solution: — cinclionine, theobromine, papaverine, narceine, jervine, more digitalin, pierotoxin, smilacin, and senegin.

On rendering the solution alkaline with ammonia:—

Petroleum spirit removes from the alkaline solution:—strychnine, brucine, quiuine, veratrine, aconitine, emetine, and the volatile alkaloids coniine, nicotine, lobeline, and trimethylamine (from putrefaction), the pimento-alkaloid, and aniline. If the presence of aconitine or emetine be suspected, the operation must be performed quickly, since these alkaloids rapidly decompose in alkaline solutions.

Benzene further removes from the alkaline solution:—atropine, hyoscyamine, physostig-

mine (eserine), thebaine, codeine, narcotine, and additional quantities of stryohnine, bruoine, quinine, cincinonine, veratrine, aconitine, and emetine.

Chloroform, again, removes from the alkaline solution:—some morphine, and additional quantities of cinchonine, narcoine, and papaverine.

Amyl-alcohol finally removes from the alkaline solution:—morphine, narccine, and somo neutral bodies, such as salicin.

Not all the substances enumerated above are poisonous; but they are bodies that may be present in medicinal mixtures, and hence are likely to come under the notice of the toxicologist in foreusic analyses.

Selmi proposed another method of extracting the poisonous alkaloids, and applied it to the extraction of the ptomaines (G. 6, 153; C. J. 31, 93). The viscera are exhausted with alcohol and dilute sulphuric acid. This acidified alcoholic extract is filtered and evaporated at a temperature of 65°, again filtered, and evaporated to a syrup. The residue is tuken np with water, filtered, the filtrate treated with basio lead acetate, and the mixture exposed to the air for twenty-four hours. It is then filtered, the excess of lead removed by hydrogen sulphide gas, and the filtrate concentrated. This is then repeatedly extracted with ether. The ethereal solution is then saturated with a stream of dry carbon dioxide gas, which generally causes a pp. of droplets containing some of the alkaloids, and adherent to the side of the vessel. The ethereal solution is poured off, mixed with about half its volume of water, and a current of carbon dioxido is again passed for twenty minutes, which may cause the ppn. of other alkaloids not ppd. by dry earbon dioxide. Usually the whole of the alkaloids are thrown down by these means, but if not, the ethereal solution is dehydrated by shaking with barium oxide, and then a solution of tartaric acid in other is added to the clear liquid, taking care not to employ an excess of acid; any alkaloid that may remain in solution is thus thrown down. As a matter of precaution, the remains of the viscera or other matters operated on are mixed with barium hydrate and a little water, and agitated with pure amyl alcohol. The alkaloids may then be extracted from tho alcohol by shaking it in a etoppered tube with very dilute sulphuric acid.

Sonnerseliein (A. 104, 45) separates the alkaloids by ppn. with phosphonolybdic acid. In extracting the bases, the organic matter to be examined is repeatedly exhausted with very dilute hydrochloric acid, the mixed solutions filtered, and the filtrate evaporated in the oven at a temperature not exceeding 35° to a thin syrup; then diluted with water, cooled, and filtered. An excess of phospho-molybdic ucid is added to the filtrate, and the pp. is washed with water containing nitric and phosphomolybdic acids. The still moist pp. is washed into a flask, made distinctly alkaline by the addition of barium hydrate, and distilled into a bulb apparatus charged with hydrochloric acid, which absorbs ammonic and the volatile bases. The residue in the flask, containing the non-volatile bases, is freed from barium hydrate by a current of carbon dioxide, evaporated to drypess, and the

bases extracted by means of strong alcohol. The filtered alcoholic colution often yields on evaporation the alkaloids in a sufficient etate of purity to admit of their being weighed, converted into hydrochloridee, and eubmitted to tests. Sometimes, however, they must be purified by re-solution and re-crystallisation from absolute alcohol, ether, chloroform, &c.

The method of Usiar and Erdmann (A. 120, 121) is not much employed in this country, tho evaporation of amyl alcohol being a disagreeable operation; but it is nevertheless a valuable process for the separating of strychnine and morphine. It is practised us follows :- The suspected matter, if solid, is made into thin paste with water, slightly acidulated with hydrochloric acid, and digested at a temperature of 70° for an hour or two. It is then strained through a moist piece of cambric, and the solid residue on the cloth is well exhausted with hot very dilute hydrochloric acid. The combined liquids after filtration are made slightly alkalino with ammonia, mixed with clean sand, and evaporated to dryness on the water-bath. The residue from the evaporation is extracted three or four times with hot amyl alcohol and the mixed liquids are filtered. The filtrate is shaken violently in a stoppered tube with several times its volume of hot water acidulated with hydroehloric acid, which removes the ulkaloids, leaving colouring matters and fat in the alcohol. The alcohol is pipetted off, and the hot acid solution is repeatedly washed by agitation with fresh portions of amyl alcohol until all fat and colouring matter is removed, after which the clear uqueous liquid is concentrated by evaporation, made alkaline with ammonia, and well shaken with fresh hot amyl alcohol four or five times. These alcoholic liquids are mixed, filtered through a filter moistened with amyl alcohol to remove water, and evaporated in a tared basin, when the alkaloids will be left in a sufficiently puro condition to be dissolved and tested, previous to which they should be weighed. Occasionally a coloured residue is obtained which requires re-solution in aqueous acid, agitation with amyl alcohol, alkalisation with ammonia, and re-extraction with amyl alcohol, in order to obtain the alkaloids in a sufficiently pure state for testing. With morphine the process, though tedious, works well.

Scheibler's process is based upon the precipitation of the alkaloids by phosphotungstic acid, a reagent prepared by treating sodium tungstate in solution with hulf its weight of phosphorio acid, when crystals of phosphotungstic acid slowly form. These are dissolved, and the process in detail is carried on in the same manner as Sonnenschein's phosphomolybdic acid process,. substituting the phosphotungstic for the phosphomolybdic solution as a precipitant. It is sometimes recommended to precede the ppn. of the alkaloid by the addition of lead acetate to vemove colouring matter, and then to remove the excessof lead by hydrogen sulphide; but some of the alkaloid present is apt to be removed with the Nor should animal charcoal be used lead pps. as a decolorant, as this is still more effective in withdrawing the alkaloids from solution. Indeed the obstinacy with which the arkaloids adhere to animal charcoal has been ntilised by Graham, Hofmann, and Redwood as a means of separating stryohnine from beer and other liquids (O. J. 5, 173).

The following scheme serves for the detection of the more commonly occurring poisonous alkaloids. The alkaloids are brought into acid aqueous solution, and this is shaken with ether:-

I. The ether withdraws from the acid aqueous

solution, and leaves on evaporation:

1. Colchicine.-Its solution is yellow, and is turned to a violet colour by strong uitric acid. Its solution in HClAq, when made alkaline with caustio soda, develops an orange-red colour.

- 2. Digitalin. When dissolved in conc. H.SO. and a minute quantity of bromine water is added, a reddish-violet tint is produced, which, on the addition of water clisnges to a green.
- 3. Picrotoxin.—Reduces Felling's solution. The solution is made alkaline with sodium bicarhonate, and again shaken with other.

II. Ether removes from the alkaline solution,

and leaves on evaporation :

- 1. Nicotine. Oily droplets, having a tobaceolike odour. Its aqueous solution is not ppd. by chlorine water, nor does it become coloured when warmed. Warmed with hydrochloric acid the alkaloid becomes violet, and then on the addition of nitric acid orange-coloured.
- 2. Coniine. Oily droplets having a mousclike odour. Its aqueous solution is ppd. by chlorine water, and becomes coloured when warmed. Dry hydrochloric acid gas turns the alkaloid at first red, and then to a violet colour.

3. Lobeline .- Oily droplets yielding no very

definite chemical reactions.

- 4. Brucine.-Turned rosy-red by strong sulphuric acid not quite free from oxides of nitrogen. The alkaloid is reddened by strong nitrie acid, and the red solution changes to a bluish violet on the addition of a solution of stannous chloride.
- 5. Strychnine.—No coloration on the addition of strong H₂SO₂. On the further addition of solid potassium dichromate, MnO2, or PbO2, a violet-blue coloration is itumediately produced, passing gradually into a cherry-red, the colour only slowly disappearing.

6. Narcotine. - It become first yellow, then bluish-violet when warmed with strong sulphuric soid. Its solution in strong H.SO, becomes red on the addition of a trace of nitric acid. Sulphomolybdie acid turns the alkaloid green.

7. Veratrine. - With strong sulphurie acid it becomes yellow, orange, and finally cherry-red. Its solution in cold concentrated hydrochloric acid is colourless, but gradually changes to a

deep red when boiled.

8. Jervine. The salts of this alkaloid, except the acctate and phosphate are only very sparingly soluble in water and acid solutions. Its solution in acetic acid is ppd. by nitric acid and by

potassium nitrate.

9. Atropine. - An odour of hawthorn is doveloped when the alkaloid is warmed with strong sulphuric acid and potassium dichromate; and the solution becomes green from roduction of chromic acid. Evaporated to dryness with fuming nitric scid, and the residue tonched with an alcoholic solution of potash, a fine purple colour is produced.

physiological properties. The chemical tests for pure aconitine are not characteristic.

11. Gelsemine .- With strong sulphurio acid and potassium dichromate, a roddish-purple or cherry-red colour is devoloped, quickly passing int# bluish-green or blue.

12. Physostigmine. - Its solutions, whether acid or alkaline, become reddish on exposure, and this colour is discharged by sulphnrous acid and the thiosulphates. Treated with sulphurio acid and bromine-water, it yields a brown-red colour.

III. There remain in the alkaline aqueous solution :-

Morphine, curarine, and cytisine. The first may be separated by shaking with acetic ether, and is identified by the usual tests-nitrio acid, ferrio chloride, iodio acid and starch, and sulphomolybdie acid. Curarine may be ppd. by phosphomolybdic acid after acidification with nitric acid. The pp. is decomposed by barium hydrate, and the alkaloid extracted by absolute alcohol (v. Sonnenschein's process, ante); it gives a reddish colour with sulphuric acid, and reacts somewhat like strychnino with sulphurio acid and potassium diehromate. Curarine, however, is not precipitated from its solutions by potassium diehromate.

Cytisine yields no definite chemical reactions.

The following are the chief trustworthy tests for the more commonly occurring poisonous alkaloids, &c. :-

Aconitine. - This alkaloid, as well as the elosely allied alkalcids pseud-aconitine and japaeonitine, when pure, yield no characteristic chemical reactions. The colour reactions with sulphuric aud phosphoric acids that have been. described by authors are untrustworthy, and due to impurities.

Apomorphine .- Its salts turn green on exposure to light and air, and its solutions on boiling. With sodium biearbonate its solutions yield a pp. which turns green on standing, and forms with ether a purple, and with ehloro-form a violet, solution. Its solutions strike a red colour with ferric chloride and with nitric acid.

Atropine.-When warmed with strong sulpliuric acid-or, more quickly, when evaporated to dryness with baryta-water, and the residue heated-an odour of stale hawthorn flowers is developed.

Brucine.-It is turned of a blood-red colour by nitric acid; and the red solution becomes violet on the cautious addition of a solution of stannous chloride. It yields an orange-red with sulphomolybdic acid; and with sulphuric acid and potassium dichromate a deep orange-red colour.

Caffeine. - Crystallises in silky needles. Evaporated to dryness with dilute hydrochloric acid and a fragment of potassium chlorate, a pink residue is left, which turns violet on the addition of ammonia.

Cinchonine.-It is difficult to get any characteristic reaction for this base. Its sulphate is soluble in chloroform, and is non-fluorescent-characters which distinguish it from quinine.

Cocaine.—Is not coloured by concentrated acids. When evaporated to dryness with alco-10. Aconitine can only be identified by its hollo potast, the residue when warmed with dilute sulphuric acid evolves an aromatic odour

of benzoio acid ... Nitrio acid strikes a violet colour, which on the addition of sodium hydrate changes to a fine orange. Strong sulphurio acid dissolves the alkaloid to a greenish yellow colonr, and on the addition of a drop of dilute nitrio acid a play of colours, beginning with violet, is manifested. The subsequent addition of caustic soda yields a fine rose tint.

Coniine.-Oily, and of a mouse-like odour. Its dilnte aqueous solution fumes with strong hydrochlorio acid, and after acidification gives no pp. with bromine-water until supersaturated with sedium hydrate, when a white pp. forms.

Curarine. — Is turned blue violet when

touched with sulphurie acid and potassium dichromate added; but the colour is more persistent than in the case of strychnine.

Cytisine .- This alkaloid is insoluble in ether, benzene, chloroform, and carbon disulphido. It dissolves in sulphuric acid without colour, but the subsequent addition of nitrio acid produces an orango coloration. Nitric acid dissolves cytisino without colour, but on warming tho mixture becomes orange-red.

Emetine is said to strike a blood-red colonr with strong nitrio acid, but the writer believes that this is due to impurities, and that tests for

emetine aro desiderata.

Gelsemine.-Sulphurio acid and potassium dichromate produce an immediate but evanescent violet coloration, and the chromate is quickly reduced. The alkaloid is naturally associated with gelseniic aoid, a substance which in alkalino solution fluoresces strongly, and is yellow by transmitted, blue by reflected light.

Hyoscyamine. -- No good chemical test for this

alkaloid is known.

Jervine.—The salts of this alkaloid, especially the acetate, are ppd. by nitrio acid and by potassium nitrate.

Morphine gives with nitricacid a deep red colour, not materially altered by the subsequent addition of stannous chloride. With sulphuricacid and potassium dichromate a green coloration is gradually developed. Ferrio chloride gives a blue or blue-green coloration. It liberates iodino from iodic acid, and the mixture shaken with chloroform imparts a violet tint to this; but the brown colour is only deepeued and altered in tint by ammonia. Sulphomolybdic soid gives an immediate purple coloration.

Narcotine.—Uncoloured by sulphuric acid, except this contains a trace of nitrio acid, when a fine cherry-red colour is gradually developed, and is porsistent. Its acidulated solution when warmed with bromine-water, added drop by drop, develops a purple or violot tint.

Nicotine.-Oily, and having the odour of tobacco. Freely soluble in water. Its acidulated solutions give a copious pp. with brominewater, and this pp. disappears when excess of sodinm hydrate is added.

Physostigmine. - Sulphnrio acid gradually produces a reddish colour. Its solutions acquire a red colour on standing, and at once when treated with sodium hydrate and warmed, and on evaporation leave a bluish residue which on acidulation affords a dichroio (red and blue)

solution, which becomes permanently red on

standing.

Pilocarpine affords no very characteristic chemical reactions. With snlphuric acid and potassium dichromate & green colonr, due to reduction, is developed.

Piperine has the pungent fragrant odour of pepper. It is turned of a deep red colour by sulphuric, and of an orange colour by nitric acid.

Quinine.- Its solution in sulphurio acid is fluorescent. Treated with bromine-water and then excess of ammonia added, an emerald

green coloration is produced.

Salicin. - A neutral glucoside. It is not withdrawn from its acid or alkaline solutions by ether, benzene, or chloroform, and hence isnot obtained in the ordinary processes for the separation of the alkaloids. It is turned of a cherry-red colour by sulphuric acid, and then on the addition of potassium dichromate an odour of meadow-sweet is evolved. Fused and partially sublimed in a test-tube, and then dissolved iu water, ferrie chloride strikes a violet colour.

Strychnine.—Sulphurie acid gives no colonr till potassium dichromate, MnO₂, PbO₂, or ferrieyanide of potassium is added, when immediately a fine blue-violet colour is produced which gradually passes into reddish-violet, red, and finally cherry-red. Evaporated to dryness with furning nitrio acid, and the residue moistened with alcoholio potash a dcep orange colour is produced.

Veratrine.-Tonched with strong sulphuric acid, this alkaloid gradually develops a fine red colour, or immediately on warming. Its solution in cold strong hydrochloric acid is colourless, but becomes intensely red on boiling.

ALKAMINES, aleamines, alkines, or aloines. Names used by Ladenburg to denote substances that contain both alcoholic hydroxyl and amido-

gen, such as oxy-cthyl amine, C.H. (OH).NH.

ALKANET. The commercial name of two different plants. True alkanet is Lawsonia inermis, false alkanet is Anchusa tinctoria. The leaves of Lawsonia contain a yellow dye, its roots contain a red pigment, used as a cosmetio. The root of Anchusa (Oreannette, Radix Al-

cannæ spuriæ) contains auchusin.

Anchnsin or Alkannin C₁₁H₁₄O₈ (Bolley a. Wydler, A. 62, 141) or C₁₁H₁₂O₄ (Pelletier, A. 6, 27) or C₁₅H₁₄O₄ (Carnelutti a. Nasini, G. 1880, 283; B. 13, 1514). Obtained by extracting the root of Anchusa tinctoria with petroleum; the crude product is treated with dilute potash, the filtrate is shaken with ether, and the alkannin is ppd. by a current of CO. It is a brownish-red mass with a metallic lustre; sol. ether, chloroform, and acetic acid, sl. sol. alcohol. Softens below 100°. Alcoholic solutions give with baryta-water a blue pp. of a barium compound. NaOAc and Ac₂O produce a crystalline descetyl derivative: C₁₈H₁₂Ac₂O₄(?). Nitrio acid forma oxalic and succinic acids. Alkannin appears to bo allied to santalin. An alcoholio solution of alkannin dyes cotton mordanted with alum, violet; iron mordants give a grey colonr. Turned blue by alkalis, especially ammonia (Böttger, J. pr. 107, 46; Enz. J. 1870, 935). Alkannin, unlike resamiline, is not abstracted by cubss of gelatin from its solution. Its absorption-spec-

trum shows & bands, dividing the spectrum between D and the blue strontium line into a equal parts. On adding ammonia the red solution turns blue, how showing 2 bands, one at D, the other in the red, two thirds of the way towards the lithium line (A. Dupré, C. J. 37, 572).

ALKARSIN. Name given by Bunson to cacodyl or arsenic di-methido (q. v.), C.H.As, as being empirically alcohol in which O has been displaced by As (A. 24, 271).

ALKYL. An alcohol radiole. ALKOYL. An acid radicle. ALLANIC ACID C4II5N5O5Aq.

Formed, together with urea and allanturio acid by the action of nitrie acid in the cold on allantoin (E. Mulder, A. 159, 353). Stellato needles (from water). Sl. sol. cold water. Decomposes at 210°-220° without melting. Does not give off gas with HNO4 containing N.O4. Gives no pp. with CaCl. Aq and NH, Aq. Gives pps. with AgNO, Aq and NH, Aq, and with basic lead accetate, but not with neutral lead accetate.

Salts .- NH, A': prisms .- HO.Pb.A': ppd. by basic lead acetate. - 2PbA'25Pb(OH)2-AgA' aq: amorphous pp

ALLANTOÏC ACID

C.H., N.O., i.e. NH., CO.NH.CH(CO.H).NH.CO.NH. A solution of allantoin in aqueous potash which has stood for some days, no longer gives a pp. with acetic acid, even after some time; but if a little alcohol be also added, and the liquid be left in an exsiccator over lime, orystalline potassio allantoate, KA', separates (E. Mulder, 159, 362; Schlieper, A. 67, 231; Ponomarow,
 17. R. 11, 13. The solution of potassio allantoate gives crystalline pps. with Pb(OAc)2 and AgNO, but not with BaCl. BaCl, and alcohol give a hygroscopic curdy pp.
Salts.—NH,A'—NaA' aq.—KA'.—BaA', 2aq.

-PbA'2 aq. -AgA'.

ALLANTOIN C.H.N.O. i.e. NH.CO NII.CH.NII.CO.NH. NH.CH(OH) NH.C:N.CO.NH,

Mol. w. 158. S. 62 at 20°; 3.3 at 100°.

Occurrence.—In the allantoic liquid of the cow (Lassnigne, A. Ch. [2] 17, 301; compare Vauquelin, A. Ch. 33, 269). In urine of sucking calves (Wöhler, A. 70, 229). Occasionally in urine of dogs (Salkowski, B. 9, 721; 11, 500; Meissner a. Jolly, Z. 1865, 131). In the young leaf-buds of the plane and maple, and in the bark of the horse-chestnut tree (E. Schulze, B. 14, 1602; J. pr. 133, 147; H. 9, 425). In wheat, to the amount of 5 p.c. of the embrye (Richardson a. Crampton, B. 19, 1181).

Formation .- 1. By treating uric scid with boiling water and PhO. (Liebig a. Wöbler, A. 26, 244; E. Mulder, A. 159, 349), with KOH and potassic ferricyanide (Schliepor, A. 67, 216), or with KMnO, (Claus, B. 7, 227).-2. By heating glyoxylic acid (1 pt.) with urea (2 pts.) eight hours at 100° (Grimaux, C. R. 83, 62).-3. By the action of nitrous acid on dialuric acid (Gibbs. A. Suppl. 7, 337).—4. By heating mesocalic sold with urca at 110° (Michael, Am. 5, 198).

Properties. — Glassy monoclinic prisms

Tasteless. (Dauber, A. 71, 68). Neutral.

Readily soluble in alcohol.

Reactions.—1. Dry distillation gives ammonic carbonate and cyanide, and charcoal .- 2. Gently heated with hydrochloric or nitric acid it gives urea and allanturic soid.—3. Hot sulphuric acid forms CO, CO, and NH,—4. Boiled with baryta-water, CO2, NH3, oxalio acid, and hydantoin are got (Baeyer, A. 130, 161) .-5. Hot cone. potash forms CO2, NH2, exalio acid, and acetic soid .- 6. Cold potash slowly forms allantoïo acid (q. v.).—7. Nitric acid of S.G. 135 forms, on boiling, allanio acid (q. v.).—8. Potassic ferricyanide and KOH form allantoxanio aoid (Mulder, B. 8, 1291) .- 9. Sodium amalgam forms glycoluril, C, H, N, O, (Strecker a. Rheineck, A. 131, 119).—10. Hydric iodide reduces it to urea, and hydantoin or glycolyl-urea (Baeyer, A. 117, 178).

Tests.-1. A cono. solution of furfurol, to which a little HCl has been added, gives a violet colour with an aqueous solution of allantoin (Schiff, B. 10, 771).-2. Mercuric nitrate (but not chloride) gives a pp., as with urea. 100 g. of dry allantoin require 172 g. of mercuric oxide.

The pp. is (C₄H₈N₄O₃),5HgO.
Compounds with Bases.—These are formed by boiling aqueous solutions of allantoin with metallic oxides. They are sparingly soluble (Limpricht, A. 88, 94).

 $\begin{array}{lll} \text{Coll}_{1} & \text{Coll}_{2} & \text{Coll}_{3} & \text{Coll$

The following are described as true salts: AgC₄H₂N₄O₃, got as a pp. by ammoniacal AgNO₄.

—KC₄H₂N₄O₃: from allantoin, KOHAq, and alcohol, in exsiccator.

Nitrato C411, N4O3HNO3. Amerphous. Decomposed by water or alcohol into HNO, and

allantoin.

Constitution.-The constitutional formulæ given above are chiefly based upon Formation 2 and Reaction 10

ALLANTOXATDIN

NH.CO C₃H₂N₃O₂ aq i.e. CO(NH.C:NH

When allantoxanic acid is liberated from its salts, it at once splits up into CO, and this body (Ponomarcw, J. R. 11, 47). Glittering prisms or V. sol. boiling water, sl. sol. cold water or alcohol, insol. ether. Decomposed by heat, giving off HCN, HCNO, and NH, Acid reaction. Boiled with water, or treated with cold Na CO.Aq. it splits up into formio acid and biuret.

Salts. - KA': ppd. byalcohol. - AgA'.
ALLANTOXANIC ACID C.H.N.O. i.e.
NH.CO

NH.Ċ:N.CO₂H

Formation .- 1. Allantoin dissolved in aqueous KOH is treated with potassic ferricyanide until the colour is permanent. Acetic acid is then added, when C.H.KN₂O₄ is ppd. (Van Embden, A. 167, 39).—2. From allantoin, KOH₄ and KMnO, (Mulder, B. 8, 1292; Ponomarew, J. R. 11, 19).—3. From oxalyl-di-urelde and aqueous potash (P. B. 18, 982).

Properties .- The soid, liberated from its lead salt by H.S. splits up into allantoxaldin and CO. Salts.-NH,A': needles. -(NH,),C,HN,O.

4-12-8-17

-EA': silky needles. S.: \$6. Boiled with water, it gives CO₂, biurel, and formic acid. Reduced by sodium-anfalgam to hydroxonic acid.—

K₂C₄HN₂O₂ aq: v. sol. water, insoluble in dry aloohol.—Ba(C₁HN₂O₂); daq.—BaC₄HN₂O₄ 2aq.

—Pb(C₁HN₂O₃); ½aq: very thin needles.—Pb(C₁HN₂O₄.—AgC₄HN₃O₄: crystalline pp.—Ag,C₄HN₃O₄: gelatinous.

Ethyl ether C.H.EEN.O.: from AgA' and Etl.

ALLANTURIC ACID C.H.N.O. i.e.

NH.CO Glyoxyl-urea.

NH.CH(OH)

Formation.—1. By boiling allantoin with HNO₃, HCl or PbO₂ or heating with water at 140° (Pelouze, A. Ch. [3] 6, 71; Mulder, A. 159, 359).—2. Formed, together with glycoluril and urea, by sction of sodium-amalgam on allantoin (Reinecke, A. 134, 220).—3. By boiling allantoin acid or alloxanic acid with water (Ponomarew, J. R. 11, 15; Schlieper, A. 56, 5).—4. By oxidation of hydantoin (Baeyer, A. 117, 179; 130, 160).—5. By boiling uroxanic acid with water (Medicus, B. 9, 1162; Ponomarew, B. 11, 2155).

Properties.—A deliquescent gummy mass. Insol. alcohol. Boiling potash forms CO., NII., acetic acid, and oxalic acid (Medieus, B. 10, 544).—Salts. These are amorphous.—KA'HA' 2aq. S. 10.—BaA'. 3aa.

S. 10.—BaA', 3aq.
ALLENE. Name sometimes used instead of ALLYLENE.

ALLITURIC ACID C.H.N.O.

S. 5 or 6 at 100°. Obtained from an aqueous solution of alloxantin, mixed with HCl, by rapidly evaporating to a small bulk, and treating the resulting powder with HNO, which dissolves alloxantin but not allituric acid. The latter crystallises from water as a bulky yellowish-white powder (Schlieper, A. 56, 20). Not attacked by cone. H₂SO₄ or HNO₂. Evolves NH₃ when boiled with KOH.

ALLO.—A profix proposed by Michael (B. 19, 1878) to denote unexplained 4 somerism; thus fumaric acid would be called allo-maleic acid.
ALLOCAFFEINE C.H.,N.O..

ALLOCAFFEINE C₈H₉N₃O₃. [196°]. (E. Fischer, A. 215, 276).

Formation.—1. Obtained by action of water on the unstable product of addition of bromine to caffeine methylo-hydroxide.—2. One of the products of action of HCl and KClO, on caffeine methylo-hydroxide (Schmidt a. Schilling, A. 228, 162).

Small white trimetric crystals, a:b:c= 6953:1:5401, soluble in benzene, chloroform, and hot water, nearly insoluble in cold water,

sparingly in alcohol or ether.

Reactions.—1. Decomposed by boiling water into CO₂ and methyl-caffuric acid. Allocaffeine is therefore probably methyl-apocaffeine.—2. HNO₂ (S.G. 1-2) gives cholestrophane, methylamine, and CO₂.—3. HCl and KClO₂ form dimethyl-alloxan, amalic acid, cholestrophane, methylamine, and CO₂.—4. Bromine appears to form an addition product, but it is decomposed by water into allocaffeine, cholestrophane, and methylamine hydrobromide.—5. Boiling baryta forms sameosine, formic acid, and CO₂.

Constitution.—Inasmuch as it splits off

Constitution.—Inasmnch as it splits off NMeH, in reactions where caffeine splits off NH, the Me (and consequently OH also) must

he attached to nitrogen, the formula being either:

i.e. NH₂CO.NH.CO₂H Urea v-carboxylic acid. The free acid splits up at once into CO₂ and urea. Its ethers are formed by passing vapour of eyanic acid into alcohols: 2CONH + HOEt = NIL_CO_Et + CONH = NH_CO.NH.CO_Et. The ethers are sparingly solublo crystalline solids.

Salts.—BaA'. Obtained from the ether by cold baryta-water (Licbig a. Wöhler, A. 59, 291). When boiled with water it gives off CO., deposits BaCO., and urea is left in solution. Dry distillation produces basic cyanate, NH., and CO., It gives no pp. with AgNO.,—Salts of Ca. K, and Na have been prepared.

Methyl allophanate NH2.CO.NH.CO2Me

(Richardson, A. 23, 138).

Ethyl allophanate Eth.' [191°]. 1. From alcohol and the vapour of cyanic acid (L. a. W). 2. From ClCO_Et and urea (Wilm a. Wischin, Z. [2] 4, 5).—3. Together with oxamide and alcohol by heating urea with oxalic ether at 135°-170° (Grabowski, A. 134, 115).—4. From potassic cyanate, alcohol, and chloro-acctic ether (Saytzeff, A. 135, 230) or chloroformic ether, ClCO_Et (Wilm, A. 192, 244).—5. From potassic cyanate, alcohol, and IHCl (Amato, G. 3, 469). Small needles. Tasteless. Sl. sol. cold water, nore soluble in alcohol. V. sl. sol. cold ether (difference from carbamic ether). At 190° it slowly changes to alcohol and cyanuric acid. Alcohol at 160° converts it into carbamic ether: NII_CO.NH.CO_Et + HOEt=2NH_CO_Et (Hofmann, B. 4, 268).

Acetyl derivative, NHAc.CO.NH.CO.Et. [107°]. Silky needles (from alcohol) (Seidel, *J. pr.* [2] 32, 273).

Benzoyl derivative NHBz CO.NH.CO.Et. [163°]. Together with alcohol, lICl, and CO. from benzoyl chloride and urethane (Kretsohmar, B. 8, 104).

Propyl allophanate PrA'. [150°-160°].

(Cahours, J. 1874, 834).

A myl allophanate C,H,A'. [162°]. From eyanic acid and amyl alcohol (Schlieper, A. 50, 23). From amyl alcohol and urea (Hofmann, B. 4, 267). Unctuous pearly scales (from water).

Oxethyl altophanate HO.C.H.A. [160°]. From glycol and cyanio acid vapour. Shining laming (from alcohol) (Baever. 4, 114, 160).

lamins (from alcohol) (Baeyer, A. 114, 160).

Dioxy-propyl allophanate C₂H₃(OH)₂A'.

[160°]. From glycerin and cyanic acid vapour (B.). Plates (from alcohol). Sol. water. Heated with baryta-water, it forms BaCO₃, urea, and glycerin.

Phenyl-allophanate PhA'. Cyanic acid vapour is passed into phenol; the product is dissolved in alcohol and ppd. by ether. Sleuder crystals. At 150° it splits up into cyanic acid and phonol (Tuttle, J. 1867, 451).

Propenyl-methoxy-phenyl-allophanate C₁₂H₁, N₂O₄ i.e. NH₂CO.NH.CO.O.O.Q.H₃(C,H₃).OCH₄.

From eugenol and cyanic acid vapour (Baeyer.

4.114, 164). Needles. Insol. water. Sl. sol. in the liquid state for some degrees above the cold alcohol.

Amide of allophanic acid NH..CO.NH.CO.NH. v. Biurer.

ALLOTROPY (otherwise turned, otherwise formed, from allos = another, and Tpomos = manner) denotes the appearance of one and the same substance in several different states, distinguished from each other by different properties. The term was introduced by Berzelius in 1840 (J. No. 20 for 1839, pt. ii. p. 13), because he held the term 'isomerism' to be inadmissible where the subject of modification is an elementary substance, isomerie states being traceable to different modes of combining equal numbers of atoms of the same elements. In the view of Berzelius, accordingly, the allotropic modifications of the elements are not to be explained by differences in the arrangement of their atoms. bnt he expressed no opinion whatever about their actual cause. Since, however, he indicated it as probable that even in compounds the olements retain their allotropio states, and thereby often occasion isomeric forms of compounds (J. No. 23, p. 51; No. 21, p. 32), he appears to have been of opinion that the cause of the allotropic transformation is to be sought in a change in the atoms themselves. Now that we have learned to appreciate more correctly the doctrine of Avogadro, and so have become accustomed to consider the molecules of the majority of clements as particles composed, like those of compounds, of several atoms, the distinction introduced by Berzelius between allotropy and isomerism has lost its original meaning. But the term allotropy has been retained, being used, however, with reference not to elements only but also to compounds. Accordingly we distinguish between allotropy of elements and allotropy of compounds. The former, according to the medern use of the expression, embraces all the different forms in which an element appears; the latter only those cases in which, while the composition remains the same, there is a change in the physical, but none, or at any rate none of any consequence, in the chemical, properties, thus apparently warranting the assumption that there has been no change in the linkage of the atoms by which, doubtless, chemical behaviour is essentially determined. Allotropy of compounds is accordingly synonymous with physical, as opposed to chemical, isomerism. But since the two groups of properties are closely connected, and any change of the physical is usually accompanied by a change, however small, of the chemical also, no sharp line is to be drawn between the two kinds of isomerism.

On the other hand, the transformation of one allotropic form into another offers so many analogies to the transformation of one state of aggregation into another that, strictly speaking the three states of aggregation of any substance should be described as three allotropic modifications of it (Lehmann, Z.K. 1877. 1, 97). Hitherto, however, it has not been usual so to describe the states of aggregation, and, consequently, on this side also, the notion of allotropy is not to be defined with perfect exactness. The melting of ice, for example, is a transformation of the lighter into the heavier modification of water, for the particles of the lighter are still retained

melting point, and bring it about that the maximum of density appears not at 0° but at +4°. Something similar probably takes placa in many, if not in all, other substances, only the difficulties of observation are greater. But as it has been observed that changes in the properties of a substance usually proceed differently and follow different laws according as the substance is noar to, or more remote from, its meltingpoint (no matter whether above or below it), we may conclude that immediately below the melting point the solid substance already contains isolated portions of the liquid modification, and that above the melting point the liquid body still contains portions of the solid modification. But even if we do not account these changes of aggregation as instances of allotropy, the number of cases of alletropy as yet known, while suffering a very important diminution, will still remain pretty considerable.

I. ALLOTROPY OF THE ELEMENTS .- Allotropy, taken in the narrower sense, has hitherto been observed only in the non-metallic or semimetallic clements. Among metals preper it has been found only as regards orystalline form, in which case it is usually known as dimorphism or polymorphism. Since, to the best of our present knowledge, the gaseous molecules of the metals consist of single atenis,1 while those of the semimetals and non-inctals are composed of several atoms, the absence of allotropic meditications of the metals proper tells in favour of the present view, which is different from that of Berzelius, and is to the effect that alletropy of the elements, like isemerism of compounds, depends on differences in the mode of union of the atoms, and not on any changes in the atoms themselves. Polymorphism, occurring as it does even among metals, may be explained by supposing that there are differences in the arrangements of the atoms as well as of the molecules, while the existence of allotropic modifications in the niclted, the dissolved, or the gasified, state points to differences in the constitution of the molecules, i.e. to different modes of uniting the atoms to form molecules.

The appearance of allotropy seems to be favoured by smallness of atomic weight, for not unfrequently in one and the same natural family allotropy shows itself only in the first members, while the members with higher atomic weights exhibit it cither in some properties only or not at all. In the family of the halogens, F, Cl, Br, I, allotropy has not been observed, unless we consider as such the splitting of molecules at high temperatures into separate atoms (Victor Meyer). Hydrogen dees not exhibit allotropy. On the other hand allotropy is found very notably in the first members of the oxygen-sulphur family. Ozone exhibits much more strongly marked chemical obaracters, and morcover a greater density, than oxygen. If the molecular weight of ordinary oxygen is represented by O2, that of ozone is probably O₃, ozono being thus a polymeride of oxygen. Sulphur in each of its states of aggregation exhibits allotropic modifications, and these to some extent correspond with each other. In the solid state it is: (1) rhombic;

" It should not be forgotten that the data are most

13 (Gernez), soluble In CS₄: (2) monoclinic; D. = 1.96, M.P. 117° (Gernez), soluble in CS₂: (3) amorphous plastic; D. = 1.90 to 1.93, insoluble in CS₂: (4) according to Gernez (C. R. 98, 141) and Sabatier (C. B. 100, 1346) cryetallised in little rods with a lustredike that of mother-of-pearl. The last modification Maquenne (C. R. 100, 1499) considers to be distorted rhombic crystals, which according to Gernez are very easily produced out of the fourth modification (C. R. 100, 1584) without being identical with it. Liquid sulphur is: (1) immodiately above the melting point thin and clear: (2) at about 200° thick and dark: (3) at about 340° thin and dark. The vapour: (1) between the boiling-point (446') and about 500° has V.D. = 6.6, molecular weight = S_6 : (2) above 700° V.D. = 2.2, molecular weight = S_6 : The behaviour of selenion is analogous to that of sulphur. When solid this substance is: (1) red, amorphous, vitrous, or pulverulent, D. = 4.26, soluble in CS₂: (2) red, erystallised, monoclinie, isomorphous with sulphur, D. = 4.51, soluble in CS₂: (3) gray, granularly crystalline, D. = 4.60, insoluble in CS₂. Whether the black foliated crystals, insoluble in CS2, D. = 4.80, obtained from a solution of potassium selenide, are identical with the third modification remains to be determined. Liquid selenion is: (1) at low temperatures in a thin stratum light-red and transparent: (2) at higher temperatures, dark. Gaseous selenion under 1400° consists in part of molecules composed of more than two atoms: above 1400° all the molecules are diatomic, V.D. = 5.68, molecular weight Se... Of tellurium no allotropie form is known with certainty, yet it is worthy of remark that its electrical conductivity, like that of selenion, but contrary to that of all other conductors of the first class, increases with rising temperature. This may be explained by supposing the production of a modi fication with better conductivity. As regards the nitrogen family, the existence of any allotropic forms of nitrogen has not yet been conclusively proved, but solid phosphorus exists in three forms: (1) colourless, very easily burnt, soluble in CS2 and in many oils, crystallising out of these solutions according to the regular system, $D_* = 1.83$: (2) red, amorphous, $D_* = 2.18$: (3) darkred crystallised in rhombohedral forms, in the highest degree indifferent, D. = 2.31. The last two forms perhaps represent one and the same modification. In the liquid state there seems to be only one modification—the colourless: in tho gaseous state, on the contrary, there appear to be two, since the vapour-pressure over colourless phosphorus is greater than that over red at the same temperature, and the vapour condenses under certain circumstances into the one modification and under other circumstances into the other. Arsenic is: (1) amorphous, D. = 4.72, less easily oxidised than the following variety: (2) crystallised in rhombohedral forms, D. = 5.73, Whether explosive antimony (Gore), D. - 5.83, is a distinct modification cannot be quito definitely determined, since it cannot be obtained free from chloride. For ordinary antimony $D_1 = 6.71$. Of bismuth no allotropic modification is known.

In the carbon family carbon exists: (1) as diamond, regular, very hard, D. = 3.52: (2) ae VOL. I.

monoclinic (Clarke, Nordenskield), D. = 2.33: (3) amprihous charcoal, D. = 1.87 to 2.30, agreeing with graphite in many properties and hence perhaps not to be regarded as a distinct modi-fication. Silicons: (1) amorphous, easily oxi-dised: (2) crystal ised according to the regular system, D. = 2.20 to 2.49. The so called graphitoidal variety consists of distorted regular erystals. Of titaniilin and thorium allotropic forms are not known. Zirconium has been obtained amorphous and crystallised. Tin also appears to be dimorphous.

The element boron is probably capable of allotropic modification, yet hitherto it has been obtained pure only in the amorphous form. The crystallised always contains aluminium or earbon. Some of the platinum motals, namely iridium and pulladium are said to occur in twok

forms, regular and hexagonal.

11. Allotropy of Compounds, or Physical Isomerism, may be theoretically defined as isomerism with identity of atomic linkage. The following inorganie compounds exhibit remarkable instances of allotropy; calcium carbonate. (as eale-spar and arragonite); silica (quartz, tridymite, agate); titanium oxide (rutile, brookite anatase); the nitrates of sodium, potassium, am monium, and silver; sodium metaphosphate; arsenious and antimonious oxides; the sulphates of magnesium, iron, and copper; potassium dichromate; silver iodide; zine chloride; mercuric chloride; manganous chloride; and indeed many other substances. Many instances of allotropy have also been observed among the compounds of earbon, particularly in the following substances: benzophenone; isohydrobenzoin diacetate (Zineke); dibromopropionic acid (Tollens); tolylphenyl ketone (Van Dorp, Zineke); metachloronitrobenzene; chlorodinitrobenzene (1:3:4) (Laubenheimer); oxycamphoronic acid (Zepharowich); the benzoylated and anisylated hydroxylamines (Lossen); hydroquinone; paranitrophenol; stilbene chloride; dibromofluorens (Lehmann); tetramethyldiamido-triphenyl-methane; diphenylnaphthylmethane; pentamethylleukaniline (B. Fischer, Lehmann); dibenzoyl-diamidodibromodiphenyl (E. Lellmann). No definite and regular relation between the composition of oarbon compounds and the existence of allotropic forms of these compounds has as yet been recognised.

The production of allotropic modifications, and the transformation of one modification into another, are effected, as a general rule, by changes of temperature. Tho eases in which we are entirely ignorant of the conditions under which allotropic modifications are produced, are but few. Thomost notable is that of one of the modifications of earbon-the diamond, but on the other hand the transformation of diamond into graphite has been observed. One of the allotropic states usually corresponds to a specified interval of temperature, so that at a definite limit of temperature the one modification passes into the other. Yet we frequently succeed in cooling the modification belonging to the higher temperature below the lower limit. and sometimes also in heating the other modification above this limit, without any transformation taking place. But when such a modifiraphite, either rhombohedral (Kenngott), or cation is proserved above its fixed limit, the

unstable, and is often destroyed by very trifling causes, a particularly easy means of upsetting it being to bring the substance inte contact with a crystal of the modification that is stable at the prevailing temperature. On transformation into the stable form thereupon ensuing, heat is produced or disappears, according as contraction or expansion takes place. This thermal effect may be very considerable.

The temperature of transformation has been determined for rhombic and monoclinic sulphur by L. Th. Reicher (Z. K. 1884. 8, 6) to be 95.6°. Belew this the rhombic form is stable, above it the monoclinic, the other being unstable. The amorphous form is unstable at all temperatures below, and also for a considerable interval above, the melting point; the temperature at which it becomes stable has not been determined, but probably it lies not far below the beiling point. When cooled quickly both the monoclinie and the amorphous form may be kept a considerable time at comparatively low temperatures. One might be tempted to suppose that the modificaions that have thus become unstable would pass to the stable forms the more easily the greater he distance of their temperature from that of transformation; yet below the temperature of transformation this is not the case; on the contrary, transformation into the rhombie modification ensues the more easily the higher the temperature and therefore the nearer it comes to the temperature of transformation. This is andoubtedly due to the circumstance that the mobility of the particles increases as the temperature increases. The behaviour of seleniou is similar to that of sulphur. Amorphous selenion is produced only above the melting point, which is 217°, nevertheless when this variety is quickly cooled it remains stable for some time, and begins to pass into the grey crystalline form only at 80° (Hittorf); the progress of this change is however more rapid at 125°. The temperature of transformation of the red soluble crystals of selenion is about 110° (Mitseherlich).

The conditions under which phosphorus passes from one of its modifications into another are very remarkable. If colourless phosphorus is vaporised in a vessel too small to contain the whole of the phesphorus as vapour, the red variety is formed at 210° and upwards; the obange proceeds more rapidly at 260°, and very quickly above 300°. Conversely, red phosphorus, if it can transform itself freely into vapour, and if the vapeur is allowed to cool, is re-cenverted at 260° into the colourless form : the red modification is formed only if the vapour has been heated above as red heat and then allowed to cool (Hittorf). Arsenic vapour cendenses below 220° to form amorphous arsenie; at a higher temperature to form crystallised. At 360° the former passes into the latter with production of heat. Tin is converted by very great cold, under conditions not yet exactly determined, into loosely cohering columnar aggregations of grey colour and diminished density (Fritsche, Petri, Schertel). Light too may bring about the production of allotropic modifications; through its influence selenion and tellurium temporarily acquire a better electric conductivity—a fact which has the rhombohedral form, which is produced at been applied in telegraphy. Phosphorus be- lower temperatures. Rock-crystal and amorphous better electric conductivity-a fact which has

state of equilibrium attained by its particles is comes red through the action of light. Electriolty likewise may convert phosphorus, in vacuo, into the red modification, but perhaps the transformation may be due only to the heat produced.

Among compound substances the phenomenen of the transformation of one allotropic modification into another has been observed by many authors, but it has been studied with special attention by O. Lehmann (passim, and in later papers in Z. K.) He has proved that it obeys the same laws that hold for the elements. In most cases an unstable modification. differing from the ordinary stable one, is ebtained by raising a substance to a high temperature and then cooling it quickly to a temperature a long way below that of transformation. It is supposed that in such circumstances the particles do not find time and opportunity to assume the position of equilibrium corresponding to the lower temperature. The unstable state thus produced may be assumed alike by solid, melted, and dissolved, substances, and may be maintained, especially at pretty low temperatures, for a long time. In many eases, e.g., in that of hydrogninone, the one modification (in this ease the unstable) is obtained by melting or subliming: the other form is obtained from solutions. In other eases, either form may be obtained from the same nielted body, or from the same solution, according as it is brought into contact with a erystal of the one form or of the other. If fragments of erystals of both modifications are introduced simultaneously, both of them at first increase in size; but as soon as the two crystalline masses come into contact the form that is stable at the prevailing temperature grows into, and at the expense of, the unstable, while the latter dissolves or is consumed. As a general rule the modification that is unstable at a low temperature has a lower melting point than the stable, so that many substances on being heated are observed first to melt, then to solidify again, with transformation into the other modification. and finally to melt a second time. This phenomenon may be observed with special distinctness in the ease of dibenzoyldiamidodibromodiphenyl. because here the melting points of the two forms lie unusually far apart. The needles of thie substance crystallised out of alcohol melt at 195°, when quickly cooled the melted substance solidifies to a vitreous mass, which, when again heated, melts at 99°, re-solidifies in a crystalline form between 125° and 130°, and then melts once more at 195° (Lellmann).

Many compounds, especially inorganio compounds, behave like selenion; the form produced at high temperatmus may remain stable far below the temperature of transformation, and may become unstable only on being heated to the neighbourhood of the temperature of transformation. Arragonito, the rhombic form of calcium carbonate, which separates from hot solutions (and according to G. Rose from very dilute cold solutions also) is perfectly stable at ordinary temperatures. If, however, a crystal is heated, it breaks up, long before giving off carbon diexide, into a mass of small crystals of calc-spar (Haidinger), thus passing over into suce are persectly stable at ordinary temperatures, but at the temperatures of the porcelainkiln they are changed into tridymite, the third modification, which in turn is likewise stable at lower temperatures. As regards other substances, particularly organic compounds, the forms to be classed as unstable usually possess much less stability, but still of course they are not altogether destitute of it. This persistence in a state no longer completely stable may be explained by supposing that a certain impulse, or an increase of the proper motion of the particles, is required to change the state-to make the particles leave their respective positions and pass over into new ones. That the change is attained most easily and most surely by contact with a crystal of the stable modification, is undoubtedly due to the power of every crystal to give to the particles settling on it a dofinite and regular orientation and arrangement. L. M.

ALLOXAN C, II, N, O, aq (and 4aq.)

i.e. CO NH, CO CO. Mesoxalyl-urea. Mol.

w. I42.—Discovered in 1817 by Brugnatelli, who
named it crythric acid. Subsequently examined
by Liebig a. Wöhler (A. 26, 256), and by
Schlieper (A. 55, 253).

Formation. -1. By exidation of uric acid by HNO₃ (S.C. 1-12) diluted with water (9 pts.) at 70°. By adding SnCl., alloxantin is ppd., and, after washing, is re-exidised to alloxan by nitric acid (2 pts. of S.G. 1-52 mixed with 1 pt. of S.G. I-42) in the cold (Liebig, A. 147, 366, Bl. [2] 9,152). -2. From nric acid and aqueous Br, Cl, er I (M. E. Hardy, Bl. [2] 1, 445).—3. From xanthine, KClO₃, and HCl (E. Fischer, A. 215, 310).

Properties.—A warm saturated aqueous solution deposits on cooling trimetric efflorescent crystals (with 4aq). If the solution is kept warm while evaporating monoclinic prisms (with aq) are got. V. sol. water or alcohol, ppd. from solution by HNO₃. Astringent taste, reddens litmus, does not decompose CaCO₃. Aqueous solution turns the skin purple, imparting a peculiar smell.

Reactions.-1. Hot dilute nitrie acid forms CO, and parabanic acid, the latter then becoming CO₂ and urea.—2. Boiling potash forms mesoxalio acid and nrea.—3. Boiling very dilute sulphuric acid forms ammonic hydurilate. -4. Beiling aqueous HCl or II, SO, forms alloxantin, which separates; dialuric acid, ammonic oxalate etc., remain in solution. - 5. Boiled a long time with water, it forms CO, parabanic acid, and alloxantin.—6. By reducing agents (II.S. SnCl., Zn and IICI) it is converted into alloxantin, and finally into dialuric acid .- 7. Boiled with ammonia and sulphurous acid, it forms ammonic thionurate (q. v.). 8. KHO or baryta converts it into alloxanie acid; baryta or lime-water giving white pps. of baric or calcio alloxanate. If the alkali be in excess, the pp. centains mesoxalate. 9. Warm aqueous ammonia forms a yellow jelly of the ammonium salt of 'mycomelic acid' O.H.N.O. (L. a. W.).—IO. Ferrous sulphate gives a deep blue colour.—11. Boiled with water and PbO, there results CO, PbCO, and usea.—I2. Boiling aqueous lead acetate forms lead mesoxalate and nrea.-I3. Boiling aqueous NaNO, and acetic acid form sodic oxalurate (Gibbs, Am. S. [2] 48, 215).—14. Hydroxylamine hydrochloride forms violuric acid.—15. With a dilute solution of pyrrol it forms crystalline pyrrol allexan (Ciamician a. Silber, B. 19, 106, 1708).—16. PCl, mixed with POCl, at 130° forms tetrachlero-pyrimidine (Ciamician a. Magnaghi, B. 18,

Metallic derivatives.— C₄Ag₂N₂O₄.—
C₄H<sub>N₂O₄IgO7aq: ppd. by increaric nitrate.—
Compounds with Bisulphites.—
C₄II₁N₂O₄NaIISO₄I₂aq: largo crystals, v. sol. water.—C₄H₂N₂O₄KHSO₅aq: m. sol. cold water, v. sol. hot water.—C₄H₂N₂O₄KHSO₅aq: m. sol. cold water, prieht a. Wuth, A. 10a, 41).
ALLOXANIC ACID C₄II₁N₂O₅</sub>

i.e. NH..CO.NH.CO.CO.CO.M. Mesoxaloxyl-ured S. (alcohol) about 20. (Liebig a. Wöhler, A. 26 292; Schlieper, A. 55, 263; 56, 1; Städeler, A. 97, 122; Baeyer, A. 119, 126; 130, 159). Forme from alloxan by treatment with aqueous fixed alkalis or alkaline carbonates. White needles of warty masses. V. sol. water; sl. sol. ether.

Reactions.—1. Boiling the aqueous solution produces CO₂, lencoturie acid (q. v.), allanturie acid and hydantoin.—2. Alloxanates are converted by boiling water into mesoxalates and urea.—3. Nitrie acid forms CO₂ and parabanic acid.—4. HI reduces it to hydantoin, giving off CO₂ (Baeyer).

Salts. The alkaline alloxanates are soluble in water. The normal salts of other metals are usually insoluble. Ferrous sulphate gives a dark blue pp. with potassic alloxanate. NH₁C, H₂N₂O₃. S. about 30.—BaH₂A''₂2aq.—BaA''4aq.—CaH₂A''₂6aq. S. 5.—CaA''5aq.—CuA''4aq. S. 17 to 20.—CuA''Ca(OH)₂.—PbH₂A''₂2aq.—Pb₃H₃A''₄7aq.—PbA''aq.—PbA''aq.—PbA''₄2aq.—RjA''₄-Sq.,—NiA''2aq.—KHA'',—KA''3aq.—Ag_A''.—SrA''4aq.—ZnH₂A''₂4aq.—Zn₃ON'₄8aq.

Ise-allexante acid C,H,N,O,. Obtained by the action of alkalis upon the red substance got by heating alloxan at 260° (L. Hardy, A. Ch. [4] 2, 372). A similar body may be got by the action of bromine-water on uric acid (Magnier de la Source, Bl. [2] 22, 56). Its solution then gives with baryta-water a splendid violet pp. of baric iso-alloxanate, which, however, when exposed to moist air soon changes to colourless baric alloxanate.

Salts.—(NH₁)₂A": red powder: v. sol. water forming a purple solution, which gives with AgNO₂ an indigo blue pp., and with K₂CO₃ a violet colour.

ALLOXANTIN C₈H₄N₄O₇ 3aq. (Liebig a. Wöhler, A. 26, 262; Fritzsche, **J. pr.**

14, 237).

Formation.—I. By action of warm dilute HNO₂ on uric acid.—2. By action of electrolysis or of reducing agents on alloxan (q. v.).—3. By dissolving alloxan in a concentrated angeous solution of dialuric acid: C₄H,N,O₄ + C₄H₄N,O₄ + C₄N,O₄ + C₄N,O₄

Properties. Small oblique rhombic prisms.

Reddens litmus. V. sl. sol. cold water, Gives | class (1), when alloyed with each other, gives with baryta-water a violet pp. Reduces AgNO₂. products the conducting powers of which for Reactions—1 44 170° class hydratile acid. with baryta-water a violet pp. Reduces AgNO.

Reactions.—1. At 170° gives hydurilic acid,
oxalic acid, CO., CO. and NH.—2. Oxidation

gives alloxan .- 3. Reduction forms dialurio acid .- 4. Ammonia gas turns it red, forming mnrexide.-5. Aqueous ammonia forms a purple solution, long boiling bleaches it, uranil being formed. This is then converted into murcxido by atmospherio oxidation .- 6. The purple pp. produced by baryta-water disappears on boiling, bario alloxanato and dialurate being formed.

ALLOYS .- The word allow was originally employed to designate the product obtained by mixing gold or silvor with other metals; its application is now general, all mixtures or combounds of metals with each other being named alloys, except those containing mercury, which are termed 'amalgams.' For a detailed description of special alloys, reference must be made to one of the constituent metals; only tho general properties of the alloys will be here

considered.

On melting two metals together, or on meltng one and adding the other, complete assimiation takes place in some cases and not in thers. Thus, silver easily mixes or alloys with old, copper, or lead; but neither silver nor copper can be readily induced to unito with iron. In the cases of those metals which do not completely mix when inclted together it usually happens that a small quantity of one is taken up by the other; thus, Faraday and Stedart found that iron is able to absorb the of its weight of silver with production of a homogeneous alloy, the properties of which are considerably different from those of iron; but that if more silver than fath of the mass of the iron is present, tho greater part of the silver separates during cooling. and that which remains can be detected by the microscope. If silver is melted with addition of a small quantity of iron, the latter metal alloys to some extent; but it is impossible to obtain mixtures of these metals in any desired proportion. On the other hand, silver and copper, or silver and gold, form alloys in which the proportion of the two metals may be varied at will.

The physical properties of alloys are in some cases nearly the mean of those of their constituent metals; but in other cases a wide difference is observable between the properties of the alloy and the properties of the metals which have been used to form it. Matthiessen, to whom we owe most of our knowledge of the properties of alloys, divides all metals into two classes: (1) those which impart to an alloy their own physical properties, to a less or greater degree, according to the proportion in which they themselves exist in the alloy; and (2) those which do not come under class (1). To the first class belong the metals lead, tin, zinc, and cadmium; and to the second, in all probability, the other metals. The alloys themselves may also be divided into three groups: (a) those made of the metals belonging to class (1), (b) those made of metals of class (1) with class (2); and (c) those made of class (2) with one another. This classification is largely based on the relative conductivity for electricity of the metals and of the alloys which they form with each other.

heat and for electricity are proportional to the relative quantities by volume of the constituent metals; but that this is not the case with alloys of the metals of class (1) with those of class (2), nor with the alloys of metals of class (2) with each other. As regards conductivity for heat and for cloctricity, Wiedemann and Franz have added to our knowledge by showing that the conducting powers of metals and their alloys for heat vary in a similar manner to that in which their conductivity for electricity varies. This statement has been confirmed and amplified by

Matthiessen regards alloys of the metals of class (1) as solidified solutions of one metal in the other; but supposes that metals of class (2) enter into alloys in an allotropic form; and he further supposes that when metals are alloyed together one or more of the metals may nadergo allotropic change. Thus, ho regards as solidified solutions of the metals, alloys of lead with tin, cadmium with tin, zinc with tin, cadmium with lead, zine with cadmium, and zine with lead. He supposes that in the alloys of lead or tin with bismuth, tin or zinc with copper or with silver, one metal is dissolved in an allotropio modification of the other; and that in alloys of bismuth with gold or silver, palladium or platinum with silver, or of gold with copper or silver, both metals exist in allotropic forms. Matthiessen does not, however, ignoro the fact that certain alloys contain their constituent metals in simple atomic proportions; for example, the alloys whose composition may be expressed by the formulæ AuSn, AuSn, and AuSn; but he regards alloys of intermediate composition as solidified solutions of such definite compounds in each other. It is known that ziuc will not alloy with more than 1.2 p.o. of lead, nor will lead alloy with more than 1.6 p.c. of zine; yet, by stirring, it is possible to ob-tain mechanical mixtures of such alloys with excess of one or other metal. Such mixtures are placed by Matthiessen in a class by themselves. Most of the alloys of silver and copper with each other are regarded by him as mixtures of various solidified solutions. The hypothesis of the existence in an alloy of one of the constituent metals in an allotropic form has roceived a certain degree of confirmation from experiments by Deville and Debray, who have observed that the ridium separated by the action of an acid on an alloy of that metal with zinc explodes when heated to 300°, and is changed by the explosion into ordinary iridium. Wiedemann has suggested that the contraction of alloys after solidification, which sometimes goes on for days, is due to the gradual occurrence of an allotropic change in the constituent metals, one modification being stable at high, and the other at low, temperatures. If the hypothesis of the occurrence of allotropic change during the formation of certain alloys is tenable, it is remarkable that such allotropic modifications of metals should be producible by pressure; for Spring has succeeded in producing Wood's alloy (containing bismuth, cadmium, and tin), and also brass, but the latter only partially, by exposing mixtures of the metalic Matthiessen found that the metals placed in in fine powder to very high pressures.

On the whole, there appears to be a marked analogy between alloys and solutions. It is well known that the conductivity of water for electricity is nearly nil, but becomes considerable when the minutest trace of any salt is dissolved in it. Similarly, the conductivity of copper is greatly diminished by the admixture with it of minute quantities of other metals. Morcover, in many other cases a great modification is produced in the tenacity, mulleability, &c., of metals by very small additions of foreign substances; as, for example, by the addition of small quantities of carbon, silicon, sulphur, or phosphorus, to iron, of phosphorus to copper, or of magnesium to nickel. And just as an aqueous solution of a salt must be heated to a temperature higher than that of the boiling-point of water before the whole of the water is removed. so it has been found that alloys of zinc, sodinm, mercury, &c., must be heated to temperatures above those at which these metals volatilise before the metals in question are entirely removed from the alloys. The analogy between alloys and solutions has been strikingly shown by Guthrie. This physicist has found that that alloy of two metals which has the lowest meltingpoint does not contain the metals in atomic proportion, but is strikingly similar to an 'alloy' of two salts, such as that of nitrate of potassium and nitrato of lead. Alloys were obtained by him of bismuth and zino (Bi = 92.85 p.c. Zu = 7.15 p.c.), melting at 248° ; of bismuth and tin (Bi = 46.1 p.c. Sn = 53.9 p.c.), melting at 133° ; of bismuth and lead (Bi = 55.58 p.e. Pb = 41.42 p.c.), melting at 122.7°; and of bismuth and cadmium (Bi = 59.19 p.o. Cd = 40.81 p.c.), melting at 144°. None of these alloys contains the metals in the proportion of their atomic weights, and the melting-point of each alloy is the lowest of all possible alloys of the specified pair of metals. Such alloys are termed by Guthrie eutectic alloys; they appear to be in some sense solidified solutions, resembling cryohydrates. We are still ignorant of the true nature of such mixtures, if mixtures they bc.

Spring (B. 15, 595) has prepared several alloys by subjecting mixtures of the constituent metals to pressures of about 7,000 atmos. In this way he obtained brass, Wood's alloy (Bi, Cd, and Sn), and Rose's alloy (Bi, Pb, and Sn).

References.—Matthiessen, B. A. 1863, 37; and C. J. Trans. 1867, 201; also P. R. I. March 20th, 1868. Deville and Debray, C. R. 94, 1557. Spring, B. 15, 595. Wiedemann, W. 3, 237-250. Crookewit, A. 68, 290. Fürstenbach, Bayerisches Industrie- und Gewerbeblatt, 1869. Sundall, A. Ch. 119, 141. Crace-Calvert a. Johnson, A. Ch. 45, 454. Gutbrig, P. M. June, 1884.

ALLURANIC ACID C.H.N.O. (?). Formed by evaporating an aqueous solution of equivalent quantities of urea and alloxan (Mulder, B. 6, 1012). Orystals; sl. sol. water. AgA'2aq.

ALLYL.—The radicle CH₂:CII.CII., is called Allyl, the isomeric radicle CH₃:CII:CH being termed Propenyl.

C.E. (6°-10°) 00188; (11.9°-59.8°) 00158. S.V. 125·8 (S.); 125·7 (Z.). V.D. 2·84 (for 2·84). H.F.p. -9260. H.F.v. - 11580 (Thomsen). \(\rho_8\) I-4079. R_{\infty} 45·99 (B.). Critical temperature 23·4·4°.

Formation.—1. From allyl iodide and Na (Berthelot a. de Luca, A. 100, 361), an alloy of sodium and tin (Würtz a. Leclauché, A. Ch. [4] 3, 155), or iron (Linnemann, Bl. [2] 7, 424).—2. By heating mercury allyl iodide, IHgC₂H₄, alone (Linnemann, A. 140, 180) or with aqueous KCy (Oppenheim, B. 4, 672).

Reactions.—1. Oxidised by chronic acid mixture gives carbonic and acotic acids.—2. Oxidised by KMnO, in neutral solution gives CO,, acetic, oxalic, and succinio acids.—3. Oxidised by KMnO, in acid solution gives CO, acetic acid, and succinic acid (E. Sorokin, J. pr. 131, 1).

Constitution.— The formation of acetic acid by exidation of di-allyl seems to favour the formula CH₂CH.CH.CH.CH.CH.CH₃; while the formation of succinic acid is more in accordance with the formula CH₂:CH.CH.₂CH₂·CH₂·CH.₂CH.₃CH.₄ formula that is further supported by the conversion of di-allyl into di-propargyl. The exalic acid may be supposed to be formed by exidation of the succinic acid. Acetic acid may be considered to be formed from intermediate hydrates, CH₂ CH₃(CH₂·CH₂·CH₂·CH₃·CH₃·CH₄·CH₅·CH₄·CH₅·

and CH₂.CH₁(OH).CH₂.CH₂.CH₂.CH₁(OH).CH₃.

These bodies do, in fact, yield acetic acid when exidised. According to Sabanceff (J. R. 1885, 35) di-allyl forms two tetrabromides and must therefore be a mixture of two hydrocarbons.

Combinations.—1. When gaseous H1 is passed into strongly cooled di-allyl, combination takes place, the product Cll_CHI,CH_CH_CH_CH_being formel, v. IODO-DENANES.—2. A smaller quantity of III forms the mono-hydro-iodide, CaH_all1, (165°), also formed from preceding by alcoholic KOII (v. IODO-HEXYLENE).—3. Similarly, fuming HCl forms two hydrochlorides.—4. HOCl forus CaU_m(HOCl), di-chloro-di-oxy-hexane (q. v.).—5. Br forms tetra-bromo-hexane.

ALLYL ACETATE C₁H₂O₂ i.e. C₂H₁C₂H₂O₄ Mol. w. 100. (103²-103·5²) at 735 mm. (R. Schiff, A. 220, 109), S.G. Ψ *9276 (Brühl). S.V. 121·37 (S.). μβ 1·4105. R_∞ 42·21 (B.). ALLYL-ACETIC ACID

C₂H₂O₂ i.e. CH₁:CH₂CH₂CH₂CO₂H. Pentenole acid. (185°-188° cor.). S.G. ½ 9866; ¼ 9842; ½ 9767. M.M. 6:426 at 14° (Perkin, C. J. 49, 211). Prepared by heating allylmalonic acid (Conrad a. Bischoff, B. 13, 598) or from allyl-aceto-acetic ether (Zeidler, B. 8, 1035). Combines with Br., or HBr. Not reduced by sodium-amalgam. Oxidised by chronic acid to specinic and formic acids.

Salts.—KA': scales; v. sol. water; solution not ppd. by Fc_Cl_{*}.—CaA'₂2aq: lamins.— BaA'₂2aq.—AgA' (Mosserschmidt, A. 208, 92). Eth cr.—EtA': (142°-144°).

DI-ALLYL-ACETIC ACID C. H_{1.}O₂ i.e., (C.11)_CH_CO_H. Octivoic acid. (220°) (C.a.B.); (218°-222°) (fl.); (224°-226°) (R.); (227° cor.), S.G. 12° 9576; 12° 9555; 32° 9191. M.M. 10·844 at 16·16° (Perkin, C. J. 49°, 212).

Formation. - From di-allyl-accto-acctio ether (Wolff, A. 201, 49; Rebonl, Bl. [2] 29, 228) or from di-allyl-malonic acid (Conrad a. Bischoff,

(q. v.) by reduction (Schatzky, J. R. 17, 79). Properties. - Oil, of disagreeable odour.

Insol. water. Volatile with steam.

Reactions.—1. Conc. HBr forms, probably, an addition product (CH, CHBr.CH.), CH.CO.H which instantly splits off HBr forming

v. Bromo-oxy-octore acid (Hjelt, A. 216, 73).-2. Br in CHCl, forms, probably, an addition compound, (CH_Br.CHBr.CH_)_CH.CO.H, but this instantly splits up into HBr and a lactone .CH_..CH.CH_Br

CH_Br.CHBr.CH_.CH CO.0

v. Tri-BROMO-OXY-OCTOIC ACID. -3. HNO, (S.G 1.3) forms tri-earballylic acid (W.).

Salts.—CaA', 2aq: leaflets. -- AgA', S. ·41 at 15°. Ether.—EtA' (195°) (R.).

ALLYL-ACETO-ACETIC ETHER v. pp. 23, 25. ALLYL-ACETONE

C₆H₁₀O i.e. CH₂:CH.CH₂.CH₂.CO.CH₃

Methyl butenyl ketone. (129°). S.G. 27 834. From allyl-aceto-acetic ctlicr (Zeidler, A. 187, 35). Unpleasant smell. Forms with NaHSOs an amorphous compound, C₆II,₁₀O 2NaHSO₃ (O. Hofmann, A. 201, 81). Reduced by sodiumamalgam to hexenyl alcohol (q. v.).

DI-ALLYL-ACETONE

C₉II₁₁O i.e. (C₃H₃)₂CH.CO.CII₃. (175°). From di-allyl-aceto-acetic ether (Wolff, A. 201, 47).

ALLYL-ACETOPHENONE v. PHENYL BUTENYL

ALLYL-ACETOXIM

C₆H₁₁ON i.e. C₅H₅.CH₂.C(N.OH).CH₂ (188° corr.). Formed by the action of hydroxylamine on allyl-acetone. Liquid. Soluble in alcohol, ether, benzene, CS., ligroine, acids, and alkalis. By aqueous acids it is resolved into its constituents. It combines with bromine to form a di-bromide (Nageli, B. 16, 496).

ALLYL ALCOHOL C₂II₄O i.e. CH₂: CH.CII₂OII. [-50°]. (96.6°). S.G. 0.8706; 15.5 .8576. S.V. 74.19. C.E. (0°-20°) 00104 (Thorpo, C. J. 37, 208). S.H. 6569 (Reis, P. [2] 13, 447). H.F.p. 31,200 H.F.v. 29,750. Ro 27.09 (Brühl, A. 200, 175). H.F.p. 31,200.

Occurrence.—Crude wood spirit contains not more than one-fifth per cent. (Aronheim, B. 7, 1381; Grodzki a. Kramer, B. 7, 1492).

Formation. -- 1. Dry gascous ammonia is passed into oxalate of allyl till a solid mass of examide, saturated with allyl alcohol, is obtained. The latter is then distilled off (Zinin, A. 96, 362).—2. Produced, together with isopropyl alcohol and acropinacone (q. v.), when acrolein is treated with zinc and hydrochlorie acid (Linnemann, A. Suppl. 3, 257). 3. By the action of sodium on dichlorhydrin (Hübner a. Müller, Z. 6, 344). — 4. The two atoms of chlorine may also be removed from dichlorhydrin by sodium-amalgam (Lourenço, A. Ch. [3] 67, 323), or by copper and potassic iodide (Swarts, Z. 1868, 259).—5. Allyl iodide (1 pt.) is heated with water (20 pts.) for 60

R. 18, 598). From iodo-di-allyl-scetic acid hours in a soda-water bottle at 100°. The yield is excellent (Niedsrist, A. 196, 850).

Freparation.—Glycerine (400 pts.) is slowly distilled with orystallised exalio acid (100 pts.) and a little ammonie chloride (1 pt.), to convert any potassio oxalate into ehloride. receiver is changed at 190°, and the distillation continued up to 260°. The distillate, containing aqueous allyl alcohol, allyl formate, acrolein, and glycerin, is rectified, dried, first with K2CO2, then over solid potash, and distilled. It then boils at 90°, but whon the last traces of water are removed by quicklime, it boils at 96°. The yield is one-fifth of the weight of oxalio acid used (Tollens a. Henninger, Bl. [2] 9, 394; Brühl, A. 200, 174; Linnemann, B. 7, 854).

Theory of the Process. — Carbonio acid is first evolved freely (at 130°), but formie acid which must be produced at the same time (H₂C₂O₄ = CO₂ + H₂CO₂) reacts upon glycerin, producing monoformin: $C_3H_1(OH)_3 + H_2OO_2$ = $H_1O + C_3H_3(OH)_2(OCHO)$. The monoformin can be extracted with ether, and boils about 165° in vacuo. When distilled, monoformin splits up into allyl alcohol and carbonic acid:

 $C_3H_3(OH)_3(OCHO) = CO_2 + H_2O + C_3H_3(OH)$ (Tollens, A. 156, 140). When a large quantity of oxalio acid is used, the excess of formic acid does not produce diformin, but comes off as

formic acid (q. v.).

Properties. - A pungent liquid, with a burning tasto. It mixes with water, alcohol, and ether. Constitution .- That allyl alcohol has the formula CH2:CH.CH2OH and not CH3.CH:CH.OH may be inferred from the fact that it yields no acetio acid when oxidised by nitric acid. A similar remark applies to allyl iodido (Kekulé a. Rinne, B. 6, 386).

Reactions .- 1. Chromic acid oxidises it to CO2 and formic acid; no acrylic acid is formed, but a pungent odour, which may be due to aerolein, is observed (Hofmann a. Cahours, A. 100, 257; Rinno a. Tollens, A. 159, 110).—2. When allyl alcohol is heated, with inverted condenser, for 5 hours in a water-bath with zinc and dilute H2SO, about 16 p.c. is reduced to n-propyl alcohol: CH,:CH.CH.OH + H, == CH3.CH_CH_OH (Linnemann, B. 7, 862).—3. Solid potash at 100°-150°, in a flask with inverted condenser, forms n-propyl alcohol (by reduction), formic acid (by oxidation), ethyl alcohol, hydrogen, and other products (Tollens, A. 159, 92).—4. Potassium displaces hydrogen, forming gelatinous potassic allylate .- 5. H SO. forms (C3H3)HSO, -6. Dilute H2SO, or HCl at 100° forms an aldeliyde, C.H.O (c. 137°) (Solonina, J. R. 1885, i. 145)

Combinations .- I. With chlorine it forms u-dichlorhydrin CH_Cl.CHCl.CHI_OH (q. v.)--2. With bromine it forms a dibromide, called also dibromhydrin, CH, Br. CHBr. CH, OH, (214°). 60 grms. Br are dissolved in 300 grms. CS, and dropped slowly (in 4 hours) into a solution of 20 grms. of allyl alcohol in 100 grms. of CS, The product is distilled in vacuo (Michael a. Norton, Am. 2, 16; compare Kekulé, A. Suppl. 1, 138; Markownikoff, J. 1864, 490). Linnomann says there are two bromides (B. 7, 859). 3. When iodine is added to a solution of allyl alcohol in CHCl, it combines, and on evapora-

tion OH, LOHI.CH, OH separates as needles. Dilute Na CO, converts this into iodallyl alcohol [160] (Hübner a. Lellmann, B. 14, 207).-4. ICl unites, forming C3H5IOl(OH) (Henry, B. 3, 351). 5. With cyanogen it unitos, forming C₃H₃(CN)₂(OH), (151°) (Tollens, B. 5, 1045). -6. BaO combinos, forming BaO, 2C₃H₂O. -7. IJClO unites, forming a little chlorhydrin (q. v.).-8. Chloral combines with allyl alcohol; the compound, CCl3.Cl1(Ol1)(OC3II3) [20.5°], (116°), is analogous to chloral alcoholate (Oglialoro, G. 4, 463).

DI.ALLYL-p-AMIDO-BENZOIC ACID

C₁₃H₁₅NO₂ i.e. (C₃H₅)₂N.C₆H₄.CO₂H [127°]. From allyl iodide and potassic p-amido-benzoate (Miohael a. Wing, Am. 7, 198).

Di-allyl-m-amido-benzoic acid [90°] (Griess,

B. 5, 1041).-- IIA'HCl aq.

DI-ALLYL-AMIDO-ETHYL ALCOHOL v.

OXYETHYL-DI-ALLYLAMINE.

ALLYLAMINE C3H,N i.e. CH2:CH.CH2.NH2 Mol. w. 57. (56°) (R. Schiff, B. 19, 565); (58°) (Oeser, A. 134, 7). S.G. 12-864 (O.) S.V. 78·38 (S.). 1LF.p. - 1140. H.F.v. - 2880.

Formation .- 1. From allyl cyanato (Cahours a. Hofmann, A. 102, 301).—2. From oil of mustard, Zn, and 1ICl (0.).—3. From oil of mustard and conc. H2SO, (Hofmann, B. 1, 182;

Rinne, A. 168, 262).

Properties. - Liquid with pungent ammoniacal odour. Miscible with water. Strong base.

Dissolves ppd. Cu(OH)2 and Ag2O.

Reactions.-1. Combines with bromine.-2. H₂SO₁ at 140° forms a compound which, on pouring into water, produces oxy-propyl-amino 110.C₂II₆.NH,

Salts. - (B'HCl).PtCl,: monoclinic tables. Changed by boiling into (B'HCl).PtCl, (Liebermann a. Paal, B. 16, 530). - B', H. SO, (Andreasch, M. 5, 33).

(C,H,),NH

Di-allyl-amine U₆₂₁₁₁. (111°) (Ladenburg, B. 14, 1879). Tri-allyl-amine C_bH₁₃N i.e. (C₃H₅)N. (156° i.V.) S.G. § 8206. S.Y. 200·3 (Zander). C.E. (0°-10°) ·00103. Formed when tetraallyl-ammonium hydroxide is distilled (C. a. H.; Pinner, B. 12, 2054; Grosheintz, Bl. 31, 391).-B'HCl.-B'.H.PtCl.

Tetra-allyl-ammonium hydrate (C,H,),NOH: liquid. — $((C_3H_5)_4NCI)_2PtCl_4$. — $(C_3H_1)_1NBr$. — $(C_3H_5)_4NI$. The three last are crystalline

(C. a. H.).

ALLYL-AMYL-AMINE C,H,,N i.e.

(C₃H₅)(C₅H₁₁)N1I. (c. 150°). S.G. 18 .777. From amyl bromide and allyl-amino (Liebermann a. Paal, B. 16, 531)

ALLYL AMYL OXIDE C, II, O i.e. C, II, O.C, II, (120°) (Berthelot a. de Luca, A. Ch. [3] 48, 292).

ALLYL-ANILINE C, H, IN i.e. PhN(C, H,)11.

(209°). S.G. ²² ·982 (Schiff, A. Suppl. 3, 364).
Di-allyl-aniline C₁₂II_{1,N} i.e. Pl\N(C₃II₅)₂.
(244°). S.G. § ·9680. S.V. 225·2° (Zander, A. 214, 149). C. E. (0°-10°) ·00083.

ALLYL-BENZENE C, H, oi.e. Ph. CH: CH. Cll (?) Phenyl-propylene. Propenyl-benzene.

118. (175°) (P.); (178°) (E.). S.G. 15 92.

Formation. — 1. By-product in action of sodium-amalgam on warm aqueous cinnamyl alcohol (Fittig a. Krügener, B. 6, 214; Rüglieimer,

C. R. 78, 1153): PhO.H.Br = PhC.H. + HBr. So prepared it boils at (165°), and its di-bromide forms needles.—3. From brome-hydre phenyl-orotonic acid (Perkin, C. J. 32, 660).—4. From chloro-propyl-benzene and alcoholio potash (Errera, G. 14, 504).

Di-bromido C_aH₁₀Br₂ [66.5°]. Plates or edles. On distillation it yields an allylneedles. benzene (178°), which polymerises forming a viscid solid (330°).

Iso-allyl-benzene Ph.CII..CII:CII.(?) (155°). Chojnacki (C. R. 76, 1413) got this body from allyl iodide or bromide, benzene, and zine dust at Others have failed to get it. Allyl chloride, benzene, and Al.Cl, give di-phonyl propane, CH2.CHPh.CH.Ph, and n-propyl-benzene (q. v.) (Wispek a. Zuber, A. 218, 378).
ALLYL BENZOATE v. BENZOIC ACID.

ALLYL-BENZOYL-ACETIC ACID C12II,2O, i.e. BzCH(C₃II.)CO.H [1223-1253]. From benzoylacctic ether, NaOEt, and allyl iodidc. The resulting ether is saponified by standing for three weeks with dilute alcoholic KOII (W. H. Perkin, jun., C. J. 45, 186; 47, 240). Colourless crystals.

Reaction .- Boiled with dilute alcoholic KOH forms phenyl butenyl ketone (q. v.), benzoic acid

and (probably) allyl-acetic acid.

Ether.—EtA' (220') at 100 mm.; (241°) at 225 mm. Combines with Br.

ALLYL BORATE C.H., BO, i.e. (C.H.), BO, $(168^{\circ}-175^{\circ})$. From B_2O_3 and ally alcohol at 130° (Councler, J. pr. [2] 18, 376). Combines with bromine, forming (C₂H₁Br₂)₃EO₃.

ALLYL BROMIDE C₂H₂Br i.e. CH.: CH.CH Br

(71°). S.G. § 1·459; 18 1·436. S.V. 90·5 (Zander, A. 214, 144). C E.(0°-10°) ·00123. H.F.p. -340; H.F.v. -1500.

Formation.-1. From allyl alcohol, bromine, and phosphorus (Tollens, A. 156, 152).-2. From glycerine and PBr₄ (Henry, Z. [2] 6, 575).— 3. From allyl iodide and cupric bromide: $2C_3H_5I + 2CuBr_2 = Br_2 + Cu_2I_2 + 2C_4H_5Br$ (Oppenhenn, B. 3, 442).

Preparation .- Potassic biomide, hydric sulphate (2 pts.), and water (1 pt.) are warmed till hydric bromide begins to come off. Allyl alcohol is then dropped in (Grosheintz, Bl. 30, 98).

Combinations. - 1. With concentrated hydric bromide forms a mixture of propylene bromide (CH, CHBr.CII.Br) and trimethylene bromide (CH,Br.CH, CH,Br) which may be separated by distillation (Geromont, A. 158, 369).-2. With dry HBr it forms chiefly trimethylene bromide (q. v.).—3. With bromine it forms tribromhydrin (q. v.).—4. With ICl it forms C₂H₂IClBr.—5. With ClBr at 20° forms C₂H₂ClBr., but at 100° forms C₃H₂ClBr., but at 100° forms C₄H₂ClBr., but at 100° forms C₄H₂ClBr., but at 100° forms C₄H₂ClBr. 6. With IIClO it produces C, II, (OH) BrCl.-7. It combines with NEta.

ALLYL BUTYRATE v. BUTYRIC ACID. ALLYL-180-BUTYL-MALON1C ETHER C₁₄II₂₁O₄ i.e. (C₃II₃)C(C₄H₆)(CO₄Et)₂ (247°-250°). From di-sodio-malonic ether, allyl iodide, and

iso-butyl iodide (Ballo, B. 14, 335). On saponification it gives an acid [129°] which appears to bo propyl-iso-butyl-malonio acid.

ALLYL CARBAMINE C,H,N i.e. Cll₂:Cll.CH₂.NC (96°-106°). S.G. 11 .794. Produced by the action of silver cyanide on allyl A. 172, 129) .- 2. From propyl-benzene by Br at iodide. It is a liquid of disagreeable odour 160° and distilling the product (Radziszewski, Somewhat soluble in water (Lieke. A. 112, 316).

DI-ALLYL CARRINGL o. HEPTINYL ALCOHOL. DI-ALLYL-DI-CHLORHYDRIN C.H., Cl.O te. C.H₁₆(OH), Cl. From HClO and di-allyl (Przybytek, B. 18, 1350; Lauch, B. 18, 2288).
ALLYL CHLOBIDE C.H.Cl., C.H.; C.H.CH.Cl., Cl.

(46°) (Thorpe); (44.6°) at 744 mm. (Brühl). S.G. § 9547; 3 9379 (Brühl). C.E. (0°-10°) 00137. S.V. 84.7 (Zander). S.H. :3984 (Reis). μ_{8} 1·4225. R_{∞} 32·63 (Brühl). H.F.p. 7100. H.F.v. 5940. M.M. 6·008 at H.F.p. 19·6°.

Formation .- 1. From allyl iodide and HgCl.. From allyl oxalate, calcie chloride, and alcohol (Oppenheim, A. 140, 205).—3. From allyl alcohol and 11Cl in scaled tubes.

Preparation. - From allyl alcohol and PCl,

(Tollens, A. 156, 154).

Properties.-1. Alcoholic potash, oven below 100°, converts it into ethyl allyl oxide. Tho isomeric chloropropylene (26°) is converted by alcoholic potash at 120° into allylene. - 2. HClO unites, forming unsymmetrical dichlorhydrin, CH.Cl.CHCl.CII.OII, or dichloride of allyl alcohol. This body, when exidised by IINO, is converted into dichloropropionic acid (Henry, B. 7, 757).—
3. HCl combines, forming CH, CHCL.— 4. HBr forms CH.Br.CH.CH.Cl, together with a little CII, CHBr.CII Cl.-5. Warmed with H2SO, and then diluted and distilled, propylene chlorhydrin is produced (Oppenheim, A. Suppl. 6, 367).-6. Bromine combines, forming CaH, Cl. Br. -7. With polassic cyanide in presence of dilute alcohol it forms chiefly pyrotartaric acid, also propylene cyanide (Claus, A. 191, 38) and triallylamine (Pinner, B. 12, 2053). The reactions in this case are: (a) CH₂:CH₂CH₂CH₃:CH₄: (b) CH_:CH_CH_CN + HCN = CH_.CH(CN).CH_CN (c) CH₂.CH(CN).CH₂CN + 2KOH + 2H₂O == 2NH₃ + CH₃.CH(CO₂K).CH₂.CO₂K (pyrotartrate). The liberated ammonia forms the trially lamine. 8. With benzene, in presence of alumininn chloride, forms diphenylpropaue: C3H3Cl+2C1H6

= HCl + (C.H.), C.H. (Silva, C.R. 89, 60b). ALLYL-PSEUDO-CUMYL-PHTHAL-AMIDE. C₂₆H₂₆N₂O₂i.e.C₆H₂Me₃·NH.CO.C₆H₄·CO.NHC₄H₃. [179°]. Silky needles. Easily soluble in alcohol. Formed by the action of allylamine on phthalpseudo-cumidine (Fröhlich, B. 17, 1808).

ALLYL CYANAMIDE

C₄H₄O₂ gaq. i.e. (CN.NHC₃H₃)_x. Sinamine [100°]. From allyl-thio-nrea and Pb(OH)₂ or HgO. (Will, A. 52, 15; Andreasch, M. 2, 780; Robiquet a. Bussy, J. pr. 19, 234). Alkacompounds with HgCl., PtCl., and oxalic acid.
ALLYL CYANATE C, 11 NO i.e. C, H, N.CO.

Allyl carbinide. (82°). V.D. 3.05 (for 2.88). From allyl iodide and silver oyanate (Calionrs a. Hofmann, Tr. 1857, 555).

ALLYL CYANIDÉ C.H.N i.e. CH., CH.CH.CN. Orptonitrile. Mol. w. 67. (119° cor.). S.G. 2-8491;

*15 8351; 12 9: 8398.

Formation. - 1. By ppg. potassic myronate (q. v.) with silver nitrate and treating the pp. with hydric sulphide C.H.Ag.NS.O., +H.S.S.O.,+H.S.S.+B.+H.SO.,-2. During the fermentation of black mustard. -3. From allyl mustard oil by zine dust: C.H.NCS + $Zn = ZnS + C_3H$, CN (Sohwarz, B. 15, 2508).—4. From allyl su phocyanide and sodium (Billoter, B. 8, 465). thelot). H.F.v.-41530 (Th.).

Preparation.—Allyl lodids is hested with KCy for two days at 110°. The product is washed, dried, and heated again with KCy at It is then washed, dried over CaCl, distilled, freed from carbamine by shaking with a little HNO, and rectified (Rinno a. Tollens,

A. 159, 106).

Properties. - Liquid smelling faintly of garlic. Reactions. -1. Aqueous or alcoholic potash forms NH₂, and solid crotonic acid [72°]. The formation of this crotonic acid may be explained by the assumption that B-oxybutyrate is first of the assumption that p-daynoty are in a formed: CH.:CH.CH. F. KOII + 2II.O = NH₄ + CII₂.CII(OH).CH.:CO.K, and that this splits off water: CII₂.CII(OH).CII₂.CO₂K = II_O + CH_3.CH.CO.K, forming potassic crotonate. This view is supported by the behaviour of allyl cyanide towards IICl.-2. With fuming hydric chloride at 60° it forms B-chlorobutyrie acid: CII,:CII.CII,.CN + 2IICI + 2II,O == NII, CI + CII, CIICI.CII, CO, II. This is an unstable acid, which easily changes to crotonic acid. -- 3. IINO, forms acetic and oxalic acids. --CrO₃ forms acctic acid.

Combinations.-1. With alcohol. potassic oyanide acts on allyl iodide in alcoholic solution, a compound of allyl cyanido and alcohol, of boiling point (174°), is obtained: CII_a : CH: CH: CN + IIOEt = CII_a : CII(OEt): CII_a : CN. Saponified by strong HCl, this forms ethoxybutyramide, CH₂.CH(OEt).CH..CONH, [71°], which, when warmed with 11Cl, gives ethoxybutyric acid, CII, CII(OEt).CII CO II, boiling about 215°. Saponified by potash, the compound of allyl cyanide and alcohol (β-ethoxybutyronitrile) gives ordinary crotonic acid as follows: CH₂.CH(OEt).CH₂.CN + KOH + H₂O = NII, + HOEt + CII, CH.CO.K (Rinne, 6, 389). Dry hydrogen chloride converts the compound of allyl cyanide and alcohol into the chloride of \$\beta\$-chlorobutyrimid-other (166°), CII3.CHCl.CH., C(OEt)NII. This last compound is converted by alcoholic potash into crotonio

acid (Pinner, B. 17, 2007).

2. With allylatcohol. A similar compound, CH₃·CH(OC₃H₅).CH₂CN (96°), is formed when KCy acts on allyl chloride mixed with allyl alcohol.

Constitution. - From its mode of preparation. allyl cyanide ought to be CH.:CH.CH.CN, but from its reaction with potash it should be the nitrilo of ordinary erotonic acid CII, CII.CN. From the fact that allyl eyanide and crotonio acid both produce acetic acid on oxidation, while allyl iodide and allyl alcohol yield no acctic acid, Kekulé assumes the presence of a methyl group in the two former and its absence in the two last named (B. 6, 386). This reasoning seemed conclusive until the experiments of Pinner, mentioned above, showed that, when the eyanide is converted into crotonic acid by hydric chloride, an intermediate compound B-chlorobutyric acid) is produced, and it is therefore possible that when nitric or chromic acid is used, an unstable derivative of butyric acid (say, \$\beta\$-oxybntyric acid) is first formed, and that it is this which gives acetic acid on oxida-

ALLYLENE C.H. i.e. Mc.C:CH. Mcthylacetylene. Tropinens. Mol.w. 40. S. (other) 30 at 16°. H.F.p. -39950 (Thomsen); -37500 (Ber-

Formation. - 1. By action of alcoholic NaOH upon bromo-propylene (Sawitsch. C. R. 52, 399), chloro-propylene, or propylene bromide (Mias-nikoff, A. 118, 332).—2. By the action of Na upon CH₂CCl₂CHCl₂ (Borsche a. Fittig, A. 133, 111), CH₂Cl.CCl₂CH₂Cl, or CH₂CCl.CH.Cl (Pfoffer a. Fittig, A. 135, 357).—3. By cled rolysis of calcium incsaconate or citraconate (Aarland, J. pr. [2] 7, 142) .- 4. By heating (citra-) bromo-pyrotartarie anhydride with ammonineal AgNO, Åq at 130° (E. Bourgoin, C. R. 85, 710). Colourless gas, with unpleasant smell; burns with snicky flame. V. sol. alcohol, sol. water.
Reactions.—1. Ammoniaeal cuprous chloride

gives a canary coloured pp.-2. Absorbed by cone. H.SO, much more readily than acetylene, allylene sulphonic acid, Call SO II, being produced. An aqueous solution of this acid, when heated, yields inesitylene and acctone (A. Schroke, B. 8, 17, 367).—3. Aqueous mercuric salts form pps. containing the mereurie salt, HgO, and allylene. These pps. are decomposed by acids with formation of acetone (Kutscheroff, B. 14, 1541; J. R. 1882, 326).— 4. KMnO Aq forms, in the cold, malonic, oxalic, and formic acids (Berthelot, A. Suppl. 5, 97). -5. CrO, Aq forms propionic acid (Berthelot, A. Suppl. 8, 47).

Metallic Derivatives. - CH3.C:CNa: white powder, decomposed by water into NaOH and allylene (Berthelot, A. Ch. [4] 9, 395; J. R. 12, 288).-(C3113)2Hg: crystalline pp. formed by passing allylene into Nessler's solution (Kutscheroff, B. 17, 25).

Combinations. -1. Cold furning IICl forms EH, CCl, CII, ; 11Br, and III act similarly .- 2. Bromine forms di-bromo-propylene (q. v.) and tetra-bromo-propane (q. v.). Iodine acts simi-

Iso-allylene CH₂:C.CH₂. 1. Formed by electrolysis of potassic itaconate (A.).—2. By the action of sodium on di-chloro-propylene, CHCl:CH.CH.Cl (from symmetrical tri-chlorhydrin, Hartenstein, J. pr. [2] 6, 295).

Properties .- A gas that does not pp. ammoniacal silver or cuprous solutions. Forms a

tetrabromide.

Di-allylene C₀H₈ or CII : CII.CH ... CH ... C: CII Allyl-allylene (70°). S.G. ¹⁸ :858. V.D. 2·79 (for 2.76). Allyl-acctone is converted by PCl, into CaH, CH, CCl, CH, which is converted by alcoholic KOII into di-allyleno (L. Henry, C. R.

Reactions .- 1. Aqueous silver nitrate gives a pp. C_sH_sAg aq.—2. Ammoniacal cuprous chloride gives a canary-yellow pp. C_sH Cu aq. 3. Alcoholic AgNO₄ gives a pp. C₆II,AgEtOH.— 4. Bromine forms C₆II,Br₆.

Iso-allylene tetra-carboxylic acid v. Propane

TETRA-CARNOXYLIC ACID.

ALLYLENE DI-CHLORIDE v. DI-CHLORO-

PROPYLENE.

ALLYLENE OXIDE C.II.O (63°). by oxidising allylene with CrO, Aq (Berthelot, Bl. 14, 116). Pungent neutral liquid. Not attacked by baryta-water at 150° or by KOHAq at 300°. Reduces AgNO, Aq.

ALLYL ETHANE v. PENTINENE. ALLYL ETHER v. ALLYL OXIDE. ALLYL ETHYL OXIDE v. ETHYL ALLYL OXIDE.

DI-ALLYL HYDRATE v. HEXENTL ALCOHOL. DI-ALLYL-DI-HYDRATE v. DI-OXY-HEXANS. ALLYLIN v. GLYCERIN.

ALLYL IODIDE C.H.,I i.e. CH.,CH.CH.(102.7° i. V.). S.G. 2 - 8696. C.E. 00106. S.V. 100.9 (Zander, A. 214, 145); V. D. 5.77 (obs.).

Formation.-1. Allyl alcohol, P, and iodine (Tollens, Bl. [2] 9, 396).-2. Glycerin and PL. (Berthelot a. de Luce, A. Ch. [3] 43, 257) .- 3. Glycerin distilled with hydriodic acid; excess of the latter is to be avoided, as it would convert the allyl iodide into isopropyl iodide:

 $CH_{J}:CH.CH_{J}I + 2HI = CH_{J}.CHI.CH_{J}I + HI =$ CII_a .CH: $CH_2 + I_2 + H1 - CH_3$.CHI. $CH_3 + I_4$

(Erlenmeyer, A. 139, 211).-4. From allyl chloride and calcie iodide (Romburgh, R. 1, 151;

Spindler, A. 231, 270).

Preparation .- 200 grms. of glycerin, previously dried by heating to 280°, are mixed with 125 grms, of iodine. The tubulus of the retort is connected by a flexible tube with a flask containing 40 grms, of clear phosphorus in small pieces; this flask has also a side tube through which carbonic acid is passed until the air is cleared out of the entire apparatus. The clear phosphorus is added to the contents of the retort by tilting the flask from time to time. Allyl iodide distils over rapidly. The addition of phosphorus takes about 21 hours. The distilla-tion is then continued until the contents of the retort begin to carbonise and vapours of aerolein are given off. The distillate is washed with dilute NaOH, dried over CaCl2, and rectified. Yield 110 grms. (98° 102°).

Theory of the Process .- It is usually held

that triiodhydrin is first formed:

 $CH_2(OH)$. $CH_2(OH) + P + I_3 =$ H₃PO₃ + CH_I.CHII.CH_I and that this splits up into iodine and allyl iodide: CH_1.CHI.CH_1 - 1, + CH_2. CH_CH_1. iodide:

But the fact that allyl alcohol accompanies the allyl iodide renders it quito likely that the intermediate body is diiodhydrin:

 $CH_2I.CHI.CH_2OH = I_2 + CH_2:CII.CH_2OH.$ The allyl alcohol formed in this way being converted into iodide by HI (Henry, B. 14, 403).

Reactions .- 1. Zine and HCl reduce it to propylene .- 2. Salts of silver form silver iodide and salts of allyl .- 3. Dry hydric iodide converts it into isopropyl iodide (Simpson, Pr. 12, 533). 4. With zinc cthul at 100° it forms amylene, pentane, and diallyl (Wurtz, C. R. 56, 387) .-5. With cacodylit reacts thus: As Me + 2C, H, I = $AsMe_1I + AsMe_2(C_3II_5)_2I$ (Cahours, A. Ch. [3] 62, 291). - 6. With dry copper zinc couple at 100° it forms diallyl: 2C₃H₃I + Zn = ZnI₂ + (C₃H₃)₂ - 7. With wet copper-zine couple it forms propylene: C₁H₁I + H₂O + Zn = IZnOH + C₃H₆.—8. With zine and alcohol (S.G. 805) it also forms propylene (Gladstone a. Triec. C. J. 27, 208). -9. With HgBr, at 200° it gives HgL, HBr and propane (Montgolfier a. Girand, B. 12, 1211) .- 10. Heated at 100° for a long time with water it forms allyl alcohol (q. v.) .- 11. With KCN and alcohol it forms a di-cyanide which, when boiled with KOH, produces potassic pyrotartrate (Claus, A. 191, 38).

Combinations.—I. With chloride of iodins it unites, forming C_xH_xL_zCl (205°-210°), a colourless oil (M. Simpson, Pr. 13, 540).—2. Bromine forms C, II, Br. -3. Morcury unites with it, form-

ing C.II.IIgI, mercuric-allyl iodide.

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ALLYL-MALONIC ACID

C.H.0., i.e. C.H.CH(CO.H). [103°].

(Conrad a. Bischoff, B. 13, 597; A. 204, 166; Hjelt, A. 216, 52). Lauge prisms. V. sol. water, alcohol, and ether. A. 180° it splits np into CO. and allyl-acetic acid. Combines with HBr forming liquid (CO.LI).CH.C.H.Br which, when boiled with water gives a lactone of oxy-propylmalonic acid (q. v.). Combines with Br. forming di-bromo-propyl-malonic acid (q. v.).

Salts.— ČaA": crystalline powder.— Ag₂A".

Elher.— Et.A" (218° 225°); (194°) at 330 mm.

S.G. ½; 1018 (Č. a. B.); ½; 1014. M.M. 11·28

at 13·7° (Perkin). From sodio-malonic ether

and allyl iodide.

Di-allyl-malonic acid

 $C_yH_{12}O_4$ i.e. $(C_3H_3)_2C(CO_2H)_T$. [133°]. Rhombic prisms; $a:b:c=9916:1:1\cdot0179$ (Haushofer, $Z.\ K.\ 11,\ 147$). Sol. water, alcohol, and ether; v. sl. sol. CS_2 .

Reactions.—1. Heat splits it up into CO₂ and di-allyl-acetic acid.

Ether Et. A". (240°) (C. s. B.); (203°) at 225 nm. S.G. 14°996 (C. a. B.); 15°1000; 25°993 (Perkin). M.M. 15 at 22°. From allyl iodide and sodiomalonic ether (Conrad a. Bischoff, B. 13, 598; A. 201, 171; Hjelt, A. 216, 61).

ALLYL MERCAPTAN C, II. SII. Mol. w. 74. (90°) (Hofmann a. Cahours, A. 102, 292).— C, H, SHgCl: pearly plates (from alcohol) (Gerlich, A. 178, 88).

ALLYL METHYL ETHER v. METHYL ALLYL

ALLYL MUSTARD OIL v. ALLYL THIO-CARBIMIDE.

ALLYL NITRATE C₂11,NO₃. (106°). S.G. 12 1·09. V.D. 3·54 (for 3·56). From allyl bromide and AgNO₃ (Henry, B. 5, 452).

ALLYL NITRITE C₃H₃NO₂ i.e. C₃H₃O.NO. (44°). S.G. 2-955. Prepared by distilling glyceryl tri-nitrite with allyl alcohol. An oil. Decomposed by MeOH into allyl alcohol and methyl nitrite. Its vapour explodes at 100° (Bertoni, G. 15, 361).

ALLYL OXALATE v. OXALIC ACID.

DI-ALLYL-OXALIC ACID (so called) v. Oxy-octinoic acid.

DI-ALLYL-OXAMIDE C.H., N.O. i.e. C.H., N.H.CO.CO.NIIC., II., [154°]. (274°). White plates. Soluble in hot water. Prepared by the action of allylamine on oxalic ether.

Tetrabromide C₂O₂(NHC₄II₃Br₂)₂. Insoluble in most ordinary solvents, except hot acetic acid (Wallach a. Stricker, B. 13, 513).

DI-ALLYL OXIDE C₆11₁₀O i.e. (C₃H₃)₂O.

Allyl elher. Mol. w. 98. (82°) (Cahours a. Hofmann, A. 162, 290); (94°3° i.V.) (Zander, A. 214, 146). S.G. 3°823. S.V. 35°5. C.E. (0°-10°) '00127. H.F.p. 12460. H.F.v. 9850 (Thomsen).

DLALLYL DI-OXIDE

• ĆH₂. CH.CH₂.CH₂.ĆH₃.ĆH₄ (180°). V.D.= 3.7 (obs.). Mobile colourless fluid of slight smell and burning taste. Heavier than water. Combines with acids, and pps. magnesia from a solution of MgCl₂. Obtained by the action of solid caustic alkalis upon diallyl-di-chlorhydrin. By boiling with water it is converted into the alcohol-oxide OH₂. OH.CH₂.CH₂.CH(OH).CH₂(OH), which only by long heating with water is converted into the tetra-hydric alcohol Cl1₂(Ol1).Cl1(OH).Cl1₂.CH₂.Cl1(OH).CH₂(OH) (Pryybytck, B. 18, 1350).

0-ALLYL-PHENÓL. Methyl derivative C₁₄H₁₂O i.e. C₄H₄(OMe)-CH₄-CH₄-CH₂ (233°). S.G. 15/5 ·9972; 36/9 ·984; 45/9 ·9793. Formed by the action of Na₂OO₃Aq on the product of the union of III with the methyl derivative of (α)- or (β)-oxy-phenyl-crotonic acid (q, v,). It is an oil; combines with bromine; forms a red solid with H₂SO₄ (Perkin, C. J. 39, 425).

p Ally1-phenol. Methyl derivative. (232°) S.G. $\frac{30}{100}$ 985. Prepared as above from corresponding p-compound. Anethol (q. v.) is isomeric with this body. Anol (q. v.) is isomeric

with allyl-phenol.

ALLYL-PHENYL-THIO-UREA C₁₀H₁₂N_{1.S} i.c. C₂H₂NH₁CS.N1ll'll. [98°]. S. (alcohol) 71 at 18°. From oil of mustard and amiline (Zinin, A. 84, 348); from allyl-amine and phenyl thio-carbimide (Weith, B. 8, 1529). Monoclinic crystals; v. sol. ether, insol. water. Gyanogen passed into an alcoholic solution forms C₁₀H₁₂N₂S(CN)₂, ppd. by water (Maly, Z. 1869, 261). When this is warmed with alcohol and dilute H.SO₄ it forms the oxalyl derivative. Oxalyl Derivative

CO.NC₅H₅ CS [161°]. Lemon-yellow needles.

insol. water, sl. sol. cold alcohol.

ALLYL-PHENYL-UREA C, II, N.O i.e. C, H, NH.CO.MIPh. [97°]. Needles. Got from its oxalyl derivative by baryta (Maly, Z. 1869, 263).

Oxalyl-derivative | CO-NC₃H₃ CO.

From the oxalyl derivative of allyl-phenyl-thiourea (q. v.) and warm AgNO₃ in alcoholic solution. Long needles. Insol. water, v. sol. alcohol, benzene, and CS₂.

ALLYL-PHTHALIMIDE v. Phthalio Acid, Allylamide.

ALLYL - PROPYL ALCOHOL v. HEXENYL ALCOHOL.

ALLYL-PROPYL-AMINE C₄11₁₃N i.e. C₃11.NHC₂H., (c. 112°). S.G. 18 = .7708 Colourless fluid. S. = about 6. Prepared by the action of prepyl bromide on allylamine.

Salts: B'_II_Cl_PtCl_: orange crystals.— B'II_C_O,: sl. sol. needles.—B'_II_C_O,*: thin plates (Liebermann a. Paal, B. 16, 525).

Allyl-di-propyl-aming (C₂/I₁)₂NC₂H₃. (c. 147°), Colourless fluid. S=about 2. Formed by the action of propyl bromide on allylamine.

Salts. B'zHzCl.PtCl.: orange-red trimetric crystals, a:b:c= 9831:1:1:1217.
B'HClPtCl.: sparingly soluble yellow needles [152°]; formed by boiling the preceding salt with water (Liebermann a. Paal, B. 16, 527).

ALLYL-ISO-PROPYL-BENZENE v. Propenyl-iso-propyl-benzene.

ALLYL DI-PROPYL CARBINOL *. DECENYL ALCOHOL.

Di-allyl propyl carbinol v. Decinyl alcohol. a-ALLYL-PYRIDINE C₃H₄(C₂H₅)N. (c.190°).

B.G. 2 9595. Colourless refractive oil; sl. sol. give immediate pps. with ammoniacal AgNO, or water. Prepared by heating pure (a)-picolino with paraldehyde for 10 hours at 250°-260°. On oxidation it gives picolinic acid [183°]. On reduction in alcoholio solution by means of sodium it yields (a)-propyl-pyridino (inactive contine).

Salts.—(B'HCl)_PtCl,: [186°], sparingly soluble needles.—B'HClAuCl,*: [136°], oily pp. solidifying to small needles.—(B'HCl)_PtgCl_*: very sparingly soluble crystalline pp. (Laden-

burg, B. 19, 2578).

-ALLYL-PYRROL C,H,N i.e. C,H,N.C,11, [105°] at 48 mm. Formed by the action of allyl bromide on pyrrol-potassium. Colourless oil. Volatile with steam. Almost insoluble in water. HgCl2 gives a white pp. (Ciamician a. Dennstedt, $B.\ 15,\ 2581;\ G.\ 13,\ 17)$

ALLYL-RESORGIN $C_aH_3(C_3H_3)(OH)_2$. Monomethyl ether $C_aH_3(C_3H_3)(OMc)(OH)$. $(245^\circ-250^\circ)$; V.D. 165 (obs.); colourloss oil (Pechmann a. Cohen, B. 17, 2132).

ALLYL-SUCCINIC ACID

C₂H₁₀O₄ i.e. CO₂H.CH₂.CH(C₃H₅).CO₂H. [94°]. Plates (from alcohol). Prepared by heating allyl-ethane tri-carboxylic acid to 160°, CO. being evolved. Strong aqueous HBr converts it into the corresponding lactonic acid-

Salts: A"Ca": crystalline, soluble.—A"Ba": easily soluble, amorphous.—A"Ag $_2$ *: sparingly soluble, amorphous.-FcSO, gives a flocculent pp. (Hjelt, B. 16, 334)

ALLYL SULPHATE C3H6SO4 i.e.

C.H.O.SO., OH. Hydrogen allyl sulphate. Allyl-sulphuric acid. From allyl alcohol and H.SO. (Cahours a. Hofmann, C. J. 10, 316).

Salts.—(Szymanski, A. 230, 43.) BaA'₂.—
SrA'₂.—CaA'₂ 2aq.—CuA'₂ 4aq.—PbA'₂PbO 6aq.—
MgA'₂ 4aq.—KA'.—NaA'.—NII₄A'.
ALLYL SULPHIDE C₄H₁₀S i.e. (C₃H₃)₂S. Oil

of Garlic. M. w. 114. (140°).

Occurrence.-In the essential oils obtained by distilling, with steam, the leaves, seeds, or bulbs, of various plants (allium sativum, alliaria officinalis, allium cepa, thlaspi arvense). Often associated with allyl-thio-carbimido (q.v.) (Wertheim, A. 51, 289; 55, 297; Pless, A. 58, 36).

Formation.-From allyl iodide and alcoholic

K.S (Hofmann a. Caliours, A. 102, 291).

Properties. - A light oil, smelling of garlic. Combinations .- 1. Forms pps. with salts of Au, Hg, Pd, Pt, and Ag.—(Call.) SPtS2 (W.)-

Au, fig. Fu, Fu, Bita Ag.—(C₃H₃), 58.2 (11.7) (C₃H₃), 5AgNO₃ (Indwig, A. 139, 121). HgS((IgCl₂), 2(C,H₃), S (W.) 2. Combines with MeI (Cahours, Z. 1865, 438); ALLYL SULPHOCIANIDE C₄H₃NS & c. C,H,S.CN. Allyl thio-cyanate.

S.(4. 2 1.071; 15 1.056.

Formation .- 1. From lead salt of allyl mercaptan and cyanogen obloride in ethereal solution (Billeter, B. 8, 464):

 $(C_3H_3S)_2Pb + 2Cl(CN) = 2C_3H_3.S.CN + PbCl_3.$ 2. From ammonium sulphocyanido and a cold alcoholio solution of allyl bromide (Gerlich, A. 178, 85).

Properties .- Changes spontaneously into the isomeric allyl-thio-carbimide, especially when boiled. Alcoholic KOH forms KSCN. Does not alcoholic HgCl₂. Zn and HCl in alcohol form (C₂H₂)₂S and HCN (G.). Sodium amalgam forms Na2S and allyl oarbamine (B.).

ALLYL-SULPHONIC, ACID v. PROPILENE SULPHONIC ACID.

ALLYL SULPHYDRATE v. ALLYL MERCAPTAN. ALLYL-TAURINE C.H., NSO, i.e.

C₃H₂N1I.CH₂.CH₂.CO₃H. [190°-195°]. CILCLCII, SO, H and allymmine at 160° (Jamos, C. J. 17, 369). Prisms (from alcohol). V. e. sol. water.

ALLYL-TH10-CARBAMIC ACID Ethyl other C₆H₁₁NSO i.e. C₃H₅NH.CS.OEt. Allyl-thiourethane (210°-215°). S.G. 14 1.036. From oil of mustard and alcohol at 110°

 $C_3H_5N:CS+HOEt=C_3H_5N1I.CS.OEt$

(Holmann, B. 2, 119). Ppd. by HgCl.Aq.

Allyl-di-thio-carbamic acid C.H.NH.CS.SH. From allyl thio-carbinide and alkalino sulphydrates: $C_3H_5N:CS + HSK = C_3H_5NH.CS.SK$. The free acid is unstable.

Salts.-N11,A': unstable laminæ.-KA': large rhombic plates.—NaA' 3aq: unstable nacreous lamine.—BaA', laq: lauine; v. sol. water.—PbA',: white pp. (Will, A. 52, 30).

ALLYL THIO-CARBIMIDE C,H,NS i.e.

C, 11, N:CS. Oil of mustard, allyl mustard oil, allyl thio-cyanate, allyl iso-thio-cyanate, allyl sulphocyanide, allyl iso-sulpho-cyanide, allyl sulpho-carbinide. M. w. 99, (151°). S.G. 2 1028. S.V. 113·12 (R. Schiff, B. 19, 568). H.F.p. -45,540. H.F.v.-46,700. V.D. 3·54 (for 3·42). Occurrence.-In the oil distilled from the seeds of black mustard (sinapis nigra). Also pre-

sent in oil of garlic, and in horse-radish. Formation. -1. Seeds of black mustard contain potassic myronate, and also an unorganised nitrogenous ferment, myrosin. When treated with water, the ferment splits up the potassic

myronate thus:

 $\tilde{C}_{10}H_{18}NS_2O_{10}K = C_3H_3.NCS + C_6H_{12}O_6 + KHSO$ At low temperatures a little allyl sulphocyanide is also formed (E. Solunidt, B. 10, 187).-2. Allyl sulphoeyanido (q, v) changes, slowly at 15°, quickly on boiling, into allyl thio earbimide Consequently, when allyl iodide is distilled with alcoholic potassio sulphocyanide (Zinin, A. 95, 128; Berthelot a. De Luca, A. Ch. [3] 44, 495), or allyl sulphide (Wertheim, A. 55, 297). the product is allyl thio-carbinide.

Properties .- Oil with pungent odour and burning taste. Blisters the skiu. Sl. sol. water, v. sol. alcohol or other. Slowly decomposed by

water, sulphur being liberated.

Reactions .- 1. Zinc and hydric chlorids reduce it to allylamine and thio formic aldehyde: $C_3H_3NCS + 2H_2 = C_3H_3NH_2 + H_2CS$, the latter being partly reduced to mechane and H.S. (Hofmann, B. 1, 17').—2. HClAq at 200° forms allylamine, CO₂, and H₂S (H.).—3. Alcohol at 100°, or alcoholic potash, forms allyl-thio-carbamic ether (q. v.).—4. Aqueous alkalis, or water and the oxides BaO, PbO, Ag.O, or HgO, form di-allyl urea: $2C_2H_1NCS + 3PbO + H_2O =$ (C₁H₃)₂N₁H₂CO + 2PbS + PbCO₃, -5. K₂S at 100° forms potassio sulphocyanide and allyl-sulphide d NH phide.-6. NH, forms allyl-thio-urea (thio-sinamine).—7. Allehyde-ammonia forms seedles of C₁₄H₃₁N₃S₂O₂ [108°] (R. Schiff B. 9, 571).
—8. Furfuramide in alcoholic solution as t00° forms C, H, N, O, C, H, NOS [118°] (R. Schlif, B. 10, 1191).—9. Bolling conc. KHSO, forms C, H, NH.CS.SO, K. Psarly plates (form alcohol)

(Böhler, A. 154, 59). Combination.—C.H.NCS Ag.SO., Formed by adding AgNO, Aq to aquelus petassic myronate (Will a. Körner, A. 125, 2°7).

Additional References .- Boutron a. Rebiquet, J. Ph. 17, 296; Henry a. Plisson, J. Ph. 17, 451; Dumas a. Pelouze, A. Ch. [2] 53, 181; Aschoff, J. pr. 4, 314; Robiquet a. Bussy, A. Ch. [2] 72, 328; Boutron a. Fremy, J. Ph. 16, 112;
 Löwig a. Weidmann, J. pr. 19, 218; Will, A. 52, 1; Gerhardt, A. Ch. [3] 14, 125; Hubatka, A. 47, 153; Vollrath, J. 1871, 408; Grabowski, A. 138, 173.

ALLYL-THIO-HYDANTOIN Calla Na SO i.e.

HN:C N(C,II,) CO ---CII2

Formation. -(1) By the action of chloroacetic acid on allyl-thio-urea in aqueous solution at 100°. (2) By the action of allyl-cyanamide on thio-glycollic acid.

Minute needles. Sol. hot, sl. sol. cold, water. The hydrochloride (B'HCl) forms glistening prisms. (Andreasch, B. 15, 326; M. 2, 775).

ALLYL-THIO-PARABANIC ACID v. Thio-

PARABANIC ACID.

ALLYL-THIO.URAMIDO.BENZOIC ACID. C₁₁H₁₂N₂SO₂i.e. C₃H₃NH.CS.NH.C₆H₄CO₂H. [1:3] [189° uncor.]. Formed by boiling m-annidobenzoic acid with an alcoholic solution of allylthio-carbimide. Plates. (Aschan, B. 17, 431.)
ALLYL-THIO-UREA. C, H, N, S i.e.

C.H. NH. CS. NH. Thiosinamine. M. w. 116. [74°].

Formation .- From allyl mustard oil and ammonia (Dumas a. Pelouze, A. Ch. [2] 53, 181; Aschoff, J. pr. 4, 314; Löwig a. Weidmann, J. pr. 19, 218; Robiquet a. Bussy, J. pr. 19, 232; Will, A. 52, 1).

Properties .- Prisms, without smell. M. sol. water, v. sol. alcohol, and ether.

Reactions .- 1. HgO or PbO converts it into allyl-eyanamide (q. v.).-2. Warm AgNO, Aq

forms allyl-urea.

Combinations. - (C4H8N2S HCl)2PtCl4. C.H.N.S2HgCl.:curdywhitepp.—C.H.N.SAgNO.,—C.H.N.SBr. [147], six-sided columns; sol. water, and alcohol. Converted by moist Ag.O into alkaline C.H.N.S BrOH, whence HCl forms nto akaine C₄H₃N₂S Broth, whence hot forms C₄H₃N₃SBrCl [130°]. (Maly, Z. 1667, 42).—
(C₄H₃N₃SBr₂PtCl₄.— (C₄H₃N₂SBrCl)₂PtCl₄.—
C₄H₃N₃SCl₄Dr₄.— C₄H₃N₃SI₂ [90°].—
C₅H₃N₃SCl₄.—C₄H₃N₂SCy₂: converted by hot dilute H₂SO₄ into allyl-thio-parabannic acrib (q. v.). (Maly, Z. 1869, 259).—C₄H₃N₂SICyAgCy.
Ethyloiodide C₄H₃N₂SEtl. [72°] (Welzien, 4. 94, 103; M.J.

ALLYL UREA C.H. N.O. i.e. C.H. NH. CO.NH., Allyl carbamide. [85°]. Formation. -1. From allyl cyanate and hot NH. Aq (Hofmann a. Caliours, Tr. 1857, 555).-2. From allylamine sulphate and potassic eyanate (Andreasch, M. 5, 34).-3. From allyl thio-nrea and AgNO, Aq, the liberated HNO3 being neutralised by baryta. The yield is 92 p.c. of the theoretical.

Properties -- Needles. V. c. sol. water, and elcohel, v. sl. sol. CHCl₂, and ether. Br₂ forms di-bromo-propyl-nrea.—Salt.—B'HNO₃.

Di-allyl-uroz (CaHaNH)2CO. Sinapoline.[100°].

Formation. I. By action of water and PbO or baryta on allyl thio-carbimide (Simon, P. 50, 877; Will, A. 52, 25) .- 2. By heating allyl cyanats with water or aqueous potash.

Properties. - Unctuous shining laminm. V. sol. alcohol, ether or hot water; volatile with stsam. Its equeous solution is ppd. by HgCl, and PtCl. Dry HCl liquefies sinapoline forming B'HCl.

ALMONDS.—Bitter almonds contain a glucoside, amygdalin (q. v.), and a nitrogenous unorganised ferment, emulsin. Sweet almonds contain amygdalin but not emulsin. When bitter almonds are ground up with cold water, the amygdalin is split up by emulsin:

C₂₆H₂₇NO₁₁ + 2H₂O = C₇H₆O + CNII + 2C₆H₁₂O₆. The essential oil of bitter almonds is obtained by distilling the preduct with steam. It contains benzoic aldehyde, prussic acid and mandelonitrile, the product of their union. The presence of mandelonitrile is indicated by the formation of phonyl-ethylamino when the oil is acted upon by nascent hydrogen (Fileti, G. 9, 446). Beth sweet and bitter almonds yield by pressure a fixed oil, S.G. 12 913; this consists of clein with some stearin and palmitin. It is called oil of almonds.

ALNEIN. - A golden yellow colouring matter in the alder, birel, and beech (Savigny a. Col-

lineau, C. C. 1881, 703; C. J. 42, 309).

ALOES.—Tho thickened juico of various species of aloc.

ALOETIC ACID C, H, N,O, aq, i.e. C, H, (NO2), O, aq(?). Tetra-nitro-anthraquinone(?) Obtained, together with ehrysammic acid, by warming aloes with HNO₃ (Schunck, A. 39, 24; 65, 235; G. J. Mulder, A. 72, 286; Finck, A. 134, 236). Yellow amorphous powder. Sl. sel. cold water, m. sol. het water or alcohol, forming purple solutions, which become yellow on addition of acids, and red again when neutralised. It has a bitter taste. Boiling HNO₃ converts it into chrysammic acid, and ultimately into pierie acid. Warm potassium or ammonium sulphide containing excess of alkali forms an indigo-bluo gelatinous mass. Salts: BaC_{1,}H₂N₄O₁₀. Ag₂A": inseluble, dark rsd pewder.

ALOIN .- The purgative principle in aloss. There are several varieties classified by Shenstone (Ph. [3] 13, 461) as follows: (1) Nataloins .- Are not reddened by IINO, but converted by it into pieric and oxalic acids.—(2) Barbaloins-are reddened by IINO, aloetic, chrysam. mic, pieric and exalic acids being formed. These may be subdivided into (a)-barbaloins, reddened by cold HNO₃ (S.G. 1·4), (β)-barbaloins, reddened by fuming HNO₃, not by celd HNO₄ (S.G. 1·4). The aloins may be extracted by het water of het spirit from the various alocs, and purified by re-crystallisation. They dissolve in caustic and carbonated alkalis, forming orangs solu-Their solutions are ppd. by lead tions. subacctate.

(a)-Barbaloin, CieHinO, and (B) - Barbaloin, Zanzaloin or Socaloin C16H18O, occur in aloss from Barbadoes, Socotrina, Zanzibar and Jafferabad (T. a. H. Smith, Chem. Gaz. 1851, 107; Stenhouse, P.M. [3] 37, 481; Tilden, C. J. 25 489; 28, 1270).

Reactions.-1. By distilling with sine due a very little methyl-anthracene may be got (Schmidt, B. S. 1275).—2. Potash-fusion gives ordin, p. oxybenzoic acid, and aloreinic acid.—3. Boiling dilute sulphuric acid forms p-ecumaric acid.—4. HOl and KClO₂ form tri-chloro-aloin, . C₁₆H₁₅Cl₂O₂, yellow prisms (from alcohol).—5. Browine forms tri-bromo-aloin; yellow needles (from alcohol).—6. Ac₂O forms tri-acetyl-aldin, Cl₁₆H₁₅Ac₂O₂: amorphous.

Nataloin C_{1s}H_{1s}O₁(?). Occurs in Cape aloes (Flückiger, Ar. Ph. [2] 149, 11; Bl. 17, 328; Tilden, C. J. 25, 153). Thin bright yellow scales; sl. sol. water, benzene, ether, CS₂, and CHCl₂. Its solution in H₂SO₄ is turned green by KNO₂, the colour changing through red to blue.

Additional References.—T. B. Groves, Ph. 16, 128; Orlowski, Fr. 5, 309; Hlasiwetz, A. 134, 287; Rembold, A. 138, 186; Bornträger, Fr. 20, 234; R. H. Groves, Ph. [3] 11, 1045; Lenz, Fr. 21, 220; Plenge, Ph. [3] 15, 330; Cripps a.

Dymond, Ph. [3] 15, 633).

ALORGIC ACID C,H₁₀O₃ aq. [97°]. C,JI₁₀O₃ [115°] i.e. C₀H₂(OH)Mc₂CO₂H. Formed in small quantity, along with p-oxy-benzoic acid and crein, by fusing aloes with KOH (Weeslsky, A. 167, 65). Long needles. Sl. sol. cold water, v. sol. hot water, alcohol, and ether. The aqueous solution is not coloured by Fc₂Cl₂Aq, but alkaline solutions are turned cherry-red by air. Hypochlorites turn the aqueous solution of the acid purple-red, colour destroyed by excess. Basic, but not neutral, lead acetate gives a pp. Alorcic acid reduces silver vitrate and Fchiing's solution. Fused with potash it forms orein and K₂CO₃.

Salts. BaA', 6aq: small needles.—CaA', needles.—CuA', 4aq.

Acctyl derivative C₉H₉AcO₃: [125°]; needles.

Anhydride C₉H₈O₂: [138°]; formed by distilling the acid.

ALOXANTHIN C₁₂H₁₀O₆ i.e. C₁₄H₂Me(OH)₄O₂. From barbaloin and socaloin, but not from nataloin, by chromic mixture. Reduced, by distillation with zinc dust, to methyl-anthracene. Its alkaline solutions are cherry-red. Nitric acid converts it into aloetic and chrysammic acids (Tilden, Ph. [3] 8, 231; C. J. 32, 903). It forms an acetyl derivative C₁₂H₂AcO₂.

ALPHA. To find compounds beginning with this prefix, remove the prefix and look for the remaining word.

ALPININ C₁, H₁₂O₆. {174°}. Light yellow needles (+H₂O). Occurs, together with camphoride and galangin in the galanga-root (Jahns, B. 14, 2810).

ALSTONIDINE. [181°]. An alkaloid occurring along with alstonine (q, v) and porphyrine (q, v) in the bark of Alston&constricta. It may be separated from porphyrine by its more sparing solubility in petroleum. It crystallises in radiating needles, sol. alcohol, chloroform, and ether. Its solutions display intense blue fluorescenec. It is not coloured by cone. H₂SO₄ or HNO₃. Its salts crystallise in colourless needles. The gold and platinum salts are golden flocculent pps. (Hosse, 4. 206, 368).

ALSTONINE C₂₁H₂₆N₂O, 33aq. Chlorogenine (F. v. Müßer a. L. Rummel, C. J. 35, 34; Oberlin a. Schlagdenhauffen, Ph. [3] 10, 1059; O. Hesse, A. 205, 360). An alkaloid in the bark of Altionic constricts (F. v. M.), from which it may

be extracted by alcohol. The extract is evaporated, treated with very dilute HCi, filtered, ppd. by NH₂, dissolved in ether and evaporated.

Properties.—Orange-yellew, brittle, pellucid, bitter, mass. It melts & 100°. When dry, it melts at 195°. It dissolves casily in alcohol, ether, chloroform, and dilute acids, but sparingly in water. All dilute solutions show blue fluorescence. Salts. 1°d. by excess of acid.—(B'HCl).PtCl, 4aq.—(B'HCl).HgCl..—B',H.Cr.O..

(B'HCl) PtCl, 4aq.—(B'HCl) HgCl, —B', H, Cr, O, ALUMINA.—Oxide of Aluminium, Al, O, v.

ALUMINATES.— Certain minerals are known, e.g. Al₂O₃MgO, Al. O₃BeO, &c., which may be regarded as derived from the hydrate Al₂O₃H₂O (c. ALUMINIUM, HYDROXIDES OF), by replacing H₂ by Mg, Be, &c. Some of these minerals have been prepared by Ebelmen (A. Ch. [3] 22, 211) by dissolving Al₂O₃ and the other metallic oxide in molten boric acid, and removing part of the solvent by long-continued heating; in this way he prepared spunelle Al₂O₄Mg, chrysoberyl Al₂O₄Be, &c. By heating Al₂F₈ with boric acid and ZnF₂, Deville and Caron obtained galmite Al₂O₄Zn.

Barium aluminate Al.O.Ba.4H.O was obtained in crystals by fusing Al.O. and carbon with BaO, BaCO, Ba2NO, or BaSO, dissolving in water and crystallising (Deville, C. R. 54, 327; v. also Gaudin, C. R. 54, 687; also Beckmann, J. pr. [2] 26, 388 a. 474). Compounds of Al.O. and BaO obtained by action of BaOAq on Al.O. and Al.Cl.Aq are described by Beckmann (B. 14, 2151).

Polassium aluminate Al₂O₄K₂3H₄O was ebtained in crystals by Fremy (A. Ch. [3] 12, 362) by fusing Al₂O₂ and KOH in a silver dish, dissolving in water, and evaporating in vacuo; this salt may be recrystallised from con. aqueous solutions, from dilute solutions Al₂O₄H₄ separates out

Sodium aluminate has not been obtained crystallised. An impure salt is used in manufactures; it is obtained by heating cryolite and CaO, or bauxite and NaOH, in steam, dissolving in water and evaporating to dryness (Morin, J. 1862, 668). Tissier (C. R. 48, 627) described four compounds to which he assigned the compositions Al₂O₆Na₆, Al₂O₈Na₇, Al₂O₈Na₈, and Al₂O₄Na₂; but no analyses are given in the original paper.

M. M. P. M.

ALUMINIUM.—Al. At w. 27-02 (Mallet, T. 171, 1003). Mol. w. unknown as V.D. has not been determined. [About 700°]. S.G. \(\frac{4}{2} \) (fused) 25-83 (Mallet, T. 171, 1003); (after pressure of 6,000 atmos.) 25-62 (Spring, A. Ch. [5] 22, 170). S.H. 218 (Longuinine, A. Ch. [5] 27, 398); (0°-100°) 225-3 (Mallet, Lc.). C.E. (lin. at 40°) 002-313; (lin. 50°) 002-336 (Excan. C. R. 68, 1125) T.C. (Ag = 100) 31-33 (Lorenz, II. 13, 422). F.C. (Hg at 0°=1) 20-97 at 0°, 16-15 at 15° (Horenz, Lc.). S.V.S. 10-4. Clief lines in spectrum, 39-60°3, 3943-4, 3612-4, 3001-9, 6081-2, 2815-3, 2630-6 (Hartley, T. 175, 101).

Occurrence.—The metal aluminium does not occur native; but as silicate (in all clays and in very many minerals, especially the felspars), and oxide (corundum, diaspore, &c.), and fluoride (crye lite), it is very widely and largely distributed, forming nearly $\frac{1}{12}$ of the earth's crust. Alumina was shown to be a distinct earth by Marggraff in 1754; the metal was separated by Wöller is

1328 (P. 11, 146), and in purer form in 1854 (A. 53, 422).

Preparation. -- Wohler decomposed Al, Cl. by F.; in 1854 Deville employed Na and decomposed 2NaCl.Al,Cl_s (J. pr. 61.93, 113, 219, 386). Bunsen (P. 92, 648) decomposed fused 2NaCl.Al,Cl_s by an electric current. Rose (P. 96, 152) decomposed 6NaF.Al.F (cryolite) by fusing it with KCl Basset (J. 1869. 753) reduced and Na. 2NaCl.Al.Cl. by Zn, and heated the Zn-Al alloy to white heat to remove the Zn. The Al,Cl, may also be reduced by KCN (Wagn, J. 1858. 1); a compound of Al and S may be reduced by Fe, or by hydrocarbons (Petitjean, J. 1858, 136); and in other ways. The usual method of preparation is to heat 2NaCl.Al.Cl, with about 36 p.c. Na and 40 p.c. eryolite (as a flux), on the hearth of a roverberatory furnace, and to run off the molten Al into iron moulds. The 2NaCl.Al,Cl, is prepared by heating banxite (silicate of Al containing Fe) with Na,CO,, whereby Na aluminate is formed, dissolving in water, ppg. Al₂O₃ by a stream of CO2, collecting, washing, and drying the Al.O3, mixing it with charcoal and NaCl and heating strongly in Cl whereby 2NaCl.Al.Cl, is formed and distilled off into receivers. Mallet (T. 171, 1003), prepared very pure Al by fusing Al Br. with KCl and NaCl in the ratio 2(KCl.NaCl):Al_Br, and then heating with Na in clay crucibles lined with Al2O3 and Na aluminate; the reduced metal was heated on a support of Al.O., washed with HClAq, then with water, and dried at a gentle

Properties.—A tin-white metal; grey when in powder. Odourless and tasteless. After fusion about as hard as silver; hammered metal is about as hard as soft iron. Very malleable, and ductile; very sonorous; may be highly polished. Tenacity nearly equal to that of Cn (V. Burg, D.P.J. 151, 280); less than that of Zn (Karmarsch, D.P.J. 152, 441; 172, 55). Very feebly magnetic. Melts fairly easily (about 700°) and crystallises, apparently in regular octahedra, on cooling. Non-volatile, and non-oxidisable in air; heated in oxygen becomes covered with film of Al₂O₂. Unacted on by H₂S or ammonium sulphides, and by S only at high temperatures. Searcely attacked by HNO₄Aq, but dissolved by HIOlAq, H₂SO₂Aq, KOHAq, and NaOHAq. Most specimens of Al contain Fo and Si; they are more easily oxidised than the pure metal.

Aluminium forms one series of salts the simplest formulæ of which represent them as derived from acids by replacement of H, by Al;

e.g. Al₂3SO₄, Al3NO₃ &c.

The atom of Al is trivalent in the gaseous molecule AlCl₂. Many experiments have been conducted to determine the mol. w. of Al chloride; Nilson a. Pettersson have finally shown (Z, P. C. 4, 206) that the only molecules which exist throughout a considerable range of temperature have the composition AlCl₂. Odling (P. M. [4] 29, 316) stated the V.D. of Al methide to be 36·2 at temperatures above 200° (and 72·4 at 130°); and hence mol. w. Al(Cll₂)₃. It remained, however, uncertain whether the gas was homogeneous or not (e. Wanklyn, P. M. [4] 29, 313; Williamson, ibid. 35·5; Le Boux a. Lonïse, C. R. 106, 73). Quincke (Z, P. C. 3, 164) has shown that the molecular formula of Al methide is Al(Cll₂)₃.

The atomic weight of Al has been determined; (i.) by analyses and determination of V.D. of Al₂Cl₃, Al₂Br₄, and Al₂I₅; (ii.) by measuremente of S.H. of Al; (iii.) by comparison of various compounds of Cr and Fe; (iv.) by and yses of ammonia-alum and Al₂Br₆, and by measuring the H evolved by the action of Al on NaOHAq (Mallet, T. 171, 1003 i.; v. also for older determinations Berzelius, P. 8, 187; Dumas, A.Ch. [3] 55, 151; Tissier, C.R. 46, 1105; Terreill, Bl. 31, 153).

Aluminium is a distinctly metallic element; no allotropic forms of it are known. According to the investigations of Wheatstone (J. 1855. 22), in KOHAq Al is electropositive to Cd, Sn, Pb, Fe, Cu, and Pt, and negative to Zn; in HClAq Al is positive to Sn and Pb, and negative to Zn and Cd. Al decomposes H_Q0 at 100°. It reacts with acids to form definite salts, but at the same time hydrated Al_Q0₄ dissolves in alkalis to form unstable salts in which the Al forms a part of the negative, or acid, radicle (v. Aluminames).

The thermal value of the action of acids on Al₂O₂,3H₂O is a large positive number approximately equal to the value for CdO₂H₂, and FeO₂H₂, although considerably less than the values of the corresponding actions with CaO₂H₂, SrO₂H₂, and BaO₂H₄ thus (Thomsen):—

The difference between the heat of formation of a metallic chlorido and hydroxide has usually a positive value; in the case of a nonmetallic chloride and oxide the difference is usually a negative quantity (Thomsen, Th. 3, 531); in the case of Al the difference in question has a large negative value, thus [Al², Cl²]—[Al², O³,3H O] = -22, 320. Al shows several analogies with Be; they both very readily form basic and double sults; Al_O, ie less basic than BeO; both metals readily alloy with Si; neither seems to be easily acted on by S. For fuller discussion of the chemical relations of Al v. Earrus, Metals of the

Reactions .- 1. Pure Al is unacted on by ordinary air; impure specimens of the metal become covered with a film of oxide. - 2. Water is decomposed by Al at 100° with evolution of H.-3. Con. or dilute nitric acid has no action on Al.-4. The metal is easily dissolved by hydrochloric ocid. 5. Dilute sulphuric acid evolves II, forming Al, 3SO,; hot con. H, SO, evolves SO, -6. Most carbon acids, e.g. acetic, tartaric, have little or no action on Al; but in presence of NaClAq the action becomes marked, Al, Cl, boing formed .- 7. Sulphuretted hydrogen has no action oven at high temperatures .-Aqueous potash or soda dissolves Al, evolving H, and forming an aluminate (v. Cavazzi, G. 15, 202); molten KOH or NaOH does not act on Al.-9. Sulphates, carbonates, borotes, and silicates, of the alkali metals are decomposed by Al at high temperatures. -10. Potassium nitrate oxidises Al when the two react at a white heat. 11. Alkali sulphides are without action even at high temperatures; silver sulphide when

heated with Al is partly reduced with formation of a Ag-Al alloy.-12. Many metallic oxides are deoxidised by Al at high tomperatures, e.g. oxides of Pb. Cu, Fe; oxides of Zn and Mn are not acted on .- 13. Alkaline, but not neutral or slightly acid, solutions of lead, silver, and tin, are reduced by Al with ppn. of the metals; Cu is ppd. from CuSO Aq. --14. Most metallic chlorides in solution are reduced by Al (KClAq, NaClAq are exceptions; Cossa, Z. [2] 6, 380, 443). Fused zinc chloride, but not MgCl₂, is reduced by Al (Flavitsky, B. 6, 195).

Combinations.—1. With oxygen to form Al₂O₃; only at high temperatures, and then superficially.-2. With sulphur to form an unstable compound (v. Aluminium, Sulphide or) only at very high temperatures.—3. Wöhler (P. 11, 160) states that Al combines with phosphorus, selenion, tellurium, and arsenic, when heated in the vapours of these elements; but little is known of the compounds .- 4. With boron Al forms at least two compounds (v. ALUMINIUM, BORIDES OF) .- 5. With chlorine, bromine, and iodine, Al combines to form Al₂Cl₈, Al₂Br₆, and Al₄I₆ (q. v.). -6. Al forms alloys with most of the metals; these alloys are usually formed by heating the metals together; or sometimes by heating Al₂O₃ and carbon with the other metal. The properties of many metals are considerably changed by alloying with small quantities of Al. The alloys of Al with Cn, Ag, and Sn are much used because of their colour, hardness, and stability, and the ease with which they are worked. The alloys of Al have been chiefly studied by Calvert a. Johnson (P. M. [4] 10, 240); Tissier (C. R. 43, 885; 49, 54); Debray (C. R. 44, 925); Wöhler (A. 106, 118; 113, 248; 138, 253; P. 11, 161); Michel (A. 115, 102). The alloys with copper containing from 6 to 10 per cent. Al aro yellow, hard, unacted on by moist air, water, or salt solutions, and are easily worked. The alloys with silver are also very stable, have a fine colour and may be highly polished. When a little Al is alloyed with tin the products are very hard and elastic. (These alloys will be more fully described under Copper, Silven, and Tin.) Alloys of Al with the following metals have been prepared:—Bi (a very little Bi makes Al extremely brittle); Ca (by heating Al, Na, and much CaCl., Wöhler); Cu, Cr (by heating Al with Cr. Cl. KCl, Wöhler); Au, Fe, Mg, Mn (by heating MnCl., KCl, NaCl, and Al); Hg (Cailletet, C. R. 44, 1250); Mo, Ni, Pt, Ag, Na (this alloy easily decomposes H.O); Sn, Ti, W (by heating WO₃ with Al, cryolite, and KCl, and NaCl); and Zn.

Detection. - Many Al compounds are soluble in water; most are dissolved by HClAq. Strongly heated Al2O2 is nearly insoluble in acids; it, and also the insoluble Al-containing minerals, may be dissolved by fusion with KHSO, and treatment with water (H. Rose, P. 1, 275).

Estimation.—1. Usually as Al.O.: a fairly con. solution is ppd. by a small excess of NH, Aq (if Mg salts are present a good deal of NH, ClAq is added), the free NH, is removed by warming, the pp. is washed, thoroughly dried, and strongly licated for some time. - 2. Al.O.H. may also bo ppd. by Na,S,O,Aq. This method is specially applicable in presence of Fo salts;

is dissolved in HClAq, the solution is nearly neutralised by Na.CO, Aq and diluted so that 50 o.o. do not contain more than 1 gram Al,O, a slight excess of Na, S,O, Aq is added, and after a little the liquid is boiled so long as SO, comes off, the pp. of Al,O.H. vaixed with S is filtered hot, washed with hot water, and strongly heated until all S is burnt off;

 $Fe_2Cl_0Aq + 2Na_2S_1O_3Aq =$ $\text{Fe}_{v}\text{Cl}_{1}\text{Aq} + 2\text{NaClAq} + \hat{\text{Na}}_{v}\text{S}_{1}\text{O}_{s}\text{Aq}$ (v. Chancel, C. R. 46, 987).—3. Al is also sometimes estimated as a basic acetate (v. Atkinson, Fr. 3, 329; also Schulze, J. pr. 47, 313); or as basic formate (Schulze, C. C. 1861. 3). For methods of separating Al from alkaline earths v. Deville, A. Ch. [3] 38, 5; from Co, Ni, and Zn v. Haidlen a. Fresenius, A. 43, 129; from iron v. Macivor, C. N. 29, 199; from iron and phosphoric acid v. Flight, C. J. [2] 13, 592; also Esilman, C. N. 28, 208; from chromium v. Dexter, P. 89,142.

Aluminium, Alloys of, v. Aluminium; Combinations, No. 6.

Aluminium, Arsenids of. According to Wöhler (P. 11, 160) Al combines with As when the elements are heated together; the product is a grey metal-like mass, decomposed by water with cvolution of Asll,. No analyses are given.

Aluminium, Borides of. Al seems to form two definite compounds with boron, AlB, and They may be obtained by packing a rod of Al in amorphous B in a carbon crucible, placing this in a Hessian crucible with powdered charcoal between the crucibles, covering, and heating to redness for $1\frac{1}{4}$ or 2 hours; B_2O_3 may be used in place of B (10 parts B_2O_3 to 8 parts Al). On cooling, the fused mass is treated with HClAq, whereby Al dissolves and crystals of the borides remain, which may be separated by hand (Hampe, A. 183, 75). Both borides were obtained by Wöhler and Devillo and regarded by them as crystallised boron (A. 101, 113, and 347; 141, 268); the compositions represented by the formulæ AlB, and AtB, are assigned by Hampe (l. c.). AlB2 crystallises in thin, lustrous, palo copper-coloured, six sided, plates; unchanged by heating in air; slowly dissolved by hot con. HClAq and NaOHAq, easily soluble in warm 11NO, Aq. This compound may be prepared by the action of BCl, vapour on hot A1; or by heating BF₃.KF with KCl, NaCl, and Al. AlB₁₂ forms black, monoclinic crystals, transparent in very thin plates; harder than corundum, softer than diamond S.G. = 2.534. These crystals are unacted on by con. 11ClAq or KOHAq, and very slowly by hot con. H.SO,Aq; they are soluble in hot con. 11NO₃Aq. They are oxidised by molten KOH and PbCrO₄; also by molten KHSO₄; but are not acted on by molten KNO3. Heated with Pt an easily fusible alloy is formed.

Alnminium, Borocarbide of. AlaC.B. Originally obtained by Wöhler and Deville, and supposed to be crystallised boron; examined more fully by Hampe (A. 183, 90). Prepared by long-continued heating at a very high temperature of B.O. and Al in a graphite orucible (for details, v. Hampe, l.c.). The compound crystallises in yellowish, sparkling, crystals of the dimetric system. S G. = 2.615; hardness between the two metals are ppd. as hydrates, the pp. that of diamond and corundum. The orystals

are slowly dissolved by hot con. HNO.Aq, but not by HClAq, H,SO,Aq, or KOHAq; they behave towards molten KOH, PbCrO,, and

Aluminium, Brombe of. AlBr₂ or Al, Br₃; not certain. [abt. 90 A(Weber, P. 103, 251). (263°3) at 747 mm. (Mallet, T. 1880, 1003). S.G. 2:54. V.D. 266'8 (Deville a. Troost, A. Ch. [8] 58, 257). H.F. [Al2Br6] = 239, 440 (Th. 3, 210).

Formation.—By passing Br vapour over a heated mixture of alumina and charcoal.

Preparation.—By the action of Br on excess of powdered Al, and subsequent repeated distillation from Al, and finally in atmosphere of N.

Properties.—White, lustrous, plates; fumes in air; melts to a mobile liquid; soluble in CS., alcohol, and water, in latter with production of much heat [AlBre, Aq] = 170, 600 (Th. 3, 240).

Reactions and Combinations.—Aqueous so-

lution on evaporation in vacuo yields crystals of Al₂Br₆, 12H₂O; this solution is decomposed on heating into Al₂O₆H_s and HBr. Al₂Br_e is decomposed into Al₂O₆ and HBr when heated in air; it combines with the broundes of the alkali metals to form double salts, e.g. 2KBr.Al_Br. (Weber, P. 103, 259); it absorbs NH3, also ILS, forming compounds which are decomposed by heat (Weber, l.c.).

Aluminium, Chloride of. AlCl., Mol. w.

133·13 (800°-1500°) (Nilson a. Pettersson, Z. P. C. 4, 206). V.D. 66.5 (N. a. P. l.c.). H.F. [Al,Cl*]

= 160, 980 (Th. 3, 210).

Formation.—1. By heating powdered Al to redness in dry Cl. - 2. By passing HCl mixed with CS2 vapour over crude alumina or clay heated to redness; Al₂S₂ is formed but is at once decomposed by the HCl (Curic, C. N. 28, 307).—3. By heating Al with various metallic ehlorides, e.g. ZnCl. (Flavitsky, B. 6, 195).—1. By heating Al.O. with NH.C. (Rose, P. 74, 569), or with PCl., BCl., or SiCl. (Troost a. Hautefenille, C.R. 75, 1710 and 1819).

Preparation.-100 parts Al₂O₃ are made into a thick paste with 40 parts carbon by the help of starch paste, or oil; the paste is kneaded to a oylinder, which is dried, heated in a covered crucible, and removed while hot to a porcelain tube connected with a Cl apparatus; as soon as the apparatus is full of Cl, the cylinder is heated to redness; Al₂Cl₈ distils over into a receiver. It is purified by sublimation from Al in a closed

tube bent to an obtuse angle.

Properties. Transparent, deliquescent, hexagonal plates; colourless when pure, but usually yellowish because of presence of chlorides of Fe, 8, &c. Fusible in large masses but volatilises without fusion if heated in small quantities. Soluble in water with production of much heat $[Al^2Cl^2,Aq] = 153,690 \ (Th. 3, 240);$ soluble in alcohol, and ether; insoluble in beuzene; unacted

on byeHClAq.

Reactions.—1. Fumes in moist air absorbing H2O and giving off IICi .- 2. Easily decomposed (to Al₂O₂ and HCl) by steam (Kulinheim, J. 1861. 149). -3. Partly decomposed by oxygen at a red heat with evolution of Cl (Berthelot, C. R. 86, 787).—4. Heated to redness with lime, corundum (Al₂O₃) is formed; with magnesia, spinello (Al₂O₄Mg) is produced (Daubrec, C. R. 39, 135).— 5. Decomposed below red heat by potassium or of Al₂O, in HFAq to an excess of KFAq.Al₂F.

sodium, with production of Al, and KOL or NaCL -6. Molten Al₂Ol₄ is electrolysed to Al and Cl (Buff, A. 110, 257).-7. Sulphuric anhydride forms Al₂SSO₄, SO₂, and Cl.-8. Solution in water is decomposed by heat into Al2O2 and HCl.

Combinations. - 1. Dissolves in water; solution when slowly ovaporated gives deliquescent crystals of Al₂Cl₄.12H₂O (Bonsdorff, P. 27, 279). -2. Absorbs dry ammonia to form yellow Al₂Cl_a6NH₃; when this is heated in a stream of dry H, Al₂Cl₆2NH₃ is obtained (Persoz, A.Ch. 44, 319). -3. Combines with phosphoretted hydrogen to form a yellow powder 3Al Cla 2PH as decomposed by H₂O giving off PH₃ (Rose, P. 24, 295).—4. Phosphoric chloride at 180° forms Al₂Cl₆·2PCl₅; crystalline, easily fusible, easily decomposed by H₂O (Weber, P. 107, 375).—5. Combines with some other non-metallic chlorides; chief combinations are Al₂Cl₂.2PoCl₂. (Casselmann, A. 98, 220); Al₂Cl₂.2NaCl (Weber, P. 118, 471); Al₂Cl₃.SeCl₃, and Al₃Cl₄.TeCl₄ (Weber, P. 104, 421). Forms a compound with II.S which is decomposed by II.O (Wöhler, P. 11, 160).—6. Combines with chlorides of alkali metals to form Al.Cl, 2MCl (Degen, A. 18, 332; Deville, A. Ch. [3] 43, 30); the most important is Al_Clo.2NaCl, white erystals, melting at about 185°, volatile at a red heat without decomposition; deliquescent, but much less so than Al2Cls. This salt is prepared on the large scale, as from it the metal Al is obtained (v. Aluminium, Preparation of) .-7. By evaporating solutions of the mixed chlorides the compounds Al2Cls.2PdCl2.20H2O and Al₂Cl₆.2PtCl₄.30H₂O were obtained (Welkow, B. 7, 304 and 802).-8. With sulphurous anhydride forms Al₂Cl₆.2SO₂, a heavy liquid, solidifying at -10° (Andrianowsky, Bl. [2] 31, 495).
Aluminium, Fluoride of. AlF₃. Mol. w. un-

certain, as V.D. not determined. S.G. 3.1.

Occurrence.—In combination with NaF aa cryolite Al,F. 6NaF; also with silicates in topaz, &c.

Formation .- 1. By action of HF on Al₂O₃. -2. By fusing cryolite with Al, 3SO,, and digesting in water whereby Na SO, is dissolved and Al F, remains (Deville, C.R. 42, 49).

Preparation.-A mixture of fluorspar and alumina, in a boat of graphite, is heated to fusion in a tube of graphite, in dry HCl; crystals of Al.F. sublimo and CaCl, remains (Deville, C.R.

Properties and Reactions .- White, and very obtuse rhombohedra; unacted on by air, by acids, or by aqueous alkalis (Deville, C. R. 43,

970). Volatile at bright red heat.

Combinations.-1. With hydrofluoric acid and water: by dissolving Al₂O₃ in 1LSiF₆Aq, digesting with Al,O, till SiO, is ppd. and solution is neutral, and evaporating, crystals of AlaF .7H2O are obtained; if digestion with Al.O. is stopped while liquid is distinctly acid but SiO₂ is supper while liquid is distinctly acid but SiO₂ is ppd., and alcohol is added, 3Al_F₆AHF.10H₂O is formed; if, instead of adding alcohol, the acid liquid is evaporated Al₂F₆HF.5H₂O is produced (Deville, A. Ch. [3] 61, 327).—2. With alkali fluorides compounds of the form Al, F. 6MF are formed, the most important of these are the K and the Na compounds. Aleka cKF is the K and the Na compounds. obtained as a gelatinous pp. by adding a solution

4KF is produced by adding KFAq to excess of Al.F., in HFAq; if the pp. is boiled in the liquid Al.F., 6KF is formed. Both pps. form white powders when dry, and are decomposed with loss of all their F by hoating with con. H.SO.Aq. Al.F. 6NaF occurs native as cryolite on the west coast of Greenland. S.G. = 2.96. It may be prepared by the action of HFAq on Al₂O₃ and Na₂CO₃, mixed in ratio Al₂O₃:3Na₂CO₃, drying, and heating to fusion. Forms colourless, transparent, dimetric crystals; softer than folspar; melts below red hoat, forming a colourless glass on cooling. Decomposed to CaF, and a solution of Al₂O₂ in NaOHAq, by boiling with milk of lime, or fusing with CaCO₂ and boiling in H₂O. Heated with H.SO.Aq loses HF, and produces Al. 380, and Na. 80.

Aluminium, Hydrated Oxide of, v. ALUMINIUM, HYDROXIDES OF.

Alnminium, Hydroxides of. Three compounds of Al, H, and O are known; they react rather as hydrated oxides, Al₂O₃.xII₂O where x=1, 2, and 3, than as hydroxides. Al₂O₃ which has not been strongly heated takes up H2O; but the definite compounds are obtained by indirect methods; Al₂O₃ which has been strongly heated has no action on H.O. The hydrates of Al₂O₃ lose their water at a red heat. When freshly ppd. they dissolve easily in acids, forming the Same salts Al₂X₄ where X=acid radicle (v. ALUMINIUM, SALTS or). Thomsen gives the following thermal values [Al2, O3, 3IFO] = 388,920; [Al, O³, H³] = 297,000; $\left[\frac{A^{12}O^{6}H^{6}}{3}, H^{2}SO^{4}Aq\right]$ =

 $\begin{bmatrix} Al^2O^{\bullet}\Pi^a \\ 3 \end{bmatrix}$, 2HClAq = 18,640 (Th. 3, 20,990;

240). The hydrates also dissolve in KOHAq and NaOHAq with formation of easily decomposed aluminates (q. v.). The hydrates, especially Al₂O₃.3H₂O, form a class of bodies called lakes by their action on vegetable colouring matters; they pp. these colouring matters from solutions, hence these hydrates are used as mordants.

I. MONOHYDRATE. Al₂O₂, H₂O (= Al₂O₄H₂). Occurs nativo as diaspore, in transparent, trimetrio crystals, S.G. 3-43, which crumble to powder when heated but lose all H2O only at about 360°; insoluble in water and in boiling HClAq. By heating amorphous Al₂O₃ in a closed tube with H₂O to 240°-250° Mitscherlich obtained hydrated Al₂O₄ nearly agreeing in composition with formula Al₂O₄.H₂O (J. pr. 83, 468). According to Becquerel (J. 1868, 87; J. 1874, 132) crystalline ALO, HO is produced when a tube containing Or, Cl, Aq and covered with parchment paper is suspended in a solution of ALO, in KOHAq.

II. DIHYDBATE. M.O, 2H_O (= ALO, H).

Occurs native as amorphous bauxite (Berthier, Occurs native as anarypious occurred permits, S. 35, 154). Prepared by the action of NH₄ClAq on Al₂O₄ in KOHAq, washing, and drying at 100° (Löwe, Z. 3, 247); also by ppg. hot solutions of Al salts by NH₂Aq, digesting the pp. in the warm liquid, washing, and drying at 100° (St. Gilles, A. Ch. [3] 46, 58). A dihydrate is also obtained by the slow decomposition of basic aluminium acetate, and subsequent ovaporation to drynsss at 100° (v.i.). The dihydrate is said to be nearly insoluble in acids and aqueous alkalis. Two varieties of Al₂O₂.2H₂O in aqueous solutions were obtained by Crum (C. J. 6, 225). Vol. I

and Graham (T. 1861. 183), respectively. Crum prepared basic Al acetate

(2Al2(C2H2O2)4.Al2O4H6.3H2O), dissolved this by heating with much water, heated this solution to \$100 in a closed tube for ten days, whereby the salt was decomposed into acetic acid and Al₂O₃.xII₂O, then diluted, and boiled in an open vessel until all acctic acid was removed. The solution of Al₂O₃.xH₂O thus obtained was colourless, tasteless, and neutral to litmus; it was easily coagulated by small quantities of sulphuric, citric, tartaric, and many other acids, and by various salts, not by acctic, formic, boric, and one or two other acids. solution does not act as a mordant; evaporated at 100° Al₂O₅.2H₂O insoluble in acids is oobtined; the coagulated hydrato dissolves in con. acids. Graham dissolved Al, O, xH, O in Al, Cl, Aq or in Al acetate solution, and dialysed; the aqueous solution of Al,O.xII,O thus obtained was very easily coagulated by acids, alkalis, and salts; it acted as a mordant; the coagulated hydrate dissolved easily in dilute acids.

III. TRIBYDRATE. Al.O. 3H2O (= Al2O.H.). Occurs native, in hexagonal, fibrous, crystals, as gibbsite, and hydrargyllite. Prepared by ppg. cold solution of Al salts by NH Aq, or (NH₁)₂CO₃Aq, washing, dissolving in HClAq, reppg., washing, and drying at 100°. A soft, friable, white powder; casily soluble in acids and fixed aqueous alkalis; insoluble in water. When freshly ppd. it forms a gelatinous mass, soluble to some extent in NII, Aq, but reppd. on standing in air. Heated to redness it loses all its II.O, and contracts considerably. A erystalline trihydrate is formed by the action of air on a saturated solution of Al₂O₃ in KOHAq (Bonsdorff, P. 27, 275); it is insoluble in cold acids, slowly dissolved by boiling HClAq, more rapidly by II, SO, Aq. Al, O, 3H, O is also said to be formed by the action of H₂O on Al amalgam (Cossa, Z. 13, 443).

Alnmininm, Iodide of. All, or Al, i, not certain. [abt. 185] (abt. 360°) (Weber, P. 101, 465; 103, 259). S.G. 2·63. V.D. 387·4 (Devilla a. Troost, A. Ch. [3] 58, 257). H.F. [Al', F] = 140. 780 (Th. 3, 240).

Preparation.—Al in small pieces is placed in a retort, about \(\frac{1}{2} \) to 1 p.o. of the calculated quantity of I is added, CO2 is led in, and the retort is heated until combination occurs; rather loss than the quantity of I needed to convert. all the Al into Al, Is is then added, and heating, in a steam of CO, is continued until Al, I, sublimes (Gustavson, A. 172, 173).

Properties, &c .- White, deliquescent, plates; soluble in alcohol, and CS_2 ; soluble in water with production of much heat $[Al_2^{II},Aq]=178,000$ (Th. 3, 240). Solution in II_2O on standing over H₂SO, gives Al₂I₈.12H₂O. Decomposed by heat, ing in O (Schulze, J. pr. [2] 21, 40). Forms double salts with alkali iodides; absorbs NH,; apparently does not combine with H2S (Weber, P. 101, 465; 103, 259).

Alnminium, Nitride cf. Al heated in N increases in weight, and heated with molten NaOH evolves NH, (Briegleb a. Geuther, A. 123, 238). The compound Al, N, was obtained by Mallet (C. J. [2] 15, 349) by heating Al with Na CO, to the highest temperature of a wind furnace for some hours in a graphite cruoible.

and treating the residual mass with HClAq. The compound is pale-yellow when amorphous, bright honey-yellow and very lustrous when orystallised; brittle, not hard enough to scratch glass; in moist air it sawly crumbles down to Al₂O₂ with evolution of AH₃; it is dissolved by aqueous alkalis with evolution of NH, and solution of Al; fused with KOH or NaOH, an aluminate is formed and NI₂ is givon off; heated in air, NH₃ is evolved and Al₂O₃ remains.

Aluminium, Oxide of. [There are some indications of the existence of an oxide containing more O than Al_O₃ (v. Gibson a. Morrison P.R.S.E. 119, 146, 152).] Al_O₃ (Alumina) Mol. w. unknown. S.G. (amorphous) 3·725 to 4·152 (Roso, P. 74, 429); (crystallino) 3·928. S.H. (corundum, 9°-98°) ·19762; (sapphire, 8°-97°) ·21733 (Regnault, A. Ch. [3] 1, 129).

Occurrence.—Native, nearly pure as corundum, sapphire, topas, amethyst, &c.; also in opaque variety as emery.

Formation. - I. Amorphons. - 1. By burning powdered Al in oxygen. - 2. By ppg. boiling solution of potash-alum by (NH₄),CO₃Aq, washing, and strongly heating.—3. By strongly heating ammonia-alum.-4. By digesting clays, felspathio rocks, &c., with con. KOHAq under pressure, and ppg. by CO₂.—5. By heating a mixture of cryolite and lime in steam, lixiviating, and ppg. by CO.-H. Crystalline.-1. By fusing the amorphous Al, O, in the oxyhydrogen flame (Gaudin, C. R. 49, 1342) .- 2. By heating equal parts of potash alum and K,SO, with charcoal (Gandin, A. 103, 92).—3. By fusing together Al phosphate with three or four parts of K₂SO₄ or Na₂SO₄ (Debray, C. R. 52, 895).—4. Along with Al₂O₃.H₂O₄ by heating a solution of Al.O₃ in HClAq to 350° in a closed tube (Senarmont, C. R. 32, 762).—5. By heating to bright redness equal parts of amorphous Al,O, and PhO (Deville a. Caron, A. Ch. [4] 5, 104).

Preparation.—The amorphous variety may be prepared by heating dry potash-alum for two or three hours to reduces, finely powdering the residue, washing with water, mixing with NaOHAg containing 15 as much NaOH as the potash-alum used, drying, strongly heating, and washing with water (Wöhler, A. 53, 422; Brunner, P. 98, 488). Crystalline Al₂O₃ may be prepared by heating to whiteness a mixture of one part amorphous Al₂O₃ with four parts fused borax (Ebelmen, A. Ch. [3] 33, 34). Largo crystals, 1 cm. long were obtained by Deville a. Caron (A. Ch. [4] 5, 104) by the action of Al.F., on boric acid at a high temperature. The Al.F., was placed in a graphite crucible, the boric acid being contained in a small Pt basin fixed above the Al.F., this whole was placed in a Hessian crucible and heated in a wind furnace for some time; BF, was volatilised and crystals of Al₂O₃ remained.

remained.

Properties.—Known in two forms, amorphous and crystalline. The amorphous variety is a white, soft, powder; cakes together when strongly heated, and becomes nearly as hard as corundum; infusible except in oxyhydrogen blowpipo; insoluble in water; soluble in acids and aqueons alkalis, but after strongly heating becomes insoluble in acids except con. H.SO.Aq and con. HCIAq. The crystalline variety forms colourless rhombohedra; insoluble in all acids; nearly as

hard as diamond. Both forms are undecomposed by heat, and are unacted on by Cl.

Reactions.—1. With acids amorphous Al₂O₃ resots to form Al salts—e.g. Al₂SSO₄.—2. Fused with potash or NaOH, or KHSO₄, both amorphous and crystalline Al₂O₃ form aluminatos (g. v.) whi(h are soluble in water.—3. Amorphous Al₂O₃ heated to whiteness with potassium is partly deoxidised with formation of an alloy of K and Al. 4. Heated with sal ammoniac, Al₂O₄ is formed; the same compound is produced by the action of hot BCl₃, or SiCl₄.—5. Al₂S₃ is said to be formed by the action of carbon disulphide vapour on hot Al₂O₃.—6. Water is taken up by slightly heated amorphous Al₂O₃, but no definite hydrates have been thus obtained (v. Aluminum, Hydrausders).

Aluminium, Oxychloridss of. A series of these compounds seems to exist; they may be obtained by the action of a mixture of Al₂Cl₄ vapour and O on Al; the higher the temperature the more O is there in the product; they are soluble in dilute acids and alkalis, and are decomposed by water (Hautefeuille a. Perrey, C. R. 100, 1219). Tommasi (Bl. [2] 37, 443), describes three compounds of Al₂O₆H₆ with Al₂Cl₆ obtained by the action of Al on CuCl₂Aq under different conditions.

Alumininm, Phosphide of. Described by Wöhler (P. 11, 160) as a dark groy mass, which decomposes H₂O evolving PH₂; produced by heating powdored Al to redness in vapour of P.

Aluminium, Salts of. Salts obtained by replacing II of acids by Al. These salts belong to the form Al₂X_s, or AlX_s, where X =

$$NO_3$$
, $\frac{PO_4}{3}$, $\frac{CO_3}{2}$, $\frac{SO_4}{2}$ &c. besides these, many

basic salts (compounds of normal salts with Al₂O₂,21I₂O) are known. Very many Al salts also form double salts; the most characteristic of which are the alums Al 3SO, M, SO, 24H2O, where M = an alkali metal, Ag, or Tl (v. ALUMS). The haloid salts, the normal nitrate, sulphate, and acctate, are soluble in water; most of the other normal salts, and almost all the basic salts, are insoluble in water. The soluble salts possess a sweetish, astringent tasto. Aqueous solutions of Al salts generally contain more or less free acid, which is not, however, to be detected by the ordinary tests; Erlenmeyer a Lowinstein (Z. 3, 572) add freshly ppd Mg.NH, PO,, which pps. Al₂(PO₄)₂, and forms MgSO₄ and (NiI₂)₂SO₄, the acid can then be detected by litmus &c. The number of Al salts is not very large; with some acids, c.g. sulphurous carbonic, &c., it forms no salts or very unstable ones which can hardly be obtained pure. The chief salts—described under Borates, Pros PHATES, &c. - are the borates, nitrates, phos phates, silicates, and sulphates: v. also CARBO NATES, SULPRITES, SELENATES, &o.

Aluminium, Selenide of. According t Wöhlor, Al when heated in Se vapour com bines with the latter to form a black powde which is decomposed by H₂O into Al₂O₂xH₂(and H₂Se (P. 11, 160).

Aluminium, Silicides of, Al and Si read in almost all proportions; the products seer to be of the nature of alloys. When Al i heated with silicates in presence of a flux,

portion of the silica is reduced and combines with part of the Al. An alloy containing 10.8 p.o. Si, called cast aluminium, ie grey and very brittle; an alloy with 70 p.o. Si still exhibite metallic properties. The alloye of Si and Al are much more easily acted on by reagents than either of the elements which form them.

Alnminium, Sulphide of. Al.S.

If Al is heated to glowing and S is then thrown on to it, a black mass is produced which is decomposed by ILO with evolution of H.S. A mixture of Al₂S₃ and Al₂O₃ is obtained by passing CS₂ vapour over red hot Al₂O₃. When Al₂.3SO₄ is heated in H, Al₂O₃ remains and H₂SO₄ is volatilised (Wöhler, P. 11, 160). The best method of preparing Al₂S₃ seems to be to pass S vapour over hot Al in a carbon boat placed in a porcelain tube kept full of H. It is described as yellow crystals, with a bitter taste, which melt with difficulty, and are rapidly acted on by water with formation of H2S and Al2O3.xII2O (Fremy, A. Ch. [3] 38, 322; Sabatier, A. Ch. [5] 22, 88; Reichel, J. pr. [2] 12, 55). Spring (Bl. [2] 39, 64) obtained a sulphide of Al by vory strongly compressing an intimate mixture of Al and S.

Aluminium, Telluride of. Described by Wöhler as a black powder produced by heating

together Al and To (P. 11, 160).

M. M. P. M.

ALUMINIUM AMYLATE v. AMYL ALCOHOL.

ALUMINIUM BROMIDE, Action of, on Organio Bodies.-Aluminium bromide assists the bromination of aromatic hydrocarbons. action appears to be preceded by the formation of a compound AlBr₃60_nH_{2n-6} (Gustavson, J. 1877, 400; 1878, 380; B. 18, Ref. 208). AlBr₃ also combines with butylene; thus, when HBr mixed with C.H. is passed into AlBr, at 60°, the compound AlBr, C.H. is formed; the same compound is formed from AlBr, and C.H.Br, and by passing HBr into American or Caucasian petroleum containing AlBr_s at 70° (Gustavson, J. R. 16, 95; B. 17, Ref. 163).

The compound AlBr_sC₄H₅ is an oil, insol. in light petrolcum or CS.; it is decomposed by water with formation of unsaturated hydrocarbons and by alkyl bromides at 70° with formation of paraffine (Gustavson, J. pr. [2]

34, 161).

Aluminium bromide converts alkyl chlorides into bromides.

ALUMINIUM ISO-BUTYL C12H27Al i.e. Al(C4H9)2. From Al and Hg(C4H9)2. Fuming liquid (Cahours, J., 1873, 522).

ALUMINIUM ISO-BUTYLATE v. iso-Butyl ALCOHOL.

ALUMINIUM CHLORIDE, Action on Organic Bodies.-Aluminium chloride is converted by C₂H₄ and HCl at 100° into the compound AlCl₂C₄H₈ (G.).

Aluminium chloride added to a mixture of an aromatic hydroearbon and an alkyl or alkoyl chloride, bromide or iodide, promotes the evolu-tion of HCl, HBr, or HI and is therefore a most powerful agent in organic synthosis as a means of introducing alkyl or alkoyl groups into an aromatic nucleus (Friedel a. Crafts, G. R. 84, 1392, 1450; 85, 74, 672; 86, 1368). Thus methyl-benzenes may be formed by this means from benzene and methyl ohloride; acetophenone |

from benzene and acetyl chloride. The operation is performed by dissolving the aromatic hydrocarbon and the haloid derivative in CS, or light petroleum, and adding AlCl, in successive small portione. The reaction is completed by heating on a water-bach. Condensation is also brought about by AlCl, by the removal of water; thus benzeno and Ac₂O form acetophenone; while benzene and phi alio anhydrido form benzoyl-benzoio acid. Under the influence of AlCl, other reactions also occur; thus benzene is converted by oxygen, sulphur, sulphurous aoid, and carbonic acid into phenol, phenyl-mercaptan, benzene sulphinic acid, and benzoic acid, respectively.

These reactions are perhaps due to the formation of such compounds as AlCl.3C.H. in which the benzene may be supposed to be more unstable than when in the free state; thus, we might imagine the compound to be AlH.Cl.Ph. Molecular changes may, however, take place in the alkyls; thus both n- and iso-propyl bromides are converted by AlBr, into the same isopropyl-benzene; this is because n- propyl bropropyr beuzene, this indicates the building mide is changed by AlBr., into its isomeride (Gustavson, J. 1878, 380; P. 16, 958).

In the above cases AlCl., induces the building

up of more complicated compounds, but this reaction may be reversed and alkyl groups removed instead of introduced. Thus ethylbenzene, heated with AlCl, in a stream of HCl, evolves ethyl chloride and is reduced to benzene (Jacobsen, B. 18, 338). When the alkyl chlorido is not carried off by a stream of HCl it may act by substitution upon another portion of the hydrocarbon. Thus tolueno is converted by boiling with AlCl_a into benzeno and xylene (Anschütz a. Immendorff, B. 17, 2816). Somewhat similar reductions occur in other cases, the hydrogen being derived from another portion of the hydrocarbon. Thus naphthalene gives naphthalene dillydride, by reduction, and iso-dinaplitlyl, by abstraction of hydrogen; benzene gives toluene and cthyl-benzene togother with di-tolyl; di-phenyl-methano is reduced to bonzeno and toluene (Fricdel a. Crafts, B. A. 1884, 468; C. R. 100, 692).

AlCl, acting upon n-propyl iodide gives propyleno and HI, which then react, producing propane (Köhnlein, B. 16, 560). Aluminium bromide or iodide acting on propyl chloride or iodide at 130° form propylono but no propane (Kercz, A. 231, 286). AlCl, acting upon alcohols and pheuols eliminates HCl; thus phenol gives (PhO)₃Al₂Cl₃; resorcin gives (C₆H₄O₂)Al₂Cl₄; and di-chlorhydrin gives C₈H₃Cl₂O.AlCl₂, which erystallises from CS₂. These compounds are at once decomposed by water into the alcohol, Al(OH), and HCl (Clans a. Mercklin, B. 18, 2932) Acetyl chlorido in CS₂ is converted by AlCl₃ into a white solid, C₁₂H₁,O₂Al₂Cl₃, whence water produces a liquid C₃H₂O₂ (137°) which may be acetyl-acetone (Combes, C. R. 103, 814).

ALUMINIUM ETHIDE AIC, H, i.e. AlEt, (194°). V.D. 4.5 (for 3.9) at 234°. From mercurio ethide and aluminium heated at 100° for some hours (Buckton a. Odling, Pr. 14, 20). Liquid; fumes violently in air, soon taking fire. Decomposed by water. Iodine forms Etl.

Alumininm iodo-ethide Al2CeH, I, Al₂Et₃I₃, (840°-350°), ie a fuming liquid formed [8] 58, 5; A. 114, 242). ALUMINIUM ETHYLATE v. ALCOHOL.

ALUMINIUM 10DIDE, Action on Organic Bodies. All, turns out & displacing it by I, e.g. $BCCl_4 + 4AlI_3 - 3Cl_4 + 4AlCl_3$ (Gustavson, A. 172, 173). Aluminium iodide and aluminium are eingly without action on alcohol, but together they react, fa mixing Al₂I₂(OEt)₃ and Al₂(OEt)₆. One moleculo of aluminium iodide may convert vory many molecules of aluminium into aluminium ethylate (v. Alconol). Other alcohols act similarly (v. Propyl, Butyl, and Amyl, alcohols, Cresol, Phenol, Naphthol THYMOL), but methyl and iso-propyl alcohols, glycol, and glycerin do not; the latter forms allvl alcohol (Gladstone a. Tribe, Pr. forms allyl alcohol (Gladstone a. 80, 546, C. J. 39, 2; 41, 5; 49, 25; Hodgkinson, C. N. 1877, 237). Aluminium behaves in a eimilar way towards water and ether; it does not attack these bodies until after iodine has been added, when it reacts with water thus: $3H_2O + Al = Al(OI1)_3 + H_3$, with which may be compared its action on alcohol: 3HOEt+Al= Ai(OEt), + 1I, while with ether it forms alu-

minium iodo-ethylate and Etl. ALUMINIUM METHIDE AlC, H, i.e. AlMe, [0. 0°]. (130°). V.D. 2·8 (for 2·5) ut 220°; 4 at 160°. From HgMe, and Al at 100° (Buckton a. Odling, Pr. 14, 19). Takes fire in air. Resembles AlEt, Just above its boiling-point the

compound seems to be Al.Mes. ALUMINIUM PROPYL

AlC₂H₂₁ i.e. Al(C₃H₂)₂, (240°-245°). From HgPr₂ and Al (Cahours, J. 1873, 518).

ALUMS .- Double sulphates or selenates having the composition expressed by the general formula M.3S(orSe)O.N.S(orSe)O.24H.O, where M = Al, Cr, Mn, Fc, In, or Ga; and N - Na, K, Rb, Cs, NH, Ag, or Tl. These salts crystallise in forms belonging to the regular system, usually in octahedra or cubes. The following are the bost-known alums:—I. Sulphates: (1) M=Al, and N=Na, K, Rb, Cs, NII_1 , $NH_3(C_2H_5)$, Ag, or T1; (2) M = Cr, and N = Na, K, or NH_1 ; (3) M =Mn, and $N = K_1$ or NH_4 ; (4) M = Fe, and $N = K_1$ or NH_4 ; (5) M = Ga or In, and $N = N11_1$; (6) Mark Gral, and N=K, or NH, 11. SELENATES: (1) Maral, and N=Na, K, or NH; (2) M=Cr, and N=K, or NH, IH. MIXED SELENATES AND SULPHATES, M, 3SeO, N, SO, 24H, O: (1) M=Al, and N=K; (2) M=Cr, and N=K; (3) M=Fe, and N=K. IV. MIXED SULPHATES AND SELENATES, M. SSO. N. SSO. 24H.O: (1) M=Al, and N=K; (2) M=Cr, and N=K; (3) M=Mn, and N=K; (4) M=Fe, and N=K. Besides these, there are certain double salts which resemble, but are not isomorphous with, the alums; of these psendo-alums the most important are the following:—I. Compounds of Al₂3SO₄ with (1) MnSO₄, (2) FeSO₄, (3) MgSO₄; each with 24H₂O₅ II. Compounds of Fe₂3SO₄ with (1) MgSO₄, (2) CuSO₄, (3) ZnSO₄; each with 24H₂O. III. Mn₂3SO₄.MgSO₄.24H₂O.

In naming the alums, if no prefix is used, a donble sulphate of aluminium and one of the metals represented by N in the formula is understood; thus potassium alum is

Al₂3SO, K₂SO, 24H₂O.

The sulphates containing no aluminium are

by action of Etl on Al at 130° (Cahours, A. Ch. | monium alum, &c. Similarly, the names ammonium selenio-alum, and chromium-potaseium selenio-alum are used for the members of Group II. The salts belonging to Groups III. and IV. may be called eelenio sulphuric alums; the individual bodies are best distinguished by their formulæ.

The alums are all soluble in water, their solubility in hot, being considerably greater than in oold, water; (potash alum, S. 3.29 at 0°; S. 22 at 30°; S. 31 at 60°; S. 357 at 100°). The solutions have a styptic taste and an acid reaction. Some of the alums are separated by water into their constituent salts, e.g. silver alum, and manganese-ammonium and potassium alums. Others are partially separated; indeed it appears very probable that every alum is to some extent separated into its constituents when dissolved in a considerable quantity of water. Thus, Favre and Valson (C. R. 74, 1165) find the heats of solution of one equivalent of aluminium sulpliate (1) in water, (2) in a solution of K₂SO₄, and (3) in a solution of (NII₄)₂SO₄, to be the same (about -8,000); hence no combination occurs between the two sulphates in the presence of water. Further the heats of solution in water of the alums have large negative values, and in some cases—e.g. iron-ammonium alumthis value increases considerably as temperature rises (Favre and Valson, C. R. 74, 1016). G. Wiedemann (P. 126, 1; 135, 177), by determining the specific magnetism (that is, magnetic moment developed by unit magnetising force+ mass of salt in unit volume) of various salts in solution, has shown that an uqueous solution of ferrie sulphate is partly separated into sulphnrio acid and colloidal ferric oxide, and that the amount of this separation increases the larger the quantity of water added; he has also shown that an aqueous solution of iron-ammonium alum behaves almost exactly in the same way as ferric sulphate; hence the separation of the ferric sulphate by the water is independent of the alkalino sulphate; and hence a diluto aqueous solution of this alum is to a large extent separated into its constituent salts. When an aqueous solution of potassium alum saturated at 12° is heated to 100° a precipitate is slowly formed containing varying quantities of Al₂O₃, K₂O, SO₃, and H2O; even after 30 days the precipitate continues to be produced. The decomposition of the alum is hastened by adding K2SO4 to the solution (A. Naumann, B. 8, 1630). Chromiumpotassium alum exists in two forms; as violet erystals, and as a green non-crystallisable salt. When a solution of the former salt in water is heated to 70°-80° the colour changes to green, and this change is attended with a gradual increase of the volume of the colution; as the green solution cools the colour slowly changes to violet-blue, and the volume of the liquid slowly doereases (Boisbaudran, C. R. 79, 1491). The violet erystals change at 300°-350° to the green salt with loss of all their water of erystallisa. tion; this green dehydrated alum is wholly soluble in hot water, but when heated somewhat shows 350° it suddenly changes to greenish-yellow and becomes quite insoluble in water (Lowel, A. Ch. [8] 44,313; Siewert, A. 126, 86). When a quantity of barium chlorida. spoken of as chromium-sodium alum, iron-am- sufficient to precipitate all the acid in a given

mass of chromium potassium alum is added in four equal portions to an aqueous solution of the green form of this alum, the first and second fourths of the barium are at once precipitated, but the rest only very slowly; moreover, the quantity of heat produced during the precipitation of the first and second fourths is much greater than that produced during the subsequent precipitations (Favre and Valson, C. R. 74, 1165).

Gram-units of heat produced.

Excess of BaCl . 1st fourth, 2nd fourth, 3rd & 4th fourths. 8251 4104 4102 146

These results taken together show that solution in water of the commoner (probably of all) alume is accompanied by partial separation of these compounds into their constituent salts, and also by partial decomposition of these constituente, certainly at least of the sulphate of the heavy metal; and that the amount of this separation and decomposition is increased by increasing the quantity of water, or by raising the temperature.

The alums are delightrated by the action of heat; at a higher temperature a portion of the acid radicle is usually volatilised, and a double basic sulphate remains, such as native alum stone Al₂3SO₄.K₂SO₄.2Al₂O₅.8H₂O; at a still higher temperature more acid is removed and a mixture of alkaline sulphate and oxide of Al, Cr, Mn, or Fe remains; the ammonium alums leave a residue of oxide only. For descriptions of the properties of the individual alums v. Sulphates. M. M. P. M.

DI-ALURIC ACID v. DIALURIC ACID.

AMALGAMS. Alloys of Mercury (v. ALLOYS). Amalgams are formed (1) by direct union of mercury with other metals; e.g. amalgams of alkali metals, of Zn, Pb, Sn, Au: (2) by precipitation of other metals from solutions of their salts on mercury; this is often done by placing sodium-amalgam in the solution of a metallie salt, sometimes by electrolyeing a metallic solution in presence of mercury; e.g. amalgams of Ag, Fe, Co, Ni, Mn, Ba: (3) by precipitation of mercury on another metal, sometimes it is necessary to electrolyse the mcrcurial solution, making the other metal one of the electrodes; e.g. amalgams of Cu, Ag, Au, Pt: (4) by placing the other metal in contact with mercury and a dilute acid; e.g. Zn amalgam. The formation of amalgams is not usually attended with any marked thermal change, but in the production of amalgams of the alkali metals much heat is produced. Thus, [Na, Hg] = 10,300; [Na, Hg"] = 21,600; [K, $Hg^{\frac{1}{2}}$] = 20,300; [K, $Hg^{\frac{1}{2}}$] = 34,200 (Berthelot, C. R. 88, 1110, 1335). In the formation of amalgams of Sn, Pb, and Bi, heat is absorbed. Little or no contraction of volume accompanies the formation of amalgams, except in the cases of Cu, Ag, Sn, Pb, and a few other metals. The relative conductivity for heat of some solid amalgams is considerably greater than that of either of the metals composing them; e.g. amalgams of Sn, Zn, and Bi. Many solid amalgams seem to be chemical compounds in definite proportions; thus when various amalgams containing excess of mercury were subjected to a pressure of about 70 tons on the square inch mercury was removed, and definite bodies remained, containing mercury and metal approximately in the ratios expressed by the formula CuHg, AgHg, FeHg, Zn,Hg, Pb,Hg, PtHg, (Joule, C. J. 16, 878). Again, when amal-gams of An, Ag, Cu, Pb,K, and Na, were heated near to, to, or above, lie boiling-point of mercury (360°), the following definite amalgams were obtained (De Souza, B. 8, 1616; 9, 1050):-At 310° Au, Hg Ar, Hg Cu, Hg

" 860° Au, Hg Ag, Hg Cu, Hg Pb, Hg " 440° Au, Hg Ag, Hg Cu, Hg K, Hg Na, Hg. Some of the liquid amalgams may be regarded as solutions of definite compounds in excess of mercury, e.g. liquid Na and Kamalgams; others as solutions of metals in mercury, e.g. some of the iron amalgams. For descriptions of individual amalgams see the articles on the different metals, also MERCURY. M. M. P. M.

AMALGAMATION. The process of forming

Amalgams, q.v.
AMALIC ACID C12H14N4O8

i.e. C_sH₂Me₄N₄O₆. Tetra-methyl-alloxantin. Formation.—1. On mixing a solution of di-

methyl-alloxan with one of di-methyl-dialurio acid, amalic acid is ppd. (Maly a. Andreasch, M. 3, 103) .- 2. A product in the oxidation of caffcine by chlorine or nitrie acid (Rochleder. A. 71, 1).—3. By reducing di-methyl-alloxan by H₂S (E. Fischer, B. 14, 1912).

Properties.—Transparent colourless crystals. Stains the skin red. V. sl. sol. cold water or alcohol, sl. sol. hot water. Reduces silver salts. It forms dccp violet compounds with baryta,

KOII, or NaOH.

Reactions.-1. Oxidised by nitric acid to di-methyl-alloxan, or, better, by passing chlorine into water in which it is suspended .-2. Amalic acid may be distilled without leaving a residue, but it is decomposed and in the distillate there is a crystalline acid, $C_{12}H_{14}N_1O_6$, 'desoxyamalic' acid [260°]. This acid is v. sol. chloroform or glacial acetic acid, sparingly so in cold alcohol, water or ether. Soluble in alkalis but reppd. by HCl. 1s partly decomposed when distilled. Reduces boiling ammoniacal AgNO₃. Evaporated with HNO₂ it forms dimethyl-alloxan. Chromic mixture converts it into cholestrophane. Hence it is possibly (Fischer a. Reese, A. 221, 339);

Me.N - COCO.NMe

conc - ch co

MeN - CO ĊO.NMe

3. Hydrogen sulphide forms di-methyl-dialuric acid (M. a. A.) .- 4. By boiling with water in an open vessel di-methyl-oxamide is produced:

 $C_{12}H_{14}N_4O_8 + H_2O + O_2 = 2C_4H_8N_2O_2 + 4CO_2$.

Ammonia gas turns it violet, forming murexoin, a drystalline body resembling murcxide .- 6. From the solution made by heating amalic seid (4 pts.) with cyanamide (2 pts.) and water (100 pts.) there separates out, on cooling, crystalline cyamido-amalic, acid C₁₂H₁₄N₈O₂. Long prisms, el. sol. cold water, v. sol. hot water, insol. alcohol or ether; reduces silver salts, and yields methylamine and an oxalate when boiled with alkalis. It gives off purple vapours when heated, and forms a sublimate (Andreasch, M. 3, 433).

AMANITINE v. NEURINE.

AMARINE v. BENZOIC ALDERYDE.

AMBER. A fossil resin from Pinites succinifer. S.G. 1.05 to 1.10. It coutains from 8 to 8 p.o. snecinio acid, also a resin [146°] soluble in ether but not in alcohol, a resin '105'] soluble in alcohol and other bodies. There is about 1 p.o. ash (O. Helm, P. [3] 11, 229). When amber is distilled, succinic acid and oil of amber are got. The latter (110°-260°) is a mixture of terpenes (Pelletier a. Walter, A. Ch. [3] 9, 89).

AMBREIN. [25°-30°]. Extracted from ambergris by hot alcohol. Ambergrie is found in the intectine of the spermaceti whale in tropical climates, and also floating in the sea. Ambrein is insol. water, v. sol. alcohol or ether, neutral to litmus. It cannot be saponified. It greatly resembles choleeterin (Pelletier, A. 6, 24; J. Ph. 5, 49).

AMENYL GLYCERIN v. TRI-OXY-PENTANE. AMENYL-VALERIC ACID v. DECENOIC ACID. **AMETHENIC ACID C**₇H₁₄O₂ (185°-230°): By oxidation of diamylene with ohromic mixture (Sohneider, A. 157, 209).

Properties .- A light oil.

Salts.—SrA'2, 8aq: needles.—Zn A'2: warte; aqueous solution forme a gelatinous pp. on heating.—AgA': difficultly soluble pulverulentpp.

AMIC ACIDS. Bodies represented by the formula X"(CO2H).CO.NH2, being at once amide

and acid, e.g. oxamio acid.

AMIDES. Bodies derived from ammonia by displacing one-third of its hydrogen by a monovalent acid radicle (Gerhardt a. Chiozza, A. Ch. [3] 46, 129). When two-thirds of the hydrogen is displaced by a divalent radicle the product is called an imide, but if the same amount of hydrogen is displaced by two monovalent radioles the product is still called an amide. If the whole of the hydrogen is displaced the product is also called an amide unless it is displaced by one trivalent radicle, when it is called a nitrile.

Amidce derived from ammonia by displacing one-third of the hydrogen by an acid radicle, e.g. NH, CO.X, may also be looked upon as derived from mono-basic acids, X.CO.OII, by displace-

ment of OH by NH₂.

Di-basio acids, Y"(CO.OH)₂, form, in a similar way, first amic acids, Y"(CO.OH)(CO.NIL), theu di-amides, Y"(CO.NH2)2. Thus a di-amide may be coneidered to be derived from two moleculee of ammonia by displacing one-third of the hydrogen by a di-valent acid radicle.

Formation.-1. By action of ammonia on ethere: X.CO.OEt + NH₂ = X.CO.NH₂ + HOEt.-2. By the action of ammonia on acid chlorides:

 $X.CO.Cl + 2NII_3 = X.CO.NH_2 + NH_1Cl.$ 3. By the dehydration of ammonium salts: $X.CO.ONH_4 - H_4O = X.CO.NH_2$ The rate at which thie decomposition is brought about by heat has been studied by Menechutkin (C. R. 98, 1049).—4. By action of NH, on anhydrides: $(X.CU)_2O + 2NH_3 = X.CONH_2 + X.CO.ONH_4$

The anhydridee of di-basic acide are converted by this reaction into amic acide or imides. -5. From nitrilee and cold conc. hydrochloric acid $XCN + H_2O = X.CO.NH_2 - 6$. From nitriles, hydrogen peroxide, and very dilute KOH (Radziszewski, B. 18, 355).—7. Prepared by heating acids with ammonio sulphocyanide for three or four days, e.g.: NH₄SNC+2HOA0=2AcNH₂+COS+H₂O

(J. Schulze, J. pr. [2] 27, 512).

Properties.—The amides are usually solid, and their melting-points serve to identify the several acide. They are neutral to litmus, but form oompounds with acids; their typical hydrogeu can in many cases be displaced by metals (Ag or Hg). The typical hydrogen can be dieplaced by alkoyls, by troatment with aoid chlorides, but heating with alkyl iodides does not result in the introduction of alkyls unless sodium has been previously introduced. Alkylated amides oan bo formed by the action of alkylamines upon acid chlorides or ethers.

Reactions.-1. Converted by dehydration (e.g. with P_2O_3) into nitriles: X.CO.NH₂ = $X.C_2N + H_2O_4$ 2. Converted into acids by boiling potash, boiling dilute IICl, fuming HNO₃, or nitrous acid: X.CO.NH₂+HNO₃ = X.CO.OH+H₂O+N₂O

(Franchimont, R. 2, 343).

X.CO.NH₂ + HNO. = X.CO.OH + H₂O + N₂.

3. One of the typical atoms of hydrogen may be displaced by halogens; when bromine and alkalis are both present, compounds of the form X.CO.NKBr and X.CO.NKBr, are produced; these readily split up into KBr and oyanio ethers XNCO (Hofmann, B. 17, 1106; 18, 2734).

 Phosphorus pentachloride reacts thus: X.CO.NII₂ + PCl₂ = X.CCl₂.NH₂ + POCl₃ The resulting compound splits off IICl giving X.CCl₂NH₂=IICl + XCCl:NH a chloro-imide and X.CCl:NH = HCl + XC:N.—5. PCl₃ acts upon alkylated amides, forming compounds with twice as many carbon atoms in the mole-Thus CII. CONHEt givee CII, CCI: NEt

which then acts thus:

 $2CH_{3}CCl:NEt = NEt:CCl.CH_{2}.CMe:NEt + HCl.$ CH3. CONHPh acts similarly, while PhCO.NHPh and CClaCONHEt form only chloro-imides. Formanilide, IICONPhH does not produce NPh:CCl.CII:NPh but NPh:CH.NPhH, diphenyl-formamidine (Wallach, A. 184, 1; 214,

The HCl liberated in the decomposition $X.CCl_{y}.NHY = HCl + XCCl_{z}.NY$ may often convert undecomposed amide into

amidine 2X.CO.NIIY + HCl = XCO2II + XC(NY)(NHY)IICL

Amidines may also be formed thus: XCONHY + XCCl2NHY

XC(NY)(NHY)IICI + XCOCL 6. Heated with alcohol they form alkylam. monium ealts:

 $X.CO.NH_0 + IIOEt = X.CO.O.NII_3Et$ X.CO.NHEt + HOEt = X.CO.O.NII_Et,

 $X.CO.NEt_2 + HOEt = X.CO.O.NHEt_3$ (Baubigny, C. R. 95, 646).-7. ZnEt₂ acts thus:

 $X.OQ.NH_2 + ZnEt_2 = X.CO.NZn + 2IIEt$ $Y''.(CO.NH_2)_2 + ZnEt_2 = Y''.C_2O_2N_2H_2Zn + 2HEt.$ These compounds are decomposed by water, with reproduction of the amide (H. Gal, C. R. 96, 1315).—8. Phenyl-hydrazine reacts thus: PhNH.NH.2+NH.2CO.X = PhNH.NH.CO.X + NH. (F. Juet, B. 19, 1201).-9. Dry HCl converts primary into secondary amidee:

 $2NAcH_2 + IICl = NAc_2H + NH_4Cl$

AMIDINES. The name is applied to compounds that contain amidogen and imidogen attached to the same atom of carbon. Thus, CH3.C(NH).NH2 is called acetamidine, that is to say, an imide derived from acetamide, while C.H.C(NH)NH2 is called benz-amidine, imide of benzamide. Other names for these

bodies are ethenyl amidine, benzenyl amidine, ethenyl imid-amide, bsnzenyl imid-amide, acet-The advantage imid-amide, benz-imid-amide. of the imido-amide nomenclature is chiefly seen in naming derivatives, thus CH2.C(NEt).NH2 and CH3.C(NH).NEtH may be called acet-ethyl-imido-amide, and acet-imido-ethyl-amido, respectively. But the advantage so gained is lost in the great lengthening of the names, and both bodies will therefore be called ethyl-acetamidine in this dictionary. They might be distinguished as tertiary and secondary thyl-acetamidine respectively.

Dibasio acids can give rise to a variety of amide-imides, for which Wallach (A. 214, 256) proposes the following nomenclaturo:-

 $R < {}^{CO.NH_2}_{C(NH).NH_2}$ amid-amidine.

 $R{<}^{C(\rm NH).NH_2}_{C(\rm NH).NH_2}$ amidine.

 $R{<_{C(\mathrm{NH})}^{\mathrm{CO}}}{>_{\mathrm{NH}}}$ imid-imidine.

R < C(NH) > NH imidine.

Formation .- 1. Amidines are formed by the action of amines on thio-amides or nitriles: Ph.CN + NPh.H = Ph.C(NH).NPh2 (Bernthsen, A. 184, 290, 321).—2. By the action of amines on the compounds (X.CCl:NH or X.CCl:NY) formed by the action of PCl, on amides (Wallach, B. 8, 1575).

Reactions.—1. H₂S forms thio-amides:

Ph.C(NPh).NPhH + H.S = Ph.CS.NPhH + H.NPh $Ph.C(NH).NPhII + H_2S = Ph.CS.NPhII + NII_3$ Another reaction also takes place:

 $Ph.C(NH).NPhH + H_2S = Ph.CS.NH_2 + NPhH_2$ This may be explained by supposing an intermediate compound, Ph.C.(SII)(NH₂).NPhH, to be formed by addition of H.S.

2. CS, acts thus:

Ph.C(NH).NPhH + CS₂ = Ph.CS.NPhH + HNCS Ph.C(NH).PhP₂ + CS₂ = Ph.CS.NPh₂ + HNCS Ph.C(NPh).NPhH+CS2=Ph.CS.NPhH+PhNCS

3. Action of aceto-acetic ether, v. p. 19. The reactions of the amidines are further described in such articles as FORMAMIDINE, ACETAMIDINE, BENZAMIDINE, MANDEL-AMIDINE, PHENYL-ACRTAMIDINE, and PHENYL-BENZAMIDINE.

AMIDO-ACETANILIDE v. Acetyl-PHENYLENE-DI-AMINE.

AMIDO-ACETIC ACID v. GLYCOCOLL.

Acetyl derivative v. Aceturic Acid. Benzoul derivative v. Hipporic ACID. AMIDO-ACETO-ACETIC ACID v. ACETO-

ACETIO ACID. AMIDO-ACETO-NAPHTHALIDE v. ACETYL NAPHTNYLENE DI-AMINE.

AMIDO ACETOPHENONES Call, NO i.e. C.H. (NH2).CO.CH, Amidophenyl methyl ketone. o-Amido-acstophenone (o. 249°).

Formation.-1. By reduction of o-nitroacetophenone (Gevekoht, B. 15, 2086; A. 221, 826).-2. By action of conc. H2SO4 on a solution of o-amido-phenyl-acotylene (Baeyer a. Bloem, B. 15, 2154).—3. By boiling o-amido-phonyl-propiolic acid with water (B. a. B.).

Preparation.—o.Amido - phenyl - acetylone (50 g.) is slowly dropped into cono. H.SO. (600 c.c.) diluted with water (200 c.c.). After

half an hour, the mixture is poured upon ice; neutralised with Na₂CO₃; distilled with steam; and the distillate extracted with other. 50 p.c. of the theoretical yield is got (Baeyer a. Bloem,

of the theoretical yield is got (Baeyer a. Bleem, B. 17, 964). Propertier—Thick volatile oil.
Oxim [148°] (Munchmeyer, B. 20, 512).
Salts.—B'H.SO₄: needles.—B'H.GISnCl₂:
needles, sol. alcoliol.—B'_1l._PtCl_a: yellow pp.
Raaction.—By bo.hin, with alcoholio acetophenone and some NaOHAq it is converted into flavolin or phenyl-methyl-quinoline (O. Fisoher, B. 19, 1036): C₆H₄(NH₄).CO.Mo+Ph.CO.CH₂=

H₂O + C₆H₄ CMe:CH N : CPh

C.H.(NHAe).CO.CH. Acetyl derivative C₈H₄(NHAe).CO.CH₂.
[77°]. Silky needles (from benzoline). Sol. alcohol, ether, and hot water, sl. sol. oold water.

m-Amido acetophenone [93°]. Formed by reducing the nitro compound by Sn and HCl (Buchka, B. 10, 1714; Hunnius, B. 10, 2009; Engler, B. 11, 932). Short yellow pyramids, sol. alcohol, and ether.

Salt.—B'HCl: long pointed crystals.

p-Amido-acetophenone [106°].

Formation. - From the nitro compound (q. v.) by Sn and HCl (Drewson, A. 212, 162).

Preparation. - Aniline (2 pts.), ZnCl2 (3 pts.), and Ac₂O (5 pts.) are boiled together for 5 hours; the resulting acctyl derivative is saponi-fied. Yield 55 p.c. of the theoretical (Klingel, B. 18, 2687).

Properties.-Long fan-like crystals (from water). V. sol. alcohol, ethor and hot water, sl. sol. cold water, benzeno, and benzoline.

Salts.-B'HCl: needles. $-B'_2H_2PtCl_6:$ slender yellow needles. $-B'_2H_2SO_4:$ needlos. $-B'_2H_2C_2O_4:$ crystals, v. sol. alcohol.

Acetyl derivative C,H,(NHAc).CO.CH, [167°]. Small needles, v. sol. alcohol, and hot water, sl. sol. cold water.

Ethyl derivative v. ETHYL-AMIDO-ACETO-PHENONE.

Benzyl derivative v. Benzyl - AMIDO-ACETOPHENONE.

AMIDO ACIDS .- Amidogen, when attached to carbon in an acid, behaves as it does in amides (q. v.) or as in amines (q. v.) according as that carbon does or does not belong to carbonyl; in the former case the compound is classed as an amic acid (q. v.), the term 'amido acid' is usually restricted to the latter class of bodies.

Formation .- 1. From the halogen derivatives of fatty acids, or their ethers, by the action of ammonia .- 2. From the nitro-derivatives of (aromatic) acids by reduction .- 3. I'rom alde-

Nydes, by action of hydric cyanide and NH₂:

X.CHO + HCN + NH₂ = X.CH(NH₂)CN + H₂O.

The nitrile is then converted into amide by conc. HCl, and this is saponified by hot dilute HCl. In this way a-amido acids may be prepared; alkylamido acids can be formed by using, alkylamines instead of ammonia (Tiemann, B. 14, 1982; Stephan, C. C. 1886, 470).

Properties.—Neutral bodies which combine both with acids and bases. Their neutrality is probably due to self-saturation, as may be represented by a doublod formula: R.CH.NH, O.CO

CO.O.NH. CH.R. Reactions .- 1. Converted by nitrous acid

into oxy-acids (v. p. 57, l. 6); di-zeo deriva-tives (q. v.) are first formed, and this formation may be ntilised as a test for amido-acids (Curtius, B. 17, 959).—2. When heated with lims or baryta, they legit off CO₂ forming amines. This separation of CO₂ sometimes occurs in formation 2: thus C, H, (NO2)Br(CO2H) [4:2:1] reduces to m-bromo-aniline (Scheufelen, 4. 231, 176); C₃H₃(NO₃*₂C₅H [4:1:2] rednoes to m-phenylene diamine (Wurster, B. 7, 149; Griess, B. 7, 1225); while C₅H₄(NO₂)(CO₂H)₂ [4:1:2] becomes m-amido-benzoie acid. In all these cases the CO2 is split off from the position para to NO2.—3. AcCl forms acetyl-amido aoids. 4. Excess of methyl iodide, in presence of KOH, converts (fatty) amido acids into ammonium iodides:

 $CH_2(NH_2).CO_2H + 3MeI + 3KOH =$

 $OH_2(NMe_sI).CO_2K + 2KI + 3H_2O$ (Körner a. Menozzi, G. 13, 350). p-Amido-benzoic acid is converted by MeI and KOH into the betaine C.H. CO.O., while Etl only forms di-ethyl-amido-benzoic acid (Michael a. Wing, Am. 7, 195) .- 5. Saturated with cuprio hydroxide, suspended in hot water, they form blue solutions from which on cooling the copper salt separates. This occurs with leucine, glutamio acid, and aspartie acid. In the case of leucine, a portion remains dissolved, forming a blue mother liquor. In the case of mixtures of amido-acids, the copper salts are not so readily ppd., for thoy seem to render one another soluble (Schulze a. Barbieri, J. pr. [2] 27, 351).—6. The methods of displacing amidogen by halogens in aromatic bodies are mentioned under AMINES.

AMIDO ACRYLIC ACID C,H,NO, i.e. CH(NH2):CH.CO2H, is formed by the action of alcoholio ammonia on \$-chloracrylic acid at 100° (Pinner a. Bischoff, A. 179, 97). H. W.

AMIDO-ALCOHOLS or Alkamines (q. v.) aro formed by action of bases on chlorhydrins or on alkylene oxides, e.g.:

 $CH_2Cl.CH_2OH + NH_3 = CH_2(NH_2HCl).CH_2.OH,$

 $O + NMe_2 + H_2O = CH_2(NMe_3OH).CH_2OH.$

V. Oxy-ethyl-amine, neurine, &c. AMIDO-ALIZARIN v. OXY-AMIDO-ANTHRA-

DI-AMIDO-AMARINE v. Amarine under BENZOIO ALDEHYDE.

AMIDO-AMYLALCOHOL v. OXY-AMYL-AMINE. AMIDO-AMYL-BENZENE $C_{11}H_{12}N$ C.H.(C.H.,)NH. (258°) (C.); (260°–265°) (H.). An oil. Formed by heating amyl-anilino hydrochloride at 320° (Hotmann, B. 7, 529), or by heating aniline with amyl Alcohol and ZnOl, at 270° (Calm, B. 15, 1643).

Salts.-B'2H2SO4: silky needles.

B',H,PtCl,: slender orange-yellow needles. Benzoyl derivative C.H.(O.H.1)NHBz. [0. 1496]. Pearly plates; sol. alcohol, ether, benzene.

AMIDO-ANISIC ACID v. Methyl-oxy-amido-

BENEOIO ACID. AMKOO-ANISOL v. Methyl-amido-phenol.

AMIDO-ANTHRACENE v. ANTHRAMINE. AMIDO-ANTHRAQUINONES O. H.NO. M.w. 228. Three have been described, but theory indicates only two; (s) and m - are parhaps

o-Amide-anthraquinone

 $C_{s}H_{s} < \frac{CO(1)}{CO(6)} > C_{s}H_{s}.NH_{2}$ (2). [241°].

Formation.-1. By reducing o-nitro-anthraquinone (Roemer, B. 15, 1790).

Properties .- Ruby-red iridescent needles: may be sublimed. Sol. alcohol, ether, benzene and HOAc, forming orange liquids, v. sl. sol. water. It is a weak base, dissolving in conc. HCl. Converted by nitrous acid into erythrooxy-anthraquinone.

Salt .- B'HCl: unstable white needles. Acetyl derivative C11H,O2NAeH [202°].

Orange-red needlos, sol. alcohol and cold HClAq. (a)-Amide-anthraquinone [254°].

Formation .- 1. From bromo-nitro-anthraquinone (Claus a. Hertel, B. 14, 980) or from di-bromo-nitro-anthraquinone (Claus a. Dierenfellner, B. 14, 1334) by sodium-amalgam.-2. From (a)-nitro-anthraquinone and sodiumamalgam (Böttger a. Petersen, A. 166, 149)

Properties .- Red needles, may be sublimed. Sol. benzeno and chloroform, sl. sol. alcohol, and ether. Differs from the preceding by insolubility even in fuming HClAq.

m-Amido-anthragninone

 $C_{e}H_{s} < CO (1) > C_{e}H_{3}NH_{2} (3)$. [802°].

Formation .- 1. From anthraquinone m. sulphonic acid and NH, Aq at 200° (Perger, B. 12, 1566; according to Bouchardat, Bl. 33, 264, this reaction produces amido-oxy-anthraquinone) From its acetyl derivative, which is got by oxidising acetyl-anthramino by CrO, in glacial HOAo (Liebermann, A. 212, 61).

Properties. - Red needles. Solvaqueous HCl. insoluble in alkalis. By the action of HNO₃ and boiling alcohol it is converted into anthraquinone.

Acetyl derivative C14H,O2NAcH. [257°] (P.); [263°] (L.); colourless needles. Di-amido-anthraqninones

 $C_{14}H_{10}N_{2}O_{2}$ i.e. $C_{14}H_{6}O_{2}$ (NH₂)₂. (a)-Di-amido-anthragninene [236°].

Formation.-1. From (a)-di-nitro-anthraquinone either (a) by ammonio sulphide, (b) by aqueous NH₃ at 200°, nitrogen coming off (J. Fischor, J. pr. [2] 19, 209), or (c) by SnCl₂ and NaOHAq (Böttgor a. Petersen, A. 160, 148). 2. By reduction of tetra-bromo-di-nitro-anthraquinone (Claus a. Hertel, B. 14, 981).

Properties. - Red needles (from ether), with V. sl. sol greenish reflex (when sublimed). water, m. sol. alcohol, ether or acctone, v. sol. Hardly The solutions are purple. benzene. soluble in filute acids; does not form salts.

Reactions.-1. Nitrous acid passed into its alcoholie solution forms anthraquinone (B.a.P.). 2. Nitrous acid passed into its ethereal solution forms a brownish-violet powder, O₁₄H₈N₄O₄, which detonates at about 68° (B.a.P.).—3. Potashfusion produces alizarin (Böttger a. Petersen; B. 4, 778), or some similar body (Liebermann, B. 4, 231, 779).

(β)-Di-amido-anthraquinene [above 300°]. Formation .- 1. By boiling (\$)-di-nitro-anthraquinone with SnCl, and NaOHAq (Schmidt, J. pr. [2] 9, 266).
Properties.—Reddish-brown powder; sqb-

limes in dark red needles. Sl. sol. water, v. sol. sloohol, ether, and benzene, forming red solutions. Sol. cono. acids, but re-ppd. unaltered by

(7)-Di-amido-anthraquinone.

Preparation.—Alizarin (20 grms.) is heated for 7 hours at 170° with ammonia solution 200 c.c., S.G. 915). Alcohol extracts the greater portion (6.2 grms.) of the insoluble residue (7.3 grms.). Water is added to the alcohol, and the pp. ie dried in vacuo and washed with ether (H. v. Perger, J. pr. [2] 18, 135).

Properties.-Indigo-blue powder, which acquires a coppery lustro when rubbed. When HCl is added to its blue alcobolic solution (at 0°), the liquid turns cherry-red and deposits brownred needles of a hydrochloride, which, however, is eo unstable as to be reconverted into the amorphous blue base by merely washing with water. It does not die mordanted goods.

Reactions .- 1. Boiled with potash it is converted into oxy-amido-anthraquinone (q. v.): $\mathbf{C}_{14}\mathbf{H}_{6}\mathbf{O}_{2}(\mathbf{N}\mathbf{H}_{2})_{2} + \mathbf{K}\mathbf{O}\mathbf{H} = \mathbf{C}_{14}\mathbf{H}_{6}\mathbf{O}_{2}(\mathbf{O}\mathbf{K})\mathbf{N}\mathbf{H}_{2} + \mathbf{N}\mathbf{H}_{3}$ 2. Similar reaction by boiling HCl.-3. Fused with potash, or heated with HCl at 250°, forms alizarin.-4. By passing N2Os into its alcoholio colution nntil the blue colour is changed to pure yellow, it is converted into erythro-oxy-anthraquinone, which is thrown down when water is Yield 95 p.c. added.

(8)-Di-amido-anthraquinone

(3) $N_{12} \cdot C_e H_1 < \stackrel{CO}{CO} \stackrel{(1)}{(2)} > C_e H_1 \cdot N_{12} \stackrel{(6)}{(6)}$.

[above 300°].

Formation .- By reducing the corresponding di-nitro-anthraquinone, [above 300°] (Rocmer, H. 16, 366).

Properties. -- Splendid red metallic needles (by sublimation). Sl. sol. alcohol, ether, acctone, and ohloroform, with orange colour; v. sl. bol.

water. Very weak base. Reactions.—1. Boiling potash has no action.
2. Diazotisation followed by boiling with water converts it into anthrarufin (v. Di-oxy-anthra-QUINONE).

Di-acetyl derivative C14H, O2(NHAe) Reddish yellow needlee, v. sol. alcohol, and other. AMIDO ANTHRAQUINONE SULPHONIC ACIDS

 $C_{14}H_0NSO_5$ i.e. $C_{14}H_6O_2(NH_2).SO_8H$ o-Amido-anthraquinone sulphonio acid

 $C_0H_4< \frac{CO(1)}{CO(2)} > C_0H_2(NH_2)(SO_2H)$ [1:6:2:4).

Formed by reducing nitro-anthraquinone sulphonic acid (Lifschutz, B. 17, 899). Silvery needles.

(a)-Amido-anthraquinone snlphonio aoid. Prepared by reducing (a)-nitro-anthraquinono sulphonic acid (Claus, B. 15, 1519). Sol. dilute acids, and in hot water, sl. sol. cold water, aloohol, and ether.

Salts.—NaA'1aq: small red needles.—CaA'25aq: red needles.—BaA'23aq: slender red needles. - CuA'27aq: yollowish-red needles.

(β)-amido-anthraquinone sulphonic aoid.-Formed by reducing the lead salt of (β) -nitroanthraquinone sulphonic acid with H2S (Claus, B. 15, 1520). Red powder; v. sol. water forming a red solution, sl. sol. alcohol, insol. ether. A weak acid.

(a) - Di-amido-anthragniuone sul. phonic soid

O, H, N, SO, i.e. C, H, O, (NH,), SO, H. Obtained from (a)-di-amido-anthraquinone by means of H₂SO, containing dissolved SO, (30 p.o.); ppd. by water. The solution is a splendid red. It may be orystallised from alcohol. Sol. glacial FOAo, and in acetic ether. Insol. ether, benzeno or benzoline. On passing nitrous gas into ita alcoholic solution anthraquinone (a)-sulphonic can't is formed. Potashfusion forms alizarin.

Salt .- BaA'2: insol. cold water (v. Perger,

J. pr. [2] 19, 209). AMIDO-ARACHIC ACID

C₂₀H₄₁NO₂ i.e. C₂₀H₃₉(NH₂)O₂. [59°]. From nitro-arachio acid and SnCl, (Tassinari, B. 11, 2031). Sl. sol. ether, m. sol. alcohol. Combines with neither acids nor bases.

AMIDO-AZO-COMPOUNDS v. Azo pounds.

AMIDO-BENZALDEHYDE v. AMIDO-BENZOIO ALDEHYDE.

AMIDO-BENZAMIDE v. AMIDO-BENZOIC ACID. AMIDO-BENZ-ANILIDE v. AMIDO-BENZOIG ACID.

AMIDO-BENZENE v. ANILINE.

Di-amido-benzene v. Phenylene di-amine.

con-Tri-amido-benzene C. II, N, i.e. C. H. (NH₂). [1:2:3]. [103°]. (336° cor.). Obtained by distilling tri-amido-benzoic acid with pounded glass (Salkowsky, A. 163, 23). Crystalline; v. sol. water, alcohol, and ether. Its aqueous solution is alkaline and gives with Fe₂Cl₂ first a violet, then a brown pp.; hypochlorites and nitrites give brown pps. Reduces cold ammoniacal AgNO₂Aq. H₂SO₄ containing a little HNO₂ forms a blue colour.

Salts. - B'211Cl: sl. sol. conc. hydric chloride B".-H.SO, 2aq.

i-Tri-amido-benzene C.H.3(NH2)3. [1:2:4].

[below 100°]. (o. 340°).

Formation.-1. From (a) di-nitro-aniline, Sn. and HCl (Salkowsky, A. 174, 265) .- 2. From diamido-azo-benzene p-sulphonic acid by Sn and HCl (Griess, B. 15, 2196).—3. From chrysoidin by reduction (Witt, B. 10, 658).- 4. From dinitro-benzene-azo-benzene sulphonic acid by reduction (Janovsky, M. 5, 159).

Properties.—Colourless plates. V. sol. water, and alcohol, sl. sol. ether. Gives a red (G.) or

green (J.) colour with Fe, Cl, Aq.

Salts.—B"H,SO4. Needles or prisms; sl. eol. cold water, v. sl. sol. alcohol.—B"2HCl [133°];

needles (Hinsberg, B. 19, 1253).

s-Tri-amido-benzene Calla(NH2), T1:3:51 (?) The tin double calt, CoH3(NH2)3(HCl)3SnCl2, of this base may be got from tri-nitro-benzene (got by nitration of di-nitro-benzene) by Sn and ... HCl (Hepp, A. 215, 348). Busafter removing the tin by H2S, the hydrochloride of the base recinifies, NH,Cl being formed, although evaporation in vacuo over H.SO, a very soluble white hydrochloride may be got. It gives no colour with Fe2Cla.

Tetra-amido-benzene C.H.(NH2), [2:2:4:5] Formed by reduction of di-nitro-m-phenyleno dia mine with tin and SnCl. The base is extremely oxidisable. An aqueous colution of the hydro

chlorido when treated with Fe,Cl, gives a pr of brown noedles of C,H,(NH,),(NH),H,Cl,. Salts.—B''H,Cl,: v. sol. water, el. sol. conc aqueous HCl.—B'',(H,SO),: sparingly solubl

large plates.—B"H,SO,: long sparingly soluble

needles (Nietzki a. Hagenbach, B. 20, 834).
AMIDO-BENZENE SULPHONIC ACIDS C.H.,NSO, i.e. C.H.(NH.).SO.H. Aniline sulphonic acids. In the bromination of these acids Br never takes a position to NH₂ (Limpricht, A. 191, 252).

o-Amide-benzene snlphenie acid

C₈H₄(NH₂)SO₃H • [1:2]. E. 1 at 7°. Formation.—1. From o-uitro-benzene sulphonic acid (Berndsen a. Limpricht, A. 177, 98). 2. From m-bromo-benzene sulphonic acid by nitration and reduction (Thomas, A. 186, 128).

Properties .- Dull white crystals like rhombohedra. Also, as HA' aq, in transparent shining prisms with many faces. Bromino added to a very dilute solution of the barium salt produces H₂SO₄, tri-bromo-aniline and (1, 4, 5)-bromo-amido-benzeno sulphonic acid.

Salts (Bahlmann, A. 186, 308).—KA' aq: prisms. — AgA'; needles.—BaA', (L. a. B.).— BaA', 2aq (T.) .- PbA', laq. S. 3'4 at 6°.

m-Amide-benzene snlphenic acid

C.H. (NH.,)SO.H [1:3]. S. 1.2 at 7°; 1.5 at 15°. From m-nitro-benzeno sulphonio acid by reduction (Laurent, C. R. 81, 538; Schmitt, A. 120, 164; Berndsen, A. 177, 82). Also from (1, 2, 4)- bromo-amido-benzene sulphonic acid and HIAq at 120° (Goslich, A. 180, 102).

Long slonder radiating needles. Also, with 1 aq in monoclinic prisms. Sl. sol. cold water, v. scl. hot water, insol. alcohol, and ether. Aqueous solution turns red in air. When heated

it decomposes without fusion.

Reactions.-1. Bromine added to an aqueous solution produces no tri-bromo-aniline, but (1, 3, 4, 6). di-bromo-amido-benzene sulphonic acid, (1, 3, 5, 4, 6)- tri-bromo-amido-benzene sulphonic acid, and bromanil. Culorine acts similarly (Beckurts, A. 181, 211).—2. Does not produce quinone when oxidised (Meyer a. Stüber, A. 165, 168).

Salts: BaA'₂ 6aq.—PbA'₂.

Amide C₆H₄(NH₂).SO₂NH₂. [142°]. From m-nitro-benzene sulphamide, conc. NH3Aq, and H₂S (Limpricht a. Hybbencth, A. 221, 204). White plates or long needles (from water).

Hydro-chloride.-C.H.(NH2Cl)SO2NH2

[235°]. Needles.

Nitrous acid, passed into a cold mixture of the amide with a little HNO3, produces a di-azo nitrate, C.H. (N.NO.).SO.NH., benzene sulphamide, and a diazo-amido compound

C₆H₄(SO₂NH₂).N₂.NH.C₆H₄.SO₂NH₂; the latter, [183°], is insol. water, and is split up by HClAq into C₀H₄(SO₂NH₂)Cl, N₂, and NH₂,C₀H₄,SO₂NH₂.

p-Amido-bendene sulphonic acid C₈H₄(NH₂)SO₃H [1:4]. Sulphanilic acid. S. 6 at 6°.

Founation .- 1. By heating oxanilide or aniline with H2SO, (Gerhardt, J. Ph. [3] 10, 5).— 2. By heating aniline with fuming H.SO, at 190° (Buckton a. Hofmann, C. J. 9, 259; R. Schmitt, A. 120, 129) .- 3. From aniline and p-phenol sulphonio acid (Pratesi, B. 4, 970; Kopp, B. 4, 978).—4. By reducing p-nitro-benzens sulphonio acid.—5. By hoating aniline ethyl-sulphate (Limpricht, A. 177, 80).

Preparation.—Anilino (93 g.) is slowly poured into H₂SO₄ (50 g.) diluted with water. The

solution is evaporated and the dried sulphate is mixed with H2SO4 (50 g.) and sand, and heated in a dish, with constant stirring, until it becomes solid. Crystallised from water.

Properties.—Plates or trimetrio prisms (with

aq); monoclinic (with 2 aq).

Practions.-1. Bromine-water gives tri-bromo. aniline and (1,3,2,5)-di-bromo-amido-benzene sulphonic acid.—2. Oxidised to quinone by K.Cr.O., and H.SO.Aq (Meyer a. Ador, A. 159, 7) or by MnO₂ and H₂SO₄ (Schrader, B. 8, 759).-8. KMnO, converts its potassinm salt into the azo derivative C₆H₄(SO₃K).N₂.C₆H₄.SO₃K (Laar, J. pr. [2] 20, 264), the corresponding azexycompound being also formed (Limpricht, B. 18, 1420).—4. PCl, forms C₆H₄(SO₂Cl).NH.POCl₂ [158°], which is converted by alcohol into C₆H₄(SO₂Et).NH.PO(OEt)₂ [102°], and by mothyl alcohol into C₆H₄(SO₃Mc).NH.PO(OMe)₂ [114°]. The former is split up by boiling into alcohol, sulphanilic acid, and hydro-di-cthylic phosphate. V. also Di-bromo-amido-benzene sulphonio aoid.

Salts.—NaA' 2aq. — KA' 1\frac{1}{3}aq: triclinic prisms.—NH₄A' 1\frac{1}{3}aq.—BaA'₂ 8\frac{1}{3}aq.—CuA'₂ 4aq.—Aniline sulphanilate C₂H₁N2HA'. Gives off

all its aniline at 100°.

Acetyl derivative CaH4(NHAe)(SO3H)obtained as the sodium salt by boiling sodium sulphanilato with acetic anhydride. The free acid has not been isolated, as it readily splits off acetic acid on evaporation of its solution. The sodium sait (A'Na) forms small colourless prisms very soluble in water, but less in alcohol (Nietzki a. Benekiser, B. 17, 707).

Amido-benzene di-snlphonic acids.

I. C₆H₇NS₂O₆, i.e. C₆H₃(NH₂)(SO₃H)₂. [1:3:4?] From mamido-benzene sulphonio acid and fuming H₂SO₄ at 180° (Drebes, B. 9, 552; Zander, A. 198, 21).—Rhombic octahedra, v. e. sol, water or alcohol.

 $\begin{array}{l} {\rm Salts.-(NH_4)_2A''aq.-K_2A''aq.-KHA''.-} \\ {\rm BaA''1!aq.-BaH_2A''_2; S. 2.9 \ at 8^\circ.-PbA''aq.-} \\ {\rm PbH_2A''_2.} \end{array}$

II. C₆H₃(NH₂)(SO₃H)₂2aq. [1:3:5]. From the corresponding nitro acid by reduction (Heinzelmann, A. 188, 167). Four or six sided columns, v. sol. water, and alcohol, insol. ether.

Bromine water gives a pp. of bromanil.

Salts. — (NH₁)₂A" aq. — HNH₁A" xaq. —

K₂A" 3aq. — K₂A" 4aq. — KHA" aq. — BaA" 3½aq.

— BaH₂A"₂ 5aq. — PbA" 3½aq. — PbH₂A"₂ 6aq. —

Ag2A".

III. C_sH₂(NH₂)(SO₅H)₂ [1:2:4]. anilic acid.

Formation.-1. By heating sulphanilic acid with fuming H₂SO, at 170° for 6 hours (Buckton a. Hofmann, A. 100, 164).—2. By heating o-amido-benzene sulphonic acid with fuming H₂SO₄ at 180° (Zander, A. 198, 17).—3. By reducing the corresponding nitro acid (Heinzelmann, A. 188, 170).

Properties. - Minute (red) clumps (from water). V. sol. alcohol, insol. ether. Bromine water gives tri-bromo-aniline, bromo-amido-benzene di-sulphonic acid, and di-bromo-amido-benzene sulphonio acid.

Salts.—The acid salts are less soluble in water than the neutralonos .- (NH,) A" aq: small heragonal prisms.—NH,HA"2aq: clumps.— K₂A"aq.—KHA"aq: silky nesdles.—BaA"3aq: four-sided plates.—BaH₂A"₂aq.—CaA"2aq: minute white needles.—CaH₂A"₂: slender needles.-PbA"2aq.-PbH2A"2aq: small prisms. - PbH₂A"₂6aq. — Ag₂A": prisms. — AgHA": needles or plates.

Di-amido-benzene sulphonic acids C.H.N.SO. Phenylene-di-amine sulphonic acids.

I. $C_6H_3(NH_2)_2(SO_3H)$ [1:2 or 5:3]. S. 1 at 10°. From the corresponding nitro acid (Sachse, A. 188, 148).—Rhombic tablets. V.sl.sol. alcohol, insol. ether. Turns brown in air. Metallic salts crystallise with difficulty.

Salts.-HA'HCl: needles.-HA'HClSnCl2. -HA'HBr.-H2A'2H2SO4 aq.-HA'H2SO4 2aq.

II. $C_6H_3(NH_2)_2HSO_8$ [1:2:4]. Small colourless needles.

Preparation .- 1 By sulphonation of o-phenyleno-diaminc.-2. By reduction of (1:2:4) nitro-

amido-benzene sulphonic acid.

Salts. - A', Ba + 5 H.O: easily soluble thin tables or needles.— A'_{2} Ca + $3H_{2}$ O: soluble tables or needles (Post and Hardtung, B. 13, 39; A. 205, 98).

III. C₆H₃(NH₂)₂HSO₃ [1:3:4].

Preparation.—1. By sulphonation of m-phenylene-diamine.—2. By reduction of nitroamido-benzeno sulphonic acid [1:3:4].

Dimorphous: monoclinic tables, a:b:c= 1.31:1:1.36, or triclinio prisms, a:b:c=·424 : 1 : ·928.

Salts.-A'2Ba6aq: long soluble prisms. A'₂Ca 5aq: soluble prisms or tables (Post a. Hardtung, B. 13, 40; A. 205, 104).

Di-amido-benzene di-sulphonic acid C.H.N.32.O. aq i.e. C.H.2(NH.2).2(SO.H.).2 aq. From the nitro acid by reduction (Limpricht, B. 8, 290). V. sol. water.

Salt.—SnA" aq: needles.
AMIDO-BENZENE PHOSPHONIC ACID

C₆H₈NPO₃ i.c. C₆H₄(NH₂)PO(OH)₂. From the nitro aoid, tin, and HCl. Slonder needles (from water); v. sl. sol. water, v. sol. HClAq; insol. alcohol and other. Salts. -Na, A" 3aq. -PbA' CuA".-Ag2A" (Michaelis a. Benzinger, A. 188, 282)

DI-AMIDO-BENZHYDROL v. DI-AMIDO-DI-PHENYL-CARBINOL.

AMIDO-BENZOIC ACIDS C,H,NO, C.H. (NH.).CO2H. M. w. 137. The following derivatives are described in special articles: NITRO-AMIDO-BENZOIO ACID, CHLORO-AMIDO-BENZOIC AOID, CHLORO - METRYL - AMIDO - BENZOIC ACID, METHYL - AMIDO - BENZOIO ACID, ETRYL - AMIDO-BENZOIO ACID, PHENYL-AMIDO-BENZOIO ACID.

o-Amido-benzoic acid C.11 (NH2).CO2H [1:2].

Anthranilic acid. [144°-145°].
Formation.—1. By reducing o-nitro-benzoio acid (Beilstein a. Kuhlberg, A. 163, 138).-2. By boiling indigo with KOHAq (Fritzsche, A. 39, 83).—3. From (1, 2, 3), or (1, 4, 5)-bromo-amido-bonzoic acid by sodium-amalgam (Hübner a. Petermann, A. 149, 133).—4. From its acetyl derivative and boiling cono. HCl.—5. From isatoio acid (q. v.) and boiling cono. HCl.

Properties.—Plates, or rhombio orystals (Haushofer, A. 193, 233). May be sublimed.

V. sol. water, and alcohol. Converted by nitrous acid into salicylic acid; and by sodium-amalgam into NH, and benzoio soid. HCl and KClO. form ohloranii (Hofmann, A. 52, 65). Its anhydride is described as ANTHRANIL. Salts.—HA'HCl: f. jl°]; needles (Kubel,

parts.—HA HOI: , 31]; needles (Rubel, A. 102, 236).— HA'HNO₃.— (HA'),H₂SO₂aq: needles [188°].—(HA'),H₂SO₁aq.—(HA')₂H₂CO₄. —BaA'₂: v. e. ε.λ. v. t. t. s. s. sol. alcohol.— PbA'₂.—CuA'₂.—AgA'. Ethyl ether Eth'. (260°). Liquid; its

Ethyl ether EtA'. (260°). Liquid; its hydrochloride, EtA'HCl, [170°], forms needles, insol, other, and may be sublimed.

Reactions .- 1. Nitrous acid produces salicylic acid (Gerland, A. 86, 143) or diazobenzoic acid (v. Di-Azo compounds). -2. KCNO converts the hydrochloride of o-amido-benzoio acid into uramido-benzoio acid (q. v.); Potassium sulphocyanide forms, similarly, thio-uramido-benzoio acid (q. v.).—3. Phenyl cyanate (q. v.) forms NH_Coll. CO.NPh.CO.NPhH.—4. Cyanogen passed into an aqueous solution forms C₂H₂N₂O (Griess, B. 11, 1986), while in an alooholio solution it forms C10H10N2O2, [173°] (Grioss, B. 2, 415). The latter is converted by boiling HCl into C₈H₈N₂O₂, [above 350°], which is probably NILCO sinco it can be formed by heating

O.NH. o-amido-benzoio acid with urea. It forms crystalline nitro- and amido-derivatives.

The compound C10H10N2O2 'ethoxyl cyanamidobenzoyl' is converted by alcoholic NH, at 100° into benzereatinine (q. v.).

The compound C₀H₂N₃O 'di-cyano-amido-

benzoyl,' may be represented thus:

CO.N C.H. NH.C.CN (Griess, B. 18, 2417). This body gives the following reactions .- a. Strong

NH3Aq converts it into Coll NII.C.CO.NH2 'carboxamido-cyano-amido-beuzoyl.'-b. Aqueous ammonium sulphide forms the corresponding

CO.N C.H. -c. Baryta water forms CO.N

C_eH, NII.C.CO₂H, 'carboxy-oyano-amido-benzoyl'; which is converted by dry distillation CO.N

into CoH NH.CH 'carbimido-amido-benzoyl.' -d. m-amido-benzoio acid produces the anhy-

dride of di-phonyl-guanidine dicarboxylio aoid, CO.N C₆H NH.C.NH.C₈H CO₂H. -e. p-plicnylene-

diamine produces the anhydride of amide-diphenyl-guanidine carboxylio acid CO.N

All these bodies may NH.C.NH.C.H.NH.

be looked upon as derivatives of CoH. which may be called Quinazoline.

Formul derivative C.H. (NHCHO)CO.H 1 aq. [168°]. Formed by heating isatoic or c-amide-

bensoic said with formic said (E. v. Meyer s. Bellmann, J. pr. [2] 88, 24). Hair-like needles; sol. alcohol, sl. eol. benzene.

Acetyl derivative CaH4(NHAc)CO2H. [180°]. Formed by boiling anthranil (q. v.) with Ao.O and treating the pladuct with water [Friedländer a. Henriques, B. 15, 2105). Also by oxidation of (Py. 3) methyl quinoline by KMnO. (Doebner a. Miller, Be 15, 30 35).

Preparation. — Acetyl-o-toludine (1 g.) is oxidised by KMnO₄ (2 g.) dissolved in water (200 c.c.), the liquid being kept neutral by acetic acid (Bedson a. King, C. J. 37, 752).

Properties.—Linstrous leaslets (from water). Prisms (from HOAo). Trimetric, a:b:c= 982:1:2:803 (Fletcher).

Salts .- PbA'2: flocculent pp.-AgA': needles. Tests .- Solution of sodium salt gives with lead acetate a pp. sol. in acetic acid, with CaCl, a pp. only on adding alcohol.

Di-acetyl derivative C.H.(NAc.)CO2H.

[220°].—AgA'.

Chloro-acetyl derivative C.H.(NH.CO.CH.Cl).CO.H. From acetyl derivative and PCl, (Jackson, B. 14, 888). Clumps. Di-chloro-acetyl derivative

C,H4(NH.CO.CHCl2).CO2H. [c. 178°]. Prepared like the preceding. Yellowish needles (from water).

Salt.-AgA'.

Bensoyl derivative C.H. (NHBz).CO.H. [182°]. By BzCl; or by acting on benzoyl-otoluidine with KMnO,Aq (Brückner, A. 205, 130). Long needles (from alcohol); insol. water.

Salts.—NaA' 4aq.—MgA'2 4aq.—CaA'2 3aq. -BaA'2 Baq.

Oxaloxyl derivative v. CARBOXY-PHENYL-OXAMIO ACID.

Amide C₆H₄(NH₂)CO.NH₂. Amido-benzamide. [108°]. (300°). From NH₃ and isatoic acid (q.v.). White plates (from chloroform). Sl. sol. benzene and ether. Aqueous solution of (1 mol.) of its hydrochloride gives with NaNO₂ needles of C₇H₃N₃O, [213°]. This compound forms salts, e.g. C,H,NaN₃O, and a methyl ether, C,H₄MeN₃O. The methyl ether, [123°], is also C.H. MeN.O. The methyl ether, [123°], is also formed from the methylamide of o-amido-benzoic acid. The new substance is probably

CO.NH (Weddige a. Finger, J. pr. [2] 35, 'N = N

Aoetyl-amido-benzamide

C, H, (NHAo) CONH2. [171°]. By Ao2O. Needles. Forms salts with acids. If kept melted for some time it becomes solid, changing to the anhydro-

NH.C.CH, Oxy-methyl-quinsompound, C.H. CO.N

asoline [228°]. Yellow silky needles (from alcohol). Soluble in hot water. Forms ealts with

Förmyl-amido-benzamide C.H.(NH.CHO)CONH2. [123°]. From dry formio acid and o-amido-benzamide. When heated this gives H2O and an anhydro-compound

NH.C.H C.H. CO.N Oxy-quinazoline (A. Weddige, J. pr. [2] 31, 124).

Anviide O.H. (NH.) CO.NHPh. [1800]. Front aniline and isatoic soid. Needles (from bensens).

Phenyl-hydraside
C.H. (NH.) CO.NPh.NH. [170°]. From isatols
acid and phenyl hydrazine in alcoholic solution
at 70°. Yell w needles; col. alcohol and chloroform, v. sl. sol. ether (E. v. Meyer a. Bellmann,

J. pr. [2] 33, 21). HydroxylamideC,H,(NH,)CO.NH.OH. [82°]. From isatoic acid (q.v.) and hydroxylamine solution (M. a. B.). Yellowish plates, sol.

aloohol, ether, and chloroform.

o-Oxy-phenylether C₆H₄(NH₂).CO₂.C₅H₄OH. [136°]. From isatoic acid and pyrocatechin at 130° (M. a. B.). Needlee (from water). Sol. alcohol and ether, sl. eol. water.

m-Amide-bsnzeic acid CaH4(NH2).CO4H [1:8]. [173°-174°]. S.G. 1.51 at 4°; S. 2 in cold water,

4 in boiling water or alcohol.

Formation.—1. From m-nitro-benzoio acid by reduction (Zinin, J. pr. 36, 103; Gerland, A. 86, 143; 91, 185; Schiff, A. 101, 94; Beilstein a. Wilbrand, A. 128, 265). -2. From nitrophthalic acid, Sn, and HCl (Faust, Z. [2] 5, 335).—3. From C₈H₃Br(NH₂)CO₂H [4:3:1] by sodium-amalgam (Raveill, A. 222, 180).

Properties .- Crystalline clumps; sweet taste; may be sublimed. Aqueous solutions are browned

by air.

Reactions.-1. HCl and KClO, form chloranil. Bromine forms tri-bromo-amido-benzoie acid.

 Nitrous acid forms m-di-azo-benzoic acid. 4. The solution containing m-diazo-benzoio acid gives m-oxy-benzoio acid on heating

5. Fusion with urea produces uramido-ben-

zoic acid (q. v.). 6. Boiling with CS2 and alcohol produces thio-carbonyl-di-amido-di-benzoic acid, $CS(NH.C_0H_4.CO_2H)_2$ (Merz a. Weith, B. 3, 812). This body is also formed by heating m-amido-benzoic acid with thio-urca; and, together with thio-carbimido-benzoic aoid, by heating m - amido - benzoic acid with CSCl, (Rathke a. Schäfer, A. 169, 101). Thio-carbonyldi-amido-di-benzoic acid does not melt below 300°; it is v. sl. sol. water, m. sol. alcohol or ether; converted by HgO, in presence of KOH (Griess, A. 172, 169), into carbonyl-di-amido-di-

benzoic acid. 7. CSCl, produces the last mentioned body and also thio - oarbimido - benzoic acid SCN.C.H..CO2H. This may also be prepared by boiling thic - oarbonyl - di - amido - benzoio acid with HCl. It is an amorphous insoluble powder, v. sl. sol. alcohol. Decomposes above 310°. It unites with aniline forming di-phenyl-thio-urea carboxylio acid, PhNH.CS.NH.C.H.CO.H [191°]. This body is PhNH.CS.NH.C,H,.CO,H [191°]. also formed by heating phonyl-thio-carbimide with m-amido-benzoio acid at 100' (Merz a. Weith, B. 3, 244). It forms slender needles (from water). V. sol. alcohol, and ether, sl. sol. benzene, and benzoline. AgNO, added to its alkaline solution gives a black pp. of Ag.S; Fe₂Cl₆Aq gives a yellow pp.; Pb(OAc)₂ a white pp.; and CuSO₄ a green pp. (Aschan, B. 17, 430)₆

8. Phosgene produces carbonyl-diamido-di-henzoio aoid CO(NH.C.H., CO₂H)₂ (Sarauw, B. 15, 44). This body is also formed by heat-

ing m-uramido-benzole acid (q: v.) at 200°. (Tranbe, B. 15, 2124). White powder; insol. water, alcohol, and benzene, sol. alkalis.—BaA"8aq.—Ag.A".—PbA". Its ether Et.A", [161°-162°], is formed by heating m-uramidobensoic ether (Griess, J. pr. [2] 4, 294): needles (from dilute alcohol).

9. Phenyl cyanate forms on heatifg, phenyl nramido benzoic acid, NPhH.CO.NH.C.H. CO.H. [270°]. Concentric priams, sol. alcohol, sl. sol. ethor, insol. water

(Kühn, B. 17, 2882).

10. Aqueons KCNO evaporated with m-amidobenzoic acid forms thio-uramido-benzoic acid NH2.CS.NH.C.H4.CO2H (Arzruni, B. 4, 406).

11. Phenyl cyanato (q.v.) forms di-phenylurea carboxylic acid CO₂H.C₆H₄.NH.CO.NPhH.

12. PCl, converts amido-benzoio acid into a white powder which, when extracted with water, yields a solution greatly resembling solutions of albuminous substances. Thus if a little lime-water, NaCl, or MgSO4, be added the liquid may be coagulated by heat, more ospecially if CO, be passed through the solution before heat is applied (Grimaux, C.R. 98, 231, 1336).

13. Cyanogen gas passed into an aqueous solution forms a dicyanide of amido-benzoic acid, and cyan-carbimid-amido-benzoic acid. Cyanogen passed into an alcoholic solution forms the dicyanide, guanido-di-benzoic acid, and ethoxy-

carbimid-amido-benzoio acid.

The di-eyanide (CN), NH2, C6H4, CO2H, is a yellow, crystalline powder, insol. water, v. sl. sol. alcohol or other. It does not form metallic salts (Griess, B. 11, 1985; Griess a. Leibius, A. 113, 332). On distillation it forms m-amidobenzonitrile (Griess, B. 1, 191; Hofmann, B. 1, 194). Boiling KOHAq or HCl converts it into benz-creatino (q. v.) (Griess, B. 3, 703).

Cyano-carbimidamido-benzoic acid CO2H.CeH4.NH.C(NH).CN forms elliptic plates, v. sl. sol. cold water, sol. acids and alkalis. 1t reacts as follows :- a. Nitrous acid converts it into cyano - carboxamido - benzoic acid, CO.H.O.H. NH.CO.CN. (Griess, B. 18, 2415), which forms white plates, insol. cold water, with sweetish taste; boiling water converts it into HCN, CO, and carboxy-amido-benzoic acid; dilute NH, Aq forms uramido-benzoio acid.—b. Cold dilute HCl forms small prisms of carbox amido-carbimidamido-bonzoio acid CO2H.O6H4NH.C(NH).CO.NH2; v. sol. hot water; its aurochlorido, HA'HAuCl, 12 aq, crystallisos in needles (G.).—c. Cold aqueous di-methyl-amineforms CO₂H.C₆H.,NH.C(NH).C(NH).NMo₂; six-sided plates, v. sol. hot water, sl. sol. cold water, converted by hot Na₂CO₃ aq into mcarboxy-phenyl-oxamido NH, and NMe2H.

Guanido-di-benzoic acid NH:C(NH.C.H.CO.H)2 is also formed from thiocarbonyl-di-amido-di-benzoic acid, IIgO, and NH, (Griess, A. 172, 172). It is crystalline. Salts.—BaA".—H.A"IICI.—(H.A"HCl), PtCl.

Ethoxy - carbinidamido - benzoic acid, EtO.C(NH).NH.C.H.CO.H 12aq forms needles (from water), sol. alcohol, and ether; sonverted by alkalis into alcohol and uramidobenzoic acid; nitrous acid convorts it into

Carboxy-amido-benzoic acid monoethyl ether CO.Et.NH.C.H..CO.H [189°]. This soid is also formed from amido-benzoic

soid and Cl.CO₂Es. It orystallises in plates (from water). Salts.—BaA'₂2aq.—AgA' Griess, B. 9, 796; Wachendorff, B. 11, 701).

Circles, B. 9, 170; Washendoon, J. A., 171; Wallet, CleH., NO., NH., [158°] (W.).

Salts.—HA'HCl: pisms (Cahours, A. Ch., [3] 53, 322).—(HA'HCl) PtCl.,—HA'HClSnCl.—HA'HBr.—HA'HNO.—(HA').H., SO., aq. [225°]. HA'H,PO. (Harbert, A. 123, 290).—BaA', 4aq. —CaA', 3aq. —ZuA', 2—PbA', Nocdles.—MgA', 7aq.—AgA'—NaA' (at 100°) (Voit, A. 99, 100).—SrA', 2aq.—ZnA', (when dried at 100°).

Methyl ether.—McA'. Oil (Chancel, C. R.

30, 751).

Ethylether.-EtA'. (294°). Liquid, sol. water; from m-nitro-benzoio ethor. Salts: Eth'HCl. [185°]. (Müller, B. 19, 1493).—
(EtA'HCl)_PtCl.—EtA'HNO3: prisms.

Acetyl derivative C. H. (NHAO).CO.H.

Acetyl derivative C.H. (NHAc).CO.H. 5°]. Formation.—1. By Acol or by HOAc at 140° (G. C. Foster, C. J. 13, 235).—2. From amido-benzoic acid (10 g.) and acctic other (25 c.c.) at 150°. (Pollizzari, A. 232, 148).-3. From amido-benzoic acid and acctamide (P.), or Ac20 (Kaiser, B. 18, 2946). Properties .- White powder, v. sol. hot alcohol, sl. sol. hot water, v. sl. sol. cold water and other. Dissolves in Na, HPO, Aq, but re-ppd. by HOAc. May be sublimed. Salts.
—BaΛ', 3aq: nccdles.—CaΛ', 3aq.—NaA'(at 120°).

Formyl derivative C4H4(NH.CHO).CO.H. [225°] (Pellizzari, G. 15,

555).

Heptoyl derivative C₆H₄(NH.C,H₁₃O).CO₂H. [202] (P.)

Glyeollyl derivative C₆H₄(NII,CO.CH₂OH).CO₂H. [212°]. Gives at 220° the anhydride $\langle {}^{\mathrm{CH_2}}_{\mathrm{CO}} \rangle$ N.C₆H₄.CO₂H₇ [248°].

Acetyl dorivative. [198°]. Lactyl derivative

CH, CH(OH).CO.NH.C, H, CO.H. [162°].

anhydride melts at [243°] (P.).

Benzoyl-derivative C.H. (NHBz)CO.H. [248°]. 1. From amido benzoic acid and benzamide at 180° for 2 hours .- 2. By boiling amidobenzoic acid (2 g.) with benzoic ether (4 c.c.) for 6 honrs (Pellizzari, A.232, 150) .- 3. From amidobenzoic acid and benzanilide at 230° (P.). Minute prisms (from alcohol). Soluble with ease in alcohol, less so in ether or water. Its Ca and Ba salts are soluble. Rosolved by hot KOH into benzoio and amido-benzoio acids. Amida.

—C_aH₄(NHBz)CONPhH. [225°]. By heating C_cH₄(NBzH)CO₂H with aniline for some honrs.

Oxaloxyl derivative v. CARBOXY-PHENYL-

OXAMIO ACID.

Succinyl-derivative CuH, NO, or $CH_2 \cdot C = N \cdot C_6 H_4 \cdot CO_2 H$. [235°]. Formed by • (?)-

CH.. CO.Ò melting succinoxyl-amido-benzoic aoid, or by fusing succinic and amido-benzoic acids together (Muretow, J. R. 4, 295; Pollizzari, B. 18, 215). Necdles (from alcohol). Sl. sol. cold water.

Salts.—Bah', 2aq.—Agh'.
Succinoxyl derivative C₁H₁₁NO₁ i.e.
CO₂H.CH₂CH₂CO₂H.C₄H₆CO₂H. [230°]. From the preceding by boiling with water, baryta or ammonia (M.). Plates; m. sol. water. ammonia (M.). BaA" 1 aq.

Succinul-di-amido-di-bensoic acid

C₁₁H₁₆N₂O₂ i.e. C₂H₄(CO.NH.O₄H₄.CO₂H)₂. (?) (e. 300°]. Formed together with CO₂Et.CH₂.CH₂.CO.NH.C₄H₄.CO₂H by heating i.s. O.H.(OO.NH.O.H.CO.H), (?) alcoholic succinic ether with amido-benzoic acid Aleo from sucoinyl-amido-benzoic (M.; P.). acid, alcohol, and HCl (1.). White crystalline powder. Soluble in KONAq. Salte: CaA"7aq. S. 2.—BaA"5aq: needles:

Phthalyl denie of the C₁₃H₂NO, i.e. C₆H₄:C₂O₃:N.C₆H₄:CO₄H. [282]. Formed togsther with its ether by heating amido-benzoio acid with phthalic ether (Pellizzari, B. 18, 216). Ethyl ether: A'Et. [152°]: radiating needles.

Sebacyl derivative C, H, N,O, i.e. C_sH_{1s}(CO.NH.C_sH_sCO.H)_s. [275°]. Formed together with CO.Et.C_sH_{1s}CO.NH.C_sH_sCO.H by hsating schacio ether with amido-benzoic acid in alcoholic solution (P.). White powder, el. eol. in most menstrua.

Amide C₆H₄(NH₂).CO.NH₂aq. Amido-benzamide. [75°]; when dry [above 100°]. From m-nitrobenzamide by ammonium sulphide (II. Sohiff, A. 218, 185; Chancel, A. 72, 274).

Properties .- Large, yellow crystals. Unites with acids forming compounds :- C,H,N,OHCl: needles. -C,H,N,OHNO, -(C,H,N,OHCl),PtCl, -C,H,N,OAgNO3: needlcs.

Reactions.-1. Aqueous solutions (even when very dilute) give with fatty aldehydes crystalline ppe. of the form R.CH(NH.C_sH₄.CO.NH₂)₃. These are soluble in alcohol, but give with HNO, containing CrO, a violet colour. They differ from original amido-benzamides in being no longer basic and in giving no coloured coinpounds with furfurol solution.—2. Aqueous salicylic aldehyde give yellowish needlee of C₆H₄(OH).CH:N.C₆H₄NH₂. [186°]. V. c. sol. alcohol or warm water. Thie compound, o-oxybenzylidene-amido-bonzamide, boiled with benzoio aldehyde forme a product C_{ss}H_{2s}N₄O₃, insoluble in water, alcohol, ether, toluene, chloroform or CS2, but may be crystallised from phenol (2 vols.) and alcohol (1 vol.). It may be considered to be an anhydride of

C.H. CH(NH.CO.C.H.N:CH.C.H.OH)2.

Boiled with Ac₂O it takes up 1 molecule of Ac₂O forming small needles. Dilute aqueous NH3 reproduces the compound $C_{33}H_{2a}N_{*}O_{3}$. — 3. Helicin (2 pts.), m-amido-benzamide (1 pt.), and water (10 pts.) form yellowish plates of a gluco-side of o-oxy-benzylidene-amido-benzamide [113°]: NH2.CO.C6H4.N:CH.C6H4.O.C6H4O. 2aq. 4. Boiled with an alcoholio solution of isatin it forme a crystalline powder [c. 280°] of isatamido-benzamide:

NH CON. C. H. CO.NH2. The compounds NHq.CO.C.H.N:X derived from aldchydee and amido-benzamide are decomposed by heating with aniline into amido-benzamide and PhN:X (Schiff, G. 13, 113; A. 218, 185).

Phthalyl-amido-benzamide

NH, CO.C.H. N:C.C.H.

f240°-241°1. Got by ó.co

fnsing m-amido-benzamide with phthalic anhydride, Bundles of elender needles (from alcohol). V. el. sol. water. Heated with aniline givee phenyl-phthalimide and amido-benzamide (H. Schiff, A. 218, 194).

m-Amido-bennamidoxim C.H.(NH2).C(NH2):NOH. A cryetalline solid; formed by reduction of m-nitro-benz-amidoxim with SnCl2. Salt .- B'HCl; priems (Schöpff, B. 18, 2472).

Anilide NH2.C6H4.CO.NPhH. Amidobenzanilide. [129°]. (P.); [114°] (E. a. V.). Formed by heating m-amido-benzoic acid with aniline (Piutti, B. 16, 1321) or by reducing mnitro-benzanilide (Engler a. Volkhaueen, B. 8, 35).—Silvery scales (P.) or long needloe (from water, E. a. V.). Heated with aniline at 200° it forms two isomerides, (C,H,NO), one soluble in alcohol, [225°], called 'amido-benzoide,' and the other an insoluble powder (Piutti, G. 13, 339).
Salte: C₁₃H₁₂N₂OHCl.—(C₁₃H₁₂N₂O)₂H₂SO₄
p - Amide - benzeic acid C₅H₄(NH₂)CO₂H.

[186°-187°]. Amido-dracylic acid.

Formation.—1. By reducing p-nitro-benzoio acid (G. Fiecher, A. 127, 142; Wilbrand a. Beilstein, A. 128, 264).—2. By boiling ite euccinoxyl-derivative with HCl (Michael, B. 10,

Preparation.—50 pts. of acetyl-p-toluidine ie suspended in about 2000 pts. of boiling water and oxidized by clowly adding 200 pts. of finely powdered KMnO. The colution is filtered from MnO2, the acctamido-benzoio acid ppd. by HCl, and saponified by beiling for an hour with etrong HCl (Kaiser, B. 18, 2942).

Properties. - Long white needles, not coloured by moist air. When heated with urea it forms CO(NH.C.H.CO.H)₂ (Griese, J. pr. [2] 5, 370).

Salts.-BaA 2: chining lamine, sol. water. -A'.Pb.OAc, ppd. by Ph(OAc), Aq (Ladenburg, B. 6, 130). — (HA'), H.SO,. — The copper-satt is a dark-green pp. (Geitner a. Beiletein, A. 139, 1).

Acetyl derivative C.H.(NHAc).CO.H. [250']. Formed from acctyl-p-toluidine by KMnO. (Hofmann, B. 9, 1302). Needlee, el. sol. water. Salt: AgA'.

Benzoyl derivative C.H.(NHBz)CO2H. [278°]. From benzoyl-p-toluidine, CrO3, and HOAc (Brückner, A. 205, 127).—Small needles (from alcohol). Salts. -BaA'2. -CaA'2.

Succinoxyl-derivative CO2H.CH2.CH2.CO.NH.C8H4.CO2H. [226°]. From p-tolyl-succinimide and dilute KMnO, (Michael, B. 8, 577). Yellewish needles, sl. sol. cold water. Salt. -C, H, NO, Ag.

Amide C₆H₁(NH₂).CO.NH₂. [179°]. Formed by reducing p-nitro-benzamide (Beilstein a. Reichenbach, A. 132, 144). Yellow cryetals, sl. sol, water.

Di-amido-benzeic acids C₇H₈N₂O₂ (Gricss, A. 154, 325; B. 2, 47, 434; 5, 192; 7, 1227; 77, 603; Pr. 20, 168; Wurster a. Ambihl, B. 7, 213; V. Meyer a. Wurster, B. 5, 635; A. 171, 62). These acids can be formed by reducing the corresponding di-nitro-, or nitroamido-, benzoic acide. They are soluble in water, combine both with acids and baeee, and eplit up, when dietilled with baryta, into CO2 and plienylene-diamine.

Nitrous acid converts the (a) acid into amido. di-azo-benzoic aoid, C₆H₃(NH₂)< CO N₂>O; the (β) and (γ) acids are converted by it into azimido benzoio acide, HN₃:C₆H₂.CO₂H, while the symmetrical acid becomes tri-amido-azo-bensoic acid O.H.(NH2)(CO2H).N2.C.H2(NH2)2CO2H v. Azo compounds.

s-di-amido-benzoic acid C.H.(CO.H)(NH.), aq [1:3:5]. [228°]. S. 1·1 at 8° (Voit, A. 99, 106; Hühner, A. 222, 85). Colourless needles, neutral to litmus; loses aq at 110°. Very dilute solutions are turned yellow by HNO.

Salts.—IIA'2HCl: needles.—IIA'H₂SO 5. 1'05 at 11',—BaA', 1\frac{1}{2}aq.—AgA'2aq.

Amide C₆II₄(CONH₂)(NH₂)₂ di-amido-benz-amide. [177'] (V.); [183°] (M.). Needles (Miretow, Z. [2] 6, 642). Salt.—Ch. N. O. (M.).

C,H,N,O2HCl: silky needles. Di-acetyl-derivative C₆H₃(CONAc₂)(NH₂)₂ 2aq. [Ab Thin needles, sl. sol. cold water (M.). [Above 270°].

(a)-di-amido-benzoic acid

C₆H₃(CO₂H)(NH₂)₂. [1:2:5]. Formed also from nitro-isatoic acid by Sn and HCl (Kolbe, J. pr. [2] 30, 480). Very small prisms (from water). V. sl. sol. alcohol, ether, and boiling water (G.). The free acid turns blue in air.

Salts. — HA'H2SO4: needles, v. sl. sol. water.—HA'2HCl (K.).

(β)-di-amido-benzoic acid C₄H₃(CO₂H)(NH₂)₂ [1:3:4]. [211°] (Salkowski, A. 173, 57; Griess, B. 5, 856). Plates. Sl. sol. cold water.

Salts.--11A'H2SO4: plates; v. sl. sol. hot water. - HA'HCl 1 laq.

(γ)-di-amido-benzoic acid

 $C_8H_8(CO_2H)(NH_2)_2$ [1:2:3]. Long needles. Salt. $-(HA')_2H_2SO_4$ 1_2 aq: six-sided tables or oolumns, v. sl. sól. water ; Fe₂Cl₅ colours its solution brownish-red.

Tri-amido-bsnzoic acids $C_7H_0N_3O_2$. I. $C_6H_2(CO_2H)(NH_2)_3\frac{1}{2}aq$ [1:3:4:5]. From dinitro-amido-benzoio (or chrysanisio) (Salkowski, A. 163, 12).

Needles (from water). Sl. sol. cold water, v. sl. sol. alcohol, and ether; solution is acid. Heat splits it up into CO, and tri-amido-benzene. Its solutions give a brown pp. with Fo₂Cl_e.

Salts.—HA'2HCl: silver-grey needles.— HA'(HCl)₂SnCl₂3,aq:monoclinic.—HA'ILSO,aq: sl. sol. hot water. - HA'2HNO. - CuA'2. ZnA'2 6aq.

II. C₆II₂(CO₂H)(NII₂)₃ [1:2:3:5]. Formed by reduction of p-sulpho-benzene-azo-s-di-amido-bonzolo acid (Griess, B. 15, 2200).

Colourless crystals; v. sol. hot water; sl. sol. alcohol, insol. ether. Very readily oxidised. Salt.-HA'H.SO.; small white needles, v. sl. sol. water, insol. alcohol.

References .- Chloro-, Bromo-, Iodo-, Nitro-, and Methyl-, amido-Benzolo acids and amido-SULPHO-BENZO10 ACID.

AMIDO-BENZOIC ALDEHYDES C,H,NO o-Amido-benzoio aldehyde C_eH_{*}(NH_{*}).CHO [1:2]. [40°]. Formed by oxidising its oxim with Fe₂Cl_e (Gabriel, B. 15, 2004).

Preparation.—o-nitro-benzaldeliydo (3 g.) is digested with FeSO, (50 g.) and NH, at 100° (Friedländer a. Göhring, B. 17, 456).

Properties.—Silvery plates; volatile with steam; may be distilled. V. sol. alcohol, ether. and benzene, sl. sol. water, insol. light petro-

Salt.—B'2H2PtCl6: large yellow prisms (from MClAq).

Reactions.-Very stable towards alkalis, but condensed by acids to $C_{14}H_{12}N_{2}O$, which is probably $C_{6}H_{4}(NH_{4}).CH:N.C_{6}H_{4}.CHO$; this forms small colourless needles [189°], is not volatile with steam, and possesses only weak basic pro-

perties; NH₂Aq, cono. HOlAq, and hot dilute HClAq reconvert it intramido-benzaldehyde.

Acetyl derivative C₂H₄(NHAc).CHO.

[7°] White needys, [Friedländer, B. 15, 2572).

Oxim C₃H₄C²A₃(Vi..NOII. [133°]. Formed by reducing 6-nitro-benzaldoxim (Cabriol, B. 14, 2338; 15, 3657; 16, 517). Nachlag, may 14, 2338; 15, 3057; 16, 517). Needles; may be sublimed. Sol. alcohol and ether, sl. sol. water and benzeno. Its methyl derivative C.H.(NH2)CH:NOMo melts at [58°] (R. Meyer, C. C. 1885, 516). Its acetyl-methyl derivative C₆H₄(NHAc)CH:NOMe [109°], and its diacetyl derivative C₆H₄(NHAc)CH:NOAo [128°] are crystalline, insel. acids and alkalis.

m-Amido-benzoic aldehyde.

Oxim C.H.(NH.).CH:NOH. [88°]. Formed by reducing m-nitro-benzaldoxim with FeSO, and NH, (Gabriel, B. 16, 1997). White felted needles. Sol. alcohol, ether, and hot benzenc, sl. sol. cold benzene, and henzoline. Dissolves in acids and alkalis. Salt: B'2112PtCla; orange yellow tables.

p-Amido-benzoio aldehyde C_vH₄(NH₂).CHO [1:4]. [71°]. Formed by action of acids on its oxim. Flat plates, sol. water. With acids it forms red salts.

Acetyl derivative C.H.(NHAc).CHO. [155°]. Long white needles.

Oxim. - C.II.(NH.,).CH:NOH. Formed by reducing the oxim of p-nitro-benzoio aldehyde by ammonium sulphide (Gabriel a. Herzberg, B. 16, 2000). Flat yellow crystals, sol. water, alcohol, ether, soids, and alkalis. Its acid solution is resolved, even in the cold, into hydroxylamine and p-amido-benzoic-aldehyde. Acetyl derivative C₆H₄(NHΛc).CH:NOH. [206°]. White plates.

AMIDO-BENZONITRILES C.H. N2

o-Amido-benzonitrils N11, C, H, CN [1:2]. [103°]. By reduction of o-nitro-benzonitrile (Baerthlein, B. 10, 1714). Needles; v. sol. water, alcohol, and ether.

m-Amido-bsnzonitrile NH ... C. H. CN [1:3]. [52°]; (290°).

Formation.—1. By reducing m-nitro-benzonitrile (Hofmann, Z. [2] 4, 726; Fricke, B. 7, 1321).—2. By distilling the dicyanide of m-amido-benzoic acid (v. p. 157, l. 32) (Griess, B. 1, 191).—3. By heating m-uramido-benzoic acid (q. v.) with P₂O₅ (Griess, B. 8, 861).

Properties.—Needles or prisms, sl. sol. water,

v. e. sol. alcohol.

Salts. - B'HCl. - B'2H2PtCls: fonr-sided tables. - B'AgNO, white lamine.

p-Amido-bsnzonitrile NH, C,11, CN [1:4]. [110°] (F.); [74°] (E.). Formed by reducing p-nitro-benzonitrile (Engler, A. 149, 302), or by distilling p-uramido-benzoio acid (F.).—Needles, v. sol. alcohol, ether, and boiling water.— B'HCl.—B'₂H.PtCl₂: needles. AMIDO - BENZOPHENONES Cl₃H₁₁NO

 $C_{13}H_{11}NO.$ Amido-di-phenyl-ketones.

o-Amido-benzophenone Bz.C₆H₄.NH₂ [1:2], [106°]. Formed by reducing o-nitro-benzophenone with Sn and HCl (Geigy a. Koenigs, B.

sol. dilute acids, alcohol, and ether.

m-Amido-bensophenone Bz.C.H.NH, [1:3] [89°]. From m-nitro-benzophenone and SnCl, (G. S. K.).—Yellow falted neadles, sol. alcohol, and ether, sl. sol. water -B'HCl: [187°]; long needles.

p-Amide-benzophsnone Bz.C.H., NH2 [124°]. Benzo aniline, Progred by boiling its phthalyl derivative with alcoholik KOH (Doebner, B. 13, 1011; Doebner a. Weiss, B. 14, 1886). Colourless plates, v. sol. alcohol, ether, and glacial HOAc, sl. sol. cold water. Nitrous acid converts it into p-oxy benzophenone. On fusion with ZnC!, it loses H.O forming a compound of the formula C13H,N. The latter is a very stable indifferent substance, crystallising in glistening plates [118°], and distils undecomposed at a high temperature; it is soluble in alcohol, ether, &c., eparingly in hot water, insoluble in cold.

sparingly in not water, insulated in cold.

Salts.—B'₂H₂SO₄: long sparingly soluble needlee.—B'HCl, B'₂H₂C₂O₄, and B'HNO₃ are more soluble.—(B'HCl)₂PtCl₄: yellow needles; al. sol. oold water.

Acetyl derivative C.H. CO.C.H. NHAc. [158°]. Long needles. Sol. alcohol, ether, acetic acid and benzene; insol. water.

Benzoyl derivative CaH3.CO.CaH4.NIIBz. [152°]. Plates, sol. hot alcohol, sl. sol. cold alcohol, insol. water.

Phthalyl derivative C.H.100. i.c. C.H., CO.C. H., N(C.O.C.H.). [183]. Prepared by the action of BzC on phthalanil in presence of ZnCl2. Large needles or plates, insol. water, al. sol. alcohol or other.

(a)-Di-amido-benzophenone C13H5O(NH2)2 [172°]. Prepared by reducing (a)-di-nitro benzophenone, [190°] itself got from di-nitro di phenylmethane. [183°] (Staedel a. Sauer, B. 11, 1747; White needles. A. 218, 344).

B"2HCl: largo tables. - B"2HClSnCl., B"H2SO.

($\hat{\beta}$). Di-amide-benzephenone $C_{13}H_{12}N_{2}O$. [165°]. Flavine.-From di-nitro-benzophenone [149°] by reduction (Chancel a. Laurent, A. 72, 281; Prætorius, B. 11, 744).—Slender yellow needles (from water).

Salts: B"H_Sn_Cl6; plates.-B"H_PtCl6 Acetyl derivative C₁₃H₈O(NHAc)₂:

needlss, [227°]

Oxim (CaH. NH2)2C:NOH: [178°]; crystal-

Phenyl-hydrazide (C.H., NH2)2C:N2IIPh: [183°]; yellowish needles (from hot alcohol). (Münchmoyer, B. 20, 511).

(7)-Di-amido-benzophenone, $C_{13}H_{\bullet}O(NH_{\bullet})_2$ [181°]. From the di-nitro-benzophenone [190°] obtained from benzophenone (Staedel, A. 218, 849) .- Glittering tablets.

Salt .- B"2lICl.

Acetyl derivative. — C₁₂H₈O(NAoH)₂.
7°]. Tables; insol. water; v. sol. alcohol. AMIDO-BENZOYL-CARBAMIDE v. Unga.

AMIDO-BENZOYL-FORMIC ACID v. AMIDO-

PHENYL-GLYOXYLIC ACID. AMIDO-BENZOYL-GLYOXYLIC ACID v. UINIBATIC ACID.

AMIDO-BENZOYL-UREA v. Unea. DI-AMIDO-DI-BENZYL C14H18N2 i.e.

NH2.C.H4.CH2.CH2.C.H4.NH2. [1320]. From the (p) nitro-compound (q. v.) by roduction.

18; \$408). Yallow plates, or thick drystals, v. Colouriess scales (from hot water), v. st. sol. cold water, v. sol. alcohol; msy be sublimed. Salts: B"2HOl. — B"H_PtCl. — B"H_SO.

-B"H₂C₂O₄.—B"(H₂C₂O₄)₂ Saq (Fittig a. Stelling, A. 137, 262)

AMIDO BENZYL ALCOHOL C,H,NO i.c. NH2.C4H4.CH2OH. [1:2]. [82°]. Prepared by the action of zine dust and HCl upon o-nitro-benzylalcohol, o-nitro-benzoic aldehyde, or anthranil (Friedländer a. Henriques, B. 15, 2109). White needles, insol. light petroleum; slightly volatile with steam.

AMIDO BENZYL-AMINE NH₂,C₆H₄,CH₂NH₂ [1:4]. (269°). S.G. ²⁰ 1·08. Benzylene diamine. From acetyl-p-nitro-benzylamine, Sn, and HCl: the Ac being split off in the operation (Amsel a. Hofmann, B. 19, 1287).-Colourless liquid, v. sol. water and alcohol, insol. ether; alkaline, absorbing CO, from the air.

Salts.-B"2HCl: needles, v. sol. water. B"2(IICl),PtCl,: flat needles. The nitrate and oxalate orystalliso in long white needles. AgNO. forms a double salt in large plates.

Di-amido-di-benzyl-amine C,4H,7N, (NH.,C₆II,CH₂), NH. [106°]. From the nitro compound (Strakosch, B. 6, 1060). Needles or plates; may be distilled, but not volatile with

steam. Salts.—B"'3HCl.—B"'(HCl), PtCl.,
Tri-amido-tri-benzyl-amine C., H., N. (NII C.H. CH.) N. (2020) Tri-amido-tri benzyl-amine C₂₁II.,N, i.e. (NII₂,C₆H, CH₂)₃N. [136°]. From the nitro compound (S.).—Octaliedra (from alcohol); not volatile with steam. Insol. water, v. sol. hot alcohol. Reduced by Sn and HCl to p-toluidine and the preceding body.

AMIDO-BENZYL-ANILINE C13H11N2 NH., C.H., CH., NPhH. [88°]. From the nitrocompound by NH, and H,S at 100° (Strakosoh, B. 6, 1063).—Scales, v. sol. alcohol, ether, and benzene, not volatilo with steam. Salt .-B"2HCl; v. sol. water, less so in HClAq.

AMIDO-BENZYL-BENZENE v. AMIDO-DI-PHENYL-METHANE.

AMIDO-BENZYL CYANIDE v. NITRILE OF AMIDO-PHENYL-ACETIC ACID.

DI - p - AMIDO - DI - BENZYL - MALONIC -

ETHYL - ETHER (C.H.(NH2).CH2)2:C(CO2Et). Obtained by reduction of di-nitro-di-benzylmalonic ether with SnCl2.

Salts:- $(A''Et_x)H_xCl_z$: [230°], easily soluble needles.- $(A''Et_x)H_xSO_x$: scales.- $(A''Et_y)H_xCl_z$: glistening yellow scales.- $(A''Et_z)H_zCl_z$ PtCl_: reddish-brown plates (Lellmann a. Schleich, B. 20, 436).

AMIDO-BENZYL-PHENOL C₁₃H₁₃NO t.e. C₅H₅CH₂(C₄H₃(NH₂)(OH). [1:3:4]. From the nitro-compound. Scales (Rennie, C. J. 41, 221). DI.AMIDO-BENZYL-TOLUENE C₁₁H₁₆N₂. A

crystalline powder, obtained by reducing dinitro-p-benzyl-tolueno (q. v.) (Zincke, B. 5, 684). Salts. B"2HCl.—B"H, SO.

AMIDO BROMO COMPOUNDS AMIDO-COMPOUNDS.

AMIDO-BRUCINE v. BRUCINE. AMIDO BUTYL-BENZENE v. AMIDO-PHENYL-

AMIDO BUTYRIC ACIDS a-Amido-s-butyrio acid CH, CH, CH(NH,), CO, H. S. 3 at 15°; S. (alcohol) 18 at 80°. From a-bromo-butyric acid and NH, Aq (R. Schneider, A. Suppl. 2, 71). Stellate groups of small lamins or needles (from alcohol); nsutral; sweet taste; insol. ether.

Salts.-HA'HCl; v. sol. water.-HA'HNO; tsrn-like groups of silky needles .- (HA'), H,804-HOPb.A'.-AgA'.

8-Amide-n-butyric acid

- 11 255 1 .

CH₃.CH(NH₂).CH₂.CO₂H.

Amide CH3. CH(NH2). CH2. CO.NH2. An anyrphous mass, obtained by the action of alcoholic NH₃ on β-ehloro-n-butyric ether (Balbiano, G. 10, 137; B. 13, 312). Its platino-chloride crystallises in orangs tablés, sl. sol. alcohol.

a-Amide-iso-bntyric acid CMe. (NH.).CO., H. Formation.-From acctonyl-ures and furning HCl at 160° (Urccb, A. 164, 268).—2. From diacetonamine (Heintz, A. 192, 343; 198, 46).

Preparation.—The acctone cyanhydrin, obtained by the action of dilute HCN on acctone, is heated with alcoholic NH, at 60°, and the product saponified (Tiemann a. Friedländer, R. 14, 1971):

Properties.—Plates or tables, v. sol. water, sl. sol. alcohol, insol. ether; sublimes at about 220°.

Salts.-BaA', 2aq: needles.-MgA', tbick prisms.—CuA'₂: plates, giving a violet solution.
—AgA': needles, sol. water.—HA'HCl 2aq.— HA'HCl.

Nitrile.-CMe,NH, CN. The product of -the action of alcoholic NH, on acetone-cyanhydrin (vid. sup.).

AMIDO-CAMPHOR v. CAMPHOR.

AMIDO-CAMPHORIC ACID v. CAMPHORIC ACID. AMIDO-CAPROIC ACID v. LEUCINE, and

AMIDO-HEXOIO ACID. AMIDO-CAPRYL-BENZENE v. AMIDO -

PUENYL-OCTANE. AMIDO-CAPRYLIC ACID v. Amino-octore acin.

AMIDO-CARBOSTYRIL C.H., N.O i.c. $C_0H_1 < CH:CH > CO.$ [127°]. Anhudride of

hydrazido-einnamicacid; Oxy-amido-quinoline; Amido-pseudo-carbostyril. Prepared by converting diazo-einnanio acid by Na.SO, into SO,Na.N₂.C₄H₄.CH:CH.CO₂H, then reducing this substance by acetic acid and zino dust to SO₄Na.NII.NH.C₅H₄.CH:CH.CO₂H₅ boiling this with HCl and the Alice VIII.CO₂H₅ boiling this with HCl and then adding KOH (Fiselier a. Kuzel, A. 221, 278).

Properties. Siender needles; may be sublimed; sol. alcohol, ether, and hot water. It forms salts with acids.

Reactions. -1. Does not reduce alkaline copper or silver solutions .- 2. Nitrous acid con-

verts it, even in the cold, into carbostyril.
γ-Amide-carbostyril. From carbostyril by
nitration and reduction (Friedländer a. Lazarus, A. 229, 246). Yellow plates (from glacial HOAe). Does not melt below 320°.

Methyl derivative C.H.N(NIL)(OMe). [103°], amido-(Py. 3) methoxy-quinoline. Formed from the nitro-compound by SnCl, (Feer a. Koonigs, B. 18, 2397).-Silvery plates; v. sol. alcohol and ether; m. sol. warm water. Its sthereal solution has a blaish fluorescence. KMnO₄ oxidises it to methoxy-quinolinic acid [140°]. Diluts HCl at 120° forms (γ)-amido See also Oxy-amido-quinoline.

AMIDO-CHROMATE OF POTASSIUM v. Amido-chromates, under Chromium, acids or. You I.

AMIDO-CHRYSANISIC ACID v. NITBO-DE-AMIDO-BENZOIO ACID.

AMIDO-CINNAMIC ACIDS C.H.NO..

a-Amido-cinnamic acid

CeHs.CH:C(NH2).CO2H. Obtained by saponifying

its benzoyl derivative. Silvery plates, decomposing at 240°-250° (Pochl, B. 17, 1619).

Salts.—CuA'_22p'. small blue prisms.—
(IIA'),HCl: flat prisms.—S. sol. cold water and alcolud. alcohol.

Benzoyl derivative C.H. CH:C(NHBz).CO.H [131°]. Formed by heating an acetic acid solution of benzoyl-di-amido-

C.H.CH.CH(NIIBz). hydroeinnamio laetam

Needles or prisms; sol. aleohol, ether, and hot water.

NH.CO

o-Amido-cinnamic acid

NH₂,C₆H₄,CH:C11.CO₂H [1:2]. [159°].

Preparation. -- From o-uitro-ciunamic acid (150 g.), erystallised baryta (2100 g.), water (30 litres), and ferrous sulphate (1400 g.), by heating two hours at 100° (Fischer a. Kuzel, A. 221, 266; Tiemann a. Opermann, B. 13, 2061). Ammonia may be used in place of baryta (Gabriel, B. 15, 2294; Friedländer, A. 229, 241).

Properties.—Yellow needles; sol. alcobol, ether, and hot water, sl. sol. cold water. Dissolves in aqueous alkalis and acids.

Salts.—HA'HCl; prisms.—BaA'2; prisms. Ether.—EtA' (78°). May be distilled. Yellow needles, with yellowish-green fluorescence. Its hydrochloride is sparingly soluble in excesa of cone. HCl; its acetyl derivative, [137° forms white needles, which may be distilled (Friedländer a. Weinberg, B. 15, 1422).

Ethyl derivative C, II, (NHEt).CII:CH.CO, H. From the neid (60 g.), KOH (96 e.c. of 20 p.e. solution), alcohol (240 g.),

and Et! (60g.), by boiling (F. a. K.).

Reactions.—1. Long boiling with IIClAq
forms earbostyril.—2. ZnSO,Aq gives a crystalline pp.-3. AgNO, Aq gives a white pp.-4. CuSO, Aq gives a light green pp. - 5. Pb(OAe), Aq gives a yellow pp.

m-Amide-cinnamic acid

C₀11₄(NII₂).CH:CH.CO₂H [1:3]. [181°]. preparation is similar to that of the o-compound.

Properties. - Long yellow needles; sol. aleohol, ether, and hot water. Dissolves in aqueous acids and alkalis.

Reactions.-1. CuSO,Aq gives a dull green pp.-2, 3, same as above.-4. Pb(OAc),Aq givss a white pp. sol. hot water (T. a. O.)

Salts.-HA'HCl: plates.- (liA'HCl)2PtCl4 $-\mathrm{HA'HNO_3}$: slender needles. $-\mathrm{Ba}\underline{\Lambda'_2}$ 2aq: plates. p-Amido-cinnamic acid

C₈H₄(NH₂).CH:CH.CO₂H[1:4]. [176°]. Prepared by reducing p-nitro-einnamic ether in alcoholic solution with tin and HCl; yield: 75 p.a. (Miller a. Kinkelin, B. 18, 3234). Slender yellow needles, sol. water, aleoliol, and ether. _Dissolves in aqueous alkalis and acids.

Reactions. -1. CuSO, Aq a brown pp.-2, 3, and 4, the same as for the m-compound.

Salts.-HA'HCl.-(HA'HCl),PtCl.

Acetyl derivative [260°]. Long needlss, sol. hot alcohol, sl. sol. water, v. sl. sol. ether and benzene (Gabriel a. Herzberg, B. 16, 2041).

Di-amido-einnamic acid
U_H_1(NH_2)_c.CH:CH.CO_H. [168]. Formed by
reducing (8:4:1)-nitro-amido-cinnamic acid
(Gabriel a. Herzberg, B. 16, 2042). Yellow
needles, sol. hot alcohol, and water, insol. ether,
benzene, and benzoline.

AMIDO-COMENIC A"ID v. COMENIC ACID.
AMIDO-COUMARIN & H.NO₂. 1168°-170°].
From nitro-coumarin (g. v). Noedles, v. sl. sol. cold water, v. sol. hot water.

(Frapolli a. Chiozza, 4. 95, 253);

AMIDO-COMENIC CH.NO. Mol. v. 149

AMIDO-CRESOL O, H, NO. Mol. w. 142. Ten amido-cresols are indicated by theory: four derived from ortho-, four from meta-, and two from para-cresol. The amido-cresols are readily soluble in alcohol and in ether, sparingly so in water. They dissolve in acids and in alkalis. They are formed by reducing nitro-cresols, or from nitro-toluidines by the diazo reaction.

Amide-o-cresels $C_yH_yMe(OH)(NH_x)$ [1:2:x]. Amide-o-cresel x=3. From nitro-o-cresel [69°] (Hofmann a. Miller, B. 14, 570; Zincke a.

Hebebrand, A. 226, 72).

Reaction.—1. When heated with quinone, it forms a rederystalline Base, C₂₈H₂₆N₄O₄, [285°], v. sl. sol. alcohol, sol. acids; its acetyl derivative, C₂₈H₂₆Ac₂N₄O₄, forms orango needles (from dry HOAc).—2. Heated with formic acid it forms a

methenyl compound: $C_0H_2Me <_0^N > CH$, [39°], (200°).

Methyl ether C_o11₃Me(OMe)(NH_z). (223°). Amide-o-cresol x = 4. [161°]. From nitro-ocresol [108°] (Nölting a. Collin, B. 17, 270). Also from acetyl-tolylene-di-amine, C_cH₃Me(NH₂)(NHAc) [1:2:4] (Wallach, B. 15, 2831). Colourless plates or needles. Salt.—

B'HCl: glittering plates, which sublime as needles.
Acetyl derivatives
GH, Me(OH)(NHAc). [225°]; sol. KOHAq.—
GH, Me(OAc)(NAcH). [133°] (Maasson, B.

17, 608; Wallach, A. 235, 250). Amide-o-crssel x=5. [175°].

Formation.— 1. From nitro-o-cresol [85°] (Hirsch, B. 18, 1514).—2. From nitroso-o-cresol. 3. From sulpho-benzene-azo-o-cresol by reducing with Sn and HCl (Nölting a. Kohn, B. 17, 865).—White plates or needles; may be sub-limed.—CrO, gives toluquinone. Salt.—B'HCl.

Amido-o-cresel x=6. [124°-128°]. From nitro-cresol [143°]. Stellate groups of neodles (Ullmann, B. 17, 1962). Salt.—B'HCl.

Undetermined derivatives of amilo-o-cresols.

Methylether C.H.Mc(OMe)(NH.) [1:2:5?].
[53°] (Hofmann a. Miller, B. 14, 571).

Ethyl ether C.H.Me(OEt)(NH2) [1:2:x].
From ethyl nitro-o-crosol [71°] (Staedel a. Kayser, A. 217, 217; B. 15, 1134). Salts.—
B'HClliaq.—B'2H2SO.,—B'2H2PtCl., Acetyl derivative C.H.Me(OEt)(NHAc). [108°]. Trimetric plates (from water); tables (from sther); cnbos (from benzone).

Amido-m-crssols $C_0H_3Me(OH)(NH_2)$ [1:3:x]. Amido-m-crssol x=6. [151°]. From eulphobenzene-azo-m-cresol by reduction (Nölting a. Kohn, B. 17, 367). White warts. On exidation with CrO_1 it gives toluquinone.

Undetermined derivative of an amido-m-cresol. Ethyl ether C₄H₂Me(OEt)(NH₂) [1:3:x]. An oil formed by reducing ethyl-nitro-m-oresol

[54°] (Staedel, A. 217, 219). Salt.—B',H.C.O., Acetyl dsrivative. [114°]. Mass of needles (from water).

Amido-p-crssols O_cH_sMe(OH)(NH₂) [1:4:x].

Amido-p-orssol x=2. [141°].

Formation. — 1. From nitro-p-oresol [78°] (Knecht, A. 215, 91).—2. From nitro-toluidine C_s[1_sMe.(NH.)(NO.) [1:4:2] (Wallach, B. 15,2833). Properties.—Colourless plates by sublimation.

Actyl derivatives

C₀H₃Me(OH)(NHAo) [178°]; sol. KOHAq. — C₀H₃Me(OAc)(NHAc)[129°](Maassen, B.17,608).

Methy t'ether C.H.Me(OMe)(NH.). [47°]. From the nitro-compound (K.); needles, volatile with steam.

Amide-p-cresol x=3. [135°].

Formation. — 1. From nitro-p-cresol [33°] (Wagner, B. 7, 1270; Hofmann a. Miller, B. 14, 572). — 2. By reducing benzene-azo-p-cresol or sulpho-benzene-azo-p-cresol (Nölting a. Kohn, B. 17, 350).

Properties.—White plates or needles; gives

a red colour with Fe.Cl. Salt.—B'HCl. Reactions.—1. Gives a methenyl derivative when heated with formic acid (H. a. M.).—2. Gives, when heated with Ac.O and NaOAc, an ethenyl derivative which is converted by boiling dilute H₂SO, into an acetyl derivative.

Acetyl derivative [160°]. Long needlos. Methylether C.H.Mc(OMe)(NH.) [38°].

Ethyt ether Callame(OEt)(N11). [41°]. From the nitro-compound (Staedel a. Kayser, B. 15, 1134). Needles (from water) or plates (from other solvents). Salts.—B'HClllaq.—B'_11SO_12aq.—Acetyl derivative [107°]. Di-amide-p-cresel.

Ethyl ether C.H.Me(OEt)(N11.), [1:4:3:5]. From the nitro-compound (Staedel a. Kayser, A. 217, 221). Pleasant-smelling oil—B'HCl:

silky needles.

Di-amide-cresel C₃H₂Me(OH)(NH₂)₂ [1:x:2:4]. From antido-toluene-azo-amido-cresol (Gracff, A. 229, 349); decomposes when liberated from its salts.—B"I₁SO₄aq; slender grey needles (from alcohol-ether).

AMIDO-CRESYL- v. Amino-tolyl-.

AMIDO-CROTONIC ETHER. A name applied to the imide of aceto-acetic ether (v. p. 19).

AMIDO-CUMENE v. COMIDINE.

Di-amido-cumens C₀H₁₁N₂ i.e. Fr.C₆H₃(NH₂)₂, [47°]. From the nitro-compound (Hofmann, J. 1862, 354).

Di amide - pssudo - cumene C_eHMe_s(NH₂)_s [1:3:4:5:6]. [92⁵]. Formation. — 1. By reducing nitro-pseudo-cumidine [47°] (Edler, B. 18, 630). 2. By reducing amido-azo-cumeno (Nölting a. Baumann, B. 18, 1147). Properties.—Needlss or plates; gives with Fe_sCl_s a brownish-rsd colour and a quinone-like smell; also gives Ladenburg's aldehydine reaction.

AMIDŌ-ψ-CUMENOL C.H.13NO i.e.

C. HMe.1(OH)(NH.2) [1:3:4:6:2] [167°]. Amidopseudo-cumenol; Oxy-cumidine. Obtained by
reducing benzene-azo-ψ-cumenol (Liebermann a.

Kostanecki, B. 17, 886); or nitro-ψ-cumenyl
nitrate (Auwers, B. 17, 2980). White nssdles
(by sublimation); sol. KOHAq. Fe.Cl. gives a
red colouration. Di-acetyl derivative

C. HMa. (OA)(NHA) (186°2) needles.

C.HMe,(OAc)(NHAo), [186°]: needles. AMIDO-CUMINIC ACID C, H;2NO,

m-Amido-cuminic soid Pr.O.H. (NH.)CO.H [1:2:4]. [129°]. Amido-isopropyl-benzoic acid. Prepared by reducing m-nitro-cuminio acid [158°] (Paterno a. Fileti, G. 5, 383; Lippmann a. Lange, B. 13, 1661).— Tables; some of it occasionally crystallises from water in thin plates [104°] (Fileti, G. 10, 12). Fe.Cl. give a violet-blue colourati... with the hydrochlorido. EtI at 100° gives a syr.py ethyl-amido-cuminic acid.

Salts.—AgA': white pp.—ZnA', 3aq: needles. HA'HCl.—(HA'IICl), PtCl.,—(HA'), H.SO., Acetyl derivative. [248°-250°]. Slender

needles, sl. sol, boiling alcohol, saponified by water at 230°.

Ethylether EtA'. Heavy oil

Nitrile Pr.C.H. (NH2).CN. [45°]. (305°). From nitro-cumino-nitrile.—Needles (from water) (Czumpelik, B. 2, 183). Salt.—(B'HCl), PtCl,

o-Amido-cuminic acid $\mathbf{Pr.C_cH_3(NH_c).CO_cH.}$ [1:3:4], [115°]. Prepared by reducing o-nitro-cuminic acid with FeSO, and NH₃ (Widman, B. 19, 270). —Plates or tables.

Acetyl derivative C₁₆II₁₂AcNO. [246°]. Slonder needles, sl. sol. alcohol, and other; may be sublimed (Widman, B. 16, 2579).

Di-amido-cuminic acid C,0H,1N,O, Pr.C.H.(NH.).CO.H. [192]. Formed by reducing di-nitro cuminic acid (Boullet, C. R. 43, 399; Lippmann, B. 15, 2141).—Yellowish plates, sol, hot water, alcohol, ether, alkalis and acids. Crystallises from water with aq.

Salts. - AgA'aq. - HA'HCl aq: large prisms. AMIDO-CUMYL-ACRYLIC ACIDS C, H1, NO.

I. Pr.C₆H₂(NH₂).CII:CH.CO₂H [I:3:4]. From the nitro-acid, FeSO, and NH, (Widman, B. 19, 262).—Flat yellow prisms (from alcohol). Salt.—HA'HCl'3aq: very slender needles, v. sl. sol water; converted by boiling water into cumostyril or (B. 3)-iso-propyl-(Py. 3)-oxy-quinoline [169°]. Acetyl derinative PrC₂H₂(NHAc).CH:CH.CO₂H, [220°]: very thin

ncedlos (from alcohol).

II. PrC.H.(NII.).CH.CH.CO.H [1:2:4]. From the nitro-acid (Widman, B. 19, 415). Sixsided tables (from ether). Warm II.SO, gives a magenta colour. Salts,--HA'IICl: flat needles. -(HA'HCl), PtCl, 2aq. --(HA'), H, SO, 5aq. Acetyl derivatives $C_{12}H_{14}Ac_{1}NO_{2}$, [240°]: needles (from alcohol).— $C_{12}H_{13}Ac_{2}NO_{2}$. [236°].

AMIDO-4-CUMYLENE-ACETAMIDINE

 $C_{11}H_{15}N_{3}$ i.e. $C_{6}Mc_{3}(NIL_{2}) < N_{H} > CMc[1:3:4:2:5]$.

Ethenyl - tri - amido - tri - methyl - benzene. [215°-218°]. From acctyl-di-nitro-pseudo-cumidine by reduction with Sn and HCl (Auwers, B. 18, 2663). - Rosettes of plates or yellowish prisms (from water) containing 2aq.

Salts. - B"2HClaq. - B"HCl 2aq. -B"H2PtCl, aq.

AMIDO-CUMYLPHENYLKETONE v. PHENYL AMIDO-CUMYL EKTONE.

m-AMIDO-CUMYL-PROPIONIC ACID

13

C_{1.}H_{1,}NO₂ i.e. PrC₂H₄(NH₂).CH₂.CH₂.CO₂H [1:3]. [103°-105°]. From m-amido-oumyl-acrylic acid, NaOHAq, and sodium-amalgam (Widman, B. 19. 418). Acetyl derivative C12H18AcNO2 [168°]; prisms (from alcohol).

AMIDO-DI-CYANIC ACID C.H.N.O i.e. NH, CO.NH.CN or HN:C NH CO. phano-nitrile; carbc nyl guanidine; carbimidocyanamide.

Formation. ... From di cyano-di-amide and baryta-water.—2. Fry. potussio cyanate and cold aquecus cyanamics (Hallwachs, A. 153, 239; Wunderheb. 19, 48).

Properties.—18, 465. It decomposes car-

bonates and be saves as a strong acid. Produces biuret when warmed with II,SO, (I vol.) and water (2 vols.) at 70° (Baumann, B. 8, 708).

Salts.—Nac, III,No, —KA'.—BaA', 3aq.—CuA'., 4aq.—CuC, iIN,O 2aq.—AgA'.

AMIDO.CYANURIC ACID v. AMMELIDE.

Di-amido-cyannric acid v. AMMELINE.

AMIDO-CYMENE v. CYMIDINE. Diamido-cymene C.II.Mel'r(NH.),[I:4:3:6].

Ilydrochloridc. Formed by reducing the di-oxim of thymoquinone (Liebermann a. Ilinski,

B. 18, 3200). AMIDO-DRACYLIC ACID = p-AMIDO-BENZOIC ACID.

DI-AMIDO-DURYLIC ACID v. DI-AMIDO-TRI-METHYL-BENZOIO ACID.

AMIDO ETHANE v. ETHYLAMINE.

Di-amido-ethane v. ETHYLENE DI-AMINE. AMIDO-DI-ETHYL-ACETIC ACID v. AMIDO-HEXOIC ACID.

AMIDO - ETHYL ALCOHOL v. OXYETHYL-AMINE.

AMIDO-ETHYL-BENZENE v. AMIDO-PHENYL.

AMIDO-ETHYL METHYL KETONE METHYL AMIDO-ETHYL RETONE

DI-AMIDO-DI-ETHYL OXIDE *C.HI 12N.O i.e. (CH₂.C(NH₂)II)₂O.

Di-amido-cther. The very unstable hydro. chloride (B'2HCl) of this body is formed by passing NH₃ into an othereal solution of (ClI₃,CHCl)₂O (Hanriot, A. Ch. [5] 25, 224).

DI-AMIDO DI-ETHYL-DIPHENYL C16H20N i.c. [4:3:1] NH₂.C₆H₄Et.C₆H₃Et.NH₄ [1:3:4] (?). Formed by the action of SuCl₂ and HCl or H₂SO₄ on an alcoholic solution of o-azo-ethyl-benzene.

-B"II₂SO₄: needles, sl. sol. water, m. sol. alcohol. Acetyl derivative C₁₆H₁₆(NHAc)₂. [307°]. White needles (by sublimation), sl. sol. alcohol, sol. IIOAo.

An isomeric di-amido-di-ethyl-diphenyl is formed similarly from p-azo-ethyl-benzene. Its sulphate is a white amorphous powder (G. Schultz, B. 17, 474)

AMIDO-ETHYL-TOLUENE v. AMIDO-TOLYL-ETHANE.

AMIDO-ETHYL-TOLUIDINE v. ETNYL-TOLY-LENE DIAMINE.

AMIDO-ETHYL-m-URAMIDO-BENZOICACID C10H13N3O3 i.e. NH2.C2H1.NH.CO.NH.C0H1.CO2H. Formed by the action of ethylene diamine upon cyano-earboxamido-benzoio acid (v. AMIDO BENzoio Acid). White prisms, sl. sol. cold water. Salt. - HA'IICl 2 aq (Gricss, B. 18, 2416). • AMIDO-FLAVOLINE v. FLAVANILINE.

AMIDO-FLUORENE v. FLUORENE. AMIDO-FORMIC ACID v. CARBAMIC ACID. AMIDO-FURFUR-BUTYLENE OXIDE C₆H₁₁NO₅ i.s. C₆H₃O.C(NH₂) CMe,.

(215°-220°). Obtained by reducing CaH10N2O4, the product of addition of N.O. to furfur-butylene.

Properties.—Colourless liquid, volatile with steam, sol. water.

Salts.—B'HClaq: crystals, v. sol. water.— B'_2H_PtCl_: sol. hot. water.

Acetyl derivative C_H_O_NHAc [153°];

(305°-310°): needles, v. v. l. HClAq. Anhydride C,HNO (5142°). (300°-310°). V.D. 4.77. Formed from an in furfur-butyleno oxide on dietilling, or even on kepping. It forms large colourless cryetals, and is volatile with steam. It is a tertiary base (Tonnies a. Staub, B. 17, 854).

AMIDOGEN. The group NH2; v. AMIDEE,

Amino-acids, and Amines.

AMIDO-GLUTARIC ACID v. GLUTAMIO ACID. AMIDO-GLYCOLLIC ACID v. OXY-AMIDO-ACETIC-ACID.

AMIDO-HEMIPIC ACID v. HEMIPIO ACID.

AMIDO-HEPTOIC ACID C,H,,NO, i.e. C_sH₁₁.CH(NH₂).CO₂H. a-Amido-wnanthic acid. From brome-heptoic acid and alcoholic NH, at 100° (Helms, B. 8, 1168). Six-sided tablee or plates, v. sl. sol. cold water, ineol. alcohol.

Salts.—CuA'2: incoluble powder.—HA'HCl:

prisms, v. eol. water or alcohol.

AMIDO-HEXOIC ACID CalliaNO2.

a-Amido-n-hexoic acid v. LEUCINE.

Amido-di-ethyl-acetic acid CEt2(NH2).CO2H. Preparation .- Di-ethyl kctono cyanhydrin, CEt (OH).CN, obtained by the action of dilute HCN on di-cthyl ketone, is heated with alcoholic NH_3 , and the product is saponified by HCl.

Properties .- Thick tables or prisme (from water), v. eol. water, m. sol. alcohol, insol. ether;

may be sublimed.

Salte. - AgA': white plates. - CuA'2: violet plates.—HA'IICl: thick white prisms (Tiemann a. Friedländer, B. 14, 1975).

a-Amido-iso-butyl-acetic acid

Pr.CH., CII(NH.).CO.H. S. 85 at 12°. From iso-valeric aldeliyde-ammonia, HCN and HClAq (Limpricht, A. 94, 243; Hüfner, J. pr. [2] 1, 10). Properties.—Resembles leucine, but is opti-

cally inactive (Mauthner, H. 7, 223).

AMIDO-HEXYL ALCOHOL v. DI-ACETONE-ALCAMINE.

AMIDO-HIPPURIC ACID C.H 10 N2O3 i.e.

C.H.(NH2).CO.NH.CH2.CO2H.

m-Amido-benzoyl-glycocoll. [194°] (Conrad, J. pr. [2] 15, 258). S. 3 at 20°; S. (alcohol) 08 at 15°. From m-nitro-hippuric acid, ammonium eulphide and H2S (Schwanert, A. 112, 70).

Properties.-Plates or needles. Soluble in

alkalis and in acids. Salt .- B'HCl.

Reactions.-1. Boiling HCl forms m-amidobenzoic agi: and glycocoll .- 2. Urea forms nramido-hippuric acid C10II11N2O4 and a small quantity of carboxamido-hippuric acid C₁₀H₁₄N₁O₂ (Grices, J. pr. [2] 1, 135).

DI-AMIDO-HYDRO-ACRIDINE KETONE, so called, C13H11N3O. [223°]. Formed by reducing the o-carboxylic acid of di-nitro-di-phenyl-amine with Sn and HCl. Flat needles or thick prisme, v. sol. hot alcohol, v. sl. eol. ether, benzene, cold water, and light petroleum. Fe₂Cl₆ gives a doop yellow colour, passing into greenish-black; K₂Cr₂O, gives a red pp. Salt.—B'HCl: thin colourless neodles, sl. sol.

cold water.

Ohloro-derivative O18H10ClN2O. [0. 280] Formed by reducing ohloro-di-nitro-di-phenylamine o-carboxylio acid. Colourless crystals, sol. hot water, v. sl. sol. cold water, ether, and benzene (Jourdan, B. 18, 1450).

AMIDO-HYDRATROPIC ACID v. AMIDO-

PRENYL-PROPIONIO ACID.

v. Hydrazines.

AMIDO-HYDRO-CARBOSTYRIL CoHioNaO. Oxy-amido-di-hydro-quinoline.

(B. 3)-amido-hydro-carbostyril

CH₂CH₂CH₂[3:6] C.H.(NH2) NH.CO

Di amido-phenyl-propionic anhydride. amido-hydro-cinnamic anhydride. [211°]. Prepared by reducing di-nitro-phenyl-propionic acid with tin and HCl. It forms colourless needles or prisme, v. sol. bot water, alcohol, and HOAc, ineol. CS.

Salte.—B'HCl: needles.—B'2H2PtCl6: yellow leaflots. Bromine forms a mono-bromo-derivative, [210°] and a di-bromo-derivative, [179°], both crystallising in needles (Gabriel a. Zimmer-

mann, B. 12, 601).

(Py. 4)-Amido-hydro-carbostyril CH₂ — CH₂

C,H [143°]. N(NH.,).CO

From CaH, (NH.NH.SO3Na)CH2CH2.CO2H (v. HYDRAZIDO-PHENYL-PROPIONIC ACID) by addition of IICl (Fiecher a. Kuzel, A. 221, 282). Crystallised from water. Sol. water, v. sl. sol. alcohol. Does not reduce boiling Fehling's solution, but reduces hot Ag.O. Salt. - C. H. N.OHCl.

Reactions. - 1. An acid colution is converted

by NaNO, into hydro-carbostyril.-- 2. With EtI and alcohol at 100° it gives an ethyl derivative $C_{s}H_{s} < CH_{2} - CH_{2} > .$ This gives a nitrosamine with NaNO, and HCl.

AMIDO-HYDRO-CINNAMIC ACID v. AMIDO-

PHENYL-PROPIONIO ACID. DI - AMIDO DI - HYDRO - TERE - PHTHALIC

ACID NH2.C.CII(CO2II).CH (?) Di-imido-hexa-

H.C.CH(CO,H).C.NIL,

hydro-terephthalic acid or succino-succinic-aciddi-imide.

Ethyl ether A"Et. [181°]. Obtained by fusing di-oxy-di-hydro-terephtbalic other (succino-succinic ether) with ammonium acetate. Yellow needles; al. sol. alcohol and ether with a green fluorescence, v. sol. eliloroform. treatment with bromine in H2SO, solution it is convorted into di-amide-terephthalic ether. The hydrochloride and eulphato are colourless sparingly soluble salte (Baeyer, B. 19, 429).

AMIDO-TETRA-HYDRO-QUINOLINE

 $CH_2 - CH_2$ C,H,2N, i.e. C,H, N(NH₂).CH₂.

[56°]. (c. 255°). Prepared by reducing the nitrosamine of tetra-hydro-quinoline with zinc dust and HOAo. White crystale. Salts.—B'₂H₂SO₄ 2aq: yellow plates, sl. sol.

cold water .- The hydrochloride is y. sol. water.

Reactions.—Reduces salts of Au and Pt, and Fehling's solution. Ppd. HgO forms an azoquinoline (Hoffmann a. Königs, B. 16, 780).

AMIDO HYDRO-QUINONE C.H.NO.

NH, C, H, (OH), Di-methyl derivative $NH_2.C_6H_2(OMe)_2$ [82°]. (270°). Formed by reducing the dimethyl-derivative of nitro-hydroquinone. Pearly plates, sol. hot water, alcohol, benzene, light

Places, soil now water, account, beinded, agreement and CS₂. Very readily oxidised.

Reactions.—1. CnSO₄Aq gives a greement black colour.—Fe₂Cl₆ pps. lustrous greef shelates, which form a red solution in water. AgNO, gives a silver mirror. Salts.—*B'LCI: white needles. - *B'2H2PtCl8: brown pp.

Acetyl derivative.—C,H,(NHAc)(OMe), [91°]. Silvery seales; sol. water, alcohol, benzene, light petroleum and CS, (Magatti, B. 14, 70; G. 1881, 352; Mülhänser, A. 207, 254; Baessler, B. 17, 2119).

Ethyl derivative C.H. (NH.) (OEt) (OH). From the nitro-compound (Wesolsky a. Beucdikt,

M. 2, 370).— B'HCl.

Di-amido hydroquinone C_eH_e(OH)_e(NH_e)_e Formed by reduction of di-nitro-hydroquinone or its di-acetyl derivative with tin and IICl. Owing to its easy oxidisability the base was not isolated in the free state. B"H2Cl2.-Colourless noedles, v. e. sol. water, sl. sol. conc. HCl

Di-acetyl derivative C₀II₂(OII)₂(NIIAe)₂. [o. 240°]. Colourless needles. Is oxidised to di-acetyl-di-amido-quinone C₀II₂O₂(NIIAc)₂.

Tetra-acetyl derivative C₀H₂(OAc)₂(NHAc)₂.

[216°]; colourless needles or plates; v. sol. alcohol and acetic acid, sl. sol. water and other. Dissolves in dilute alkalis, the solution becomes oxidised on exposure to the air and deposits yellow needles of the above-mentioned di-acetyldi-amido-quinone (Nietzki a. Preusser, B. 19, 2247).

Di-methyl-derivative CaH2(NH2). (OMe)2. The hydrochloride of this body is formed by reducing the corresponding nitro compound. It erystallises in needles, [169°] (Kariof, B. 13, 1676).

Di-methyl-di-amido-hydroquinone (q. v.) is isomeric with this body.

DI-AMIDO-DI-IMIDO-BENZENE

C₆H₂(NH₂)₂(NH)₂ [1:2:4:5]. Small brown needles. Formed by oxidation of solutions of salts of tetra-amido-benzeno with Fo,Cl, &c.

Salts .- B"H2Cl2: glittering brown needles, sparingly soluble in water with a bluish-violet colour. — B"(HNO_s)₂: small greeu needles (Nietzki a. Hagenbach, B. 20, 335).

AMIDO-IMIDO-METHANE v. FORMAMIDINE. AMIDO-DI-IMIDO-(a)-NAPHTHOL

$$C_{10}H_{\nu}N_{3}O$$
 i.e. $C_{10}H_{4}(NH_{2})(OH) < NH \atop NH$ (?). Pre-

pared by the reduction of tri-nitro-(a)-naphthol with tin and HCl. Brown seales, insol. water and ether. Salts. - B'HCl: lustrous green scales, sl. sol. cold water. -B'2H2PtCl, (Diehl a. Merz, B. 11, 1663).

ducing tri-nitro-orcin with sodium-amalgam (Stenhouse A. 167, 167). Lustrous green needles, v. sl. sol. water, insol. alcohol, ether, and benzenc. NaOHAq forms a deep blue solution. Reduced by further action of sodium-amalgam to triamido-orcin. Salts.—B'HCl aq: brownish-red needles; sol. water but ppd. by HCl.— B'₂H₂SO, 2aq: purple lamine. AMIDO-DI-IMIDO-PHENOL, so called.

$$C_{e}H_{1}N_{0}O$$
 i.e. $C_{e}H_{2}(NH_{2})_{2} < \prod_{i=1}^{NH} \left[2:6:\frac{1}{4}\right]$.

Di-amido-quinone-im e (Hepp, A. 215, 351). The hydrochloride, JHCl, separates as brown needles with figer have then Fe Clais added to a cone, aqueous solution of the hydrochloride of tri-amido-phenol (Heintzel, Z. 1867, 342). It is decomposed by alkalis and by hot water; hot dilute HCl changes it into colourless needles of the hydrochloride of oxy-amido-quinone-imide (or di-amido-quinone) C.H.N.O.HCl, while H.SO. forms a corresponding sulphate crystallising in

AMIDO-IMIDO-DI-PHENYL SULPHIDE

 $C_{12}H_{10}N_2S$ i.e. $HN < C_{\alpha}H_{\alpha}(NH_2) > S$. Anido-thiodiphenyl-imide; Anido-sulphido di-phenyldi-phenyl-imide;

Formation.-1. By reduction of nitro-imidodi phenyl-sulphoxide, HN C.H. SO. 2. By heating p-amido-di-phenyl-amine with sulphur (Bernthsen, B. 17, 2858; A. 230, 101). White satiny plates (from water); m. sol. hot water, v. sol. alcohol and ether. Turns grey in moist air. Fe₂Cl₆ converts it into the following

body : Imido-imido-di-phenyl sulphide $C_{12}H_8N_2S$ i.e. $N <_{C,II}$ >s.

Small brown crystals, v. sol. alco-

hol, sl. sol. water and ether. Its salts dye silk greyish-violet. It is easily reduced to the pre-ceding body. Salts.—B'HCl 'aq.—B'₂H₂ZnCl₄: brown needles or prisms.

Di-amido-imido-di-phenyl sulphide

 $C_{12}H_{11}N_3S$ i.e. $HN < C_0H_3(NH_2) > S$. Formed by reducing (a)-di-nitro-imido-di-phenyl sulphoxide with tin and 11Cl, or Lauth's violet with ammonium sulphide (Bernthsen, B. 17, 614). Yellow needles or plates; sl. sol. water and ether. The sulphate is sl. sol. water.

Amido.imido-imido-di-phenyl-snlphide

Formation.-1. By treating a solution of pphenylene diamine hydrochloride with H.S and Fe.Cl. successively (Lanth, C. R. 82, 1441; Koch, B. 12, 592, 2069.—2. By action of Fe, Cl, on the preceding body (B.). Its alcoholic solution has a violet colour with reddish-brown fluorescence. Its solution in excess of HCl is blue; in H.SO., green changing to blue and then to violet. Long heating with MeI converts it into the methyloiodide of ponta-methyl-di-amido-imido-di-phenyl sulphide identical with that formed in the same way from methylene blue. Salt.—B'HCl: green crystals, sl. sol. cold water.

An isomeride (Bernthsen's Violet) is formed by reducing (β) di - nitro - imido di - phen¶l sul-phoxide and then oxidising the leuco base with Fe Cl. Its hydrochloride B'2HCl, forms dark needles, which dye reddish violet. H.SO, forms * violet solution.

AMIDO-DI-IMIDO RESORGIN C.H,N.O. aq

or $C_6H(OH)_2(NH_2) < \prod_{NH} (?)$. From tri-amidoresorcin hydrochloride and Fo.Cl. Instrous green needles, v. sl. sol. ater, insol. alcohol or ether: KOI1Aq forms a be solution. Dilute

HCl at 170° forms tri-ca, Histone.

Salt.—B'HCl: red needles ppd. by HCl

Now B 11

(Sehreder, A. 158, 250; Diehl a. Merz, B. 11, 1229).

AMIDO INDIGO C₁₁H₁₂N₁O₂ i.e. C₁₄H₈(NH₂)₂N₂O₂. Prepared by reducing nitroindigo with acetie acid and powdered zine (Baeyer, B. 12, I317). Dark violet pp., v. sl. sol. alcohol, ether and chloroform. Forms blue solutions in dilute acids.

AMIDO-ISATIN, so called, v. Isatinimide.

AMIDO-LACTIC ACID v. OXY-AMIDO-PROPIONIO

AMIDO-MALEÏC ACID C₁H, NO₁ i.e. CO₂H.CH:C(NIL₂).CO₂H. [182°]. Easily solublo erystals. Prepared by saponification of the amides. - A"Ag2: voluntinous pp., explosive.

Diethyl ether Et.A". [100°]. Colourless prisms. Sol. alcohol and ether, insol. water. Prepared by the action of alcoholic NH, (2 mols.) on ohloro-maleïe ether (1 mol.).

Amido-maleamic-ethyl ether

 $C_2H(NII_2)$ $< \begin{array}{c} CO.NII_2 \\ CO.Et \end{array}$. [62°]. Long white prisms. V. sol. alcohol and ether, insol. cold water. Prepared by the action of alcoholic NH₃ (3 mols.) on chloro-maleic ether (I mol.).

C₂II(NII₂)<CO.NII₂ [122°]. Diamide Colourless plates. Sol. alcohol, ether, and hot

water. Prepared by the action of an excess of strong alcoholic NH₃ on eldoro-maleïc ether (Claus a. Voeller, B. 14, 150).

AMIDO-MALONIC ACID CaH, NO, ie.

CH(NH2)(CO.H)2. Obtained from nitroso malonic acid by reducing with sodium-amalgam (Baeyer, A. 131, 295). Prisms (from water) or needles (by ppg. with alcohol). When heated alone, or in aquoous solution, it splits up into CO, and glycocoll. Iodine oxidises it, in aqueous solution, forming mesoxalic acid.

Salt.—Pb(C₃II₄NO₄)₂: crystalline pp. Amide CH(NH₂)(CO.NH₂)₂. [182°]. Formed by heating chloro-malonic ether with alcoholic NH3. Prisms, sol. hot water (Conrad a. Guthzeit,

AMIDO-MESITOL C,H,3NO i.e.

C.HMc.(NH.)(OH) [I:3:5:2:4]. A very oxidisable body formed by reduction of nitro-mesitol.—B'HCl: needles (Knecht, B. 15, 1376).

AMIDO MESITYLENE v. "TESIDINE.

Di-amide-mssitylene C₂H₁₄N₂ i.e. C₄HMe₄(NH₂)₂. [90°]. From di- or tri-nitromesitylene with tin and HCl. Long slender needles (from water), or large monoclinic crystals (from ether). Sublimes in needles. V. sol. alcohol or ether, m. sol. hot water. CrO, oxidiscs it to oxy-iso-xyloquinone, CallMe (OII)O2.

Salts.—B"HCl: square tables (from water), ppd. by HCl.—B"H.C.O.; hard grains (from water).—B"H.SO.; broad lamine (from water). Di-acetyl derivative, [above 360°]; v. sl. sol. water or cold alcohol (Fittig, A. 141, 134; 180, 27; Ladenburg, A. 179, 176). AMIDO-MESITYLENIC ACID C.H., NO.

o-Amido-mesitylenio aoid

C.H.Me.(NH.).OO.H [I:3:4:5]. [187°] (Schmitz, A. 193, 171). [190°] (Jacobsen, B. 11, 2055). From the nitro acid with tin and HCl. Long needles (from alcohol). Splits up when heated with lime into CO, and (1, 3, 4)-xylidine.

Amide-mssitylsnic acid C₆H₂Mc₂(NH₂)CO₂H [I:3.2:5]. [235°]. From the nitro acid. Long needles (from aloohol). Sl. sol. water, v. sol. hot alcohol. Gives (1, 3, 2)-xylidine when

heated with lime.

Salt .- B'IICl: long needles (Fittig a. Brückner, A. 147, 50; Jacobsen, B. 12, 608).

AMIDO - METHENYL - AMIDO - PHENYL MERCAPTAN O, H6N2S i.e. C6H4 < N C.NH2. [129°]. Prepared by heating chloro-methenylamido-phenyl mercaptan with alcoholie NH, at

160°. Nacreous laminæ. Fusion with potash produces amido-phenyl mercaptan. Salt .- It is a weak base, and has a crystalline platinochloride, B',H,PtCl, (Hofmann,

B. 12, 1129; 13, 11). AMIDO METHOXY COMPOUNDS v. Methyl

derivatives of Oxy-amido compounds. AMIDO-DI-METHYL-ACETIC ACID v. AMIDOiso-butyrio acid.

DI - AMIDO - TETRA - METHYL - DI - AMIDO -DIPHENYL v. Tetra - METHYL - tetra - AMIDO-DIPHENYL. And, in general, amido-methyl-amido compounds are described as methyl-(di)-amido compounds.

AMIDO - DI - METHYL - ANILINE v. Di-

METHYL-PHENYLENE DIAMINE.

AMIDO - METHYL - ANTHRACENE HYDRIDE C₁₃H₁₅N i.e. C₁₄H₁₀Me.NH₂. [79°]. Prepared by heating amido-methyl-anthraquinone with HI and P at 150°. Glistening lamine, which begin to sublino at 130° and are sol. alcohol, ether, chloroform, benzene, glacial acetic acid, and CS2, but v. sl. sol. water.

Reactions .- 1. Nitrous acid gives a green colour, and on adding NII, a red pp.-2. Arsenic acid gives a brownish-red mass after fusion. Salt .- B'IICl [245°]: glistening needles. Acetyl derivative C₁₅H₁₄AcN. [198°]. White needles, sol. alcohol and other (Roemer, B. 16, 1631).

AMIDO-METHYL-ANTHRANOL

C₁₃H₁₃NO i.e. C₁₂H₆Me(NH₂) [183°].

Prepared by heating amido-methyl-anthraquinone with III (S.G. 1.96) and P. Crystallises in nearly white needles, but sublimes in red needles. Sol. alcohol, ether, benzene, and glacial acetio acid, v. sl. sol. water.

Reactions.-I. H2SO4 gives a yellow solution, becoming purple-red on warming. — 2. HNO₃ gives a violet colour, turning red.—3. Air reconverts it, in alkaline solution, into amidomethyl-anthraquinone.

Di-acetyl derivative C., H., Ac, NO. [170°]. Thick white needles; its alcoholic solution exhibits blue fluorescence (Roemer a. Link, 16, 703).

AMIDO-METHYL-ANTHRAQUINONE C15H11NO2 i.e. C11H6O2(CH2)(NH2). [202°]. Propared by reduction of nitro-methyl-anthraquinone. Long dark red needles. Y. sol. alcohol, ether, benzene, acetic acid, and chloroform, v. al. sol. water.

Acetyl derivative C, H,O,(CH,)NHAc. [177°]. Small light-red needles, sol. alcohol and glacial acotic acid (Römer a. Link, B. 16, 698).

AMIDO METHYL BENZENE v. TOLUIDINE and BENZYLAMINE.

Amido-di-methyl-ber tono v. Xx IOINE. Amido-tri-rethyi-beazene g. Mesidine nd **↓**-Cumidine.

Amido-tetra-methyl-benzene v. Duriding. Amido - penta - mothyl - benzeae v. Penta-

METHYL-PHENYL-AMINE. DI-AMIDO-TETRA-METHYL-BENZIDINE

v. Tetra-METHYL-tetra-AMIDO-di-PHENYL. AMIDO METHYL BENZOIC ACID v. AMIDO-

TOLUIO ACID. Amido - di . methyl - benzoio acid v. Amido-

MESITYLENIO ACID.

Di-amido-tri-methyl-benzoic acid CieH., N.O. i.e. C₈Me₂(NH₂)₂CO₂H [6:4:3:5:2:1]. Di-amido-durytic acid. [221°]. Formed by reducing the di-mitro compound with zinc dust and diluto HOAc. Celeurless silky needles, sol. hot water and hot alcohol, v. sl. sol. other. Fe.Cl. oxidises it to pseudo-cumo-quinone carboxylio acid (Nef. B. 18, 3496; A. 237, 1).

Acetyl.derivative [275°

AMIDO.TRI.METHYL.BUTYL.LACTICACID

v. Oxy-amido-heptoic acid.

m-AMIDO-a-METHYL-CINNAMIC HYDE C10 II 11 NO i.c. C4 II (NH.). CH: CMe. CHO [60°]. Got by reducing the nitro compound with FeSO, and NII, Yellowish crystals; dissolves in aqueeus acids, and reduces ammoniacal AgNO₃. Phenyl hydrazide

C.H. (NIL.).CII:CMe.CII:N.HPh; [157°]: needles.

Acetyl derivative

C.H.(NIIAc).CII:CMc.CIIO; [120°]: short thick prisms (Miller a. Kinkerlin, B. 19, 1248).

AMIDO - (B. 2. Py. 2) - DI METHYL - (Py. 3)-ETHYL-QUINOLINE C₁₃H₁₆N₂ i.e. C₂II₁N(NII₂)Me₂Et. [149]. Plates, monoclinio

tables, prisms or flat needles. Formed by reduction of the nitro-derivative with SnCl2. Salts .-B'HCl; very soluble colourless crystals. The nitrate and sulphate are also easily soluble in water (Harz, B. 18, 3392).

AMIDO - (Py. 4)-METHYL-HYDRO-QUINO-LINE

, CH₂.CH₂ C10H14N2 i.e. C1H3(NH2)

Yellowish oxidisable Amido-kairoline. Formed by reduction of nitro-kairoline [94°] with SnCl₂. By nitrous acid it is converted into a compound $C_{10}H_{13}N_3O$, [144°] when dry, which crystallises with 5aq in splendid red needles, and dissolves in dilute acids with a deep red colour.

Salts.—The acid tartrate forms sparingly soluble crystals. B"H2Cl2PtCl4 (Feer a. Koenigs, B. 18, 2391)

AMIDO DI METHYL-HYDROQUINONE

di-methul-Amido-Hydroguinone

7052

AMIDO-TRI-METHYL-PHENYL-ACETAMI. DINE v. Amido-\psi-cumylene-acetamidine.

AMIDO-DI-METHYL-PHENYL-ACETIC AN-HYDRIDE v. Di-METHYL-OXINDOLE

AMIDO-METHYL-PROPYL-BENZENE CYMIDINE.

(B.4)-AMIDO (Py. 8)-MHTHYL-QUINOLINE C₁₀H₁₀N₂ i.e. C₁H₂(CH₂)(NH₂)N. o-Amido-quinaldine. [56°]. Formed by reduction of (B.4)nitro-(Py. 3) methyl-quinolino. Long prisms. V. sol. alcohol, ether, and hot ligroine, sparingly in water.—B'HCl: yellow needles (Doebner a. Miller, B. 17, 1701)

(B. I or 3)-Amido (Py. 3)-methyl-quinoline C₁₀H₁₀N₂aq i.e. C₃H₁(C₄₃(NH₂)N m-Amido-quin-aldine [105°] when " " Formed by reduction of (B. 1 or 3) -nitro₁ P₃", 3) -methyl-quinoline. Colour-less crystals (+H.O). V. sol. hot water, alcohol, and benzene, sparingly in ether. B'HCl: red needles (Doebner a. Miller, B. 17, 1702).

Amido - (B.2, 4.Py.3) - tri methyl quincline C12II12N(NII2). Formed by reducing nitro-trimethyl-quinoline with SnCl. Yellowish plates (from alcohol) (Panajotow, B. 20, 36).

Sulphonic acid C12111 (SO3H)(NH2).N. Small yellow needles. BaA', 3aq: silky needles, sl. sol. cold water.

AMIDO METHYL-TOLUIDINE v. METHYL-TOLYLENE DIAMINE.

AMIDO-B-METHYL UMBELLIFERON

CMe:CII C10 H2NO i.e. C4H2(NH2)(OH [247°]_ 0 . CO

From nitro-β-methyl-umbelliferon. Needles, sparingly soluble in the usual meustrua. Fe,Cl. gives an intense green colouration with the alcoholic solution.

Salt. -B',ILSO, 2aq: sparingly soluble pp. (Pechmann a. Cohen, B. 17, 2137).

AMIDO METHYL URACIL v. URAMIDO CRO-TONIC ACID.

AMIDO-NAPHTHALENE v. NAPHTHYLAMINE. Di - amido - naphthalene v. NAPHTHYLENE: DIAMINE.

Tri-amido-naphthalene C₁₀H₁₁N₂ i.e. H₃(NH₂)₂. The hydriodide, B"3HI, is C10 II (NII .)3. formed from (\$)-tri-nitro-naphthalene [2180], by redacing it (1 g.) with I (20 g.), P (4 g.) and water; it forms white needles; at 70° it becomes B"2HI. The sulphate, B"H2SO, forms silky needles. The free base is very unstable.

Benzoyl derivative C₁₇H₁₅N₃O i.e. C₁₆H₁(NII)₂(NIIBz)[2:4':1]. From benzoyl-dinitro naphthylamine.

Salts. - B"HCl: ncedles. -- B‴H,80.: needles (Lautemaun a. Aguiar, Bl. [2] 3, 263; Hübner a. Ebell, A. 208, 324).

Tetra-amido-naphthalene C₁₀H₁₀N₄ i.e. C₁₀H₄(NH₂)₄. The hydriodide B'''4HI, formed by reducing (β) tetra-nitro-naphthalene [200°] with P, I, and water, crystallises in yellowish lamino, sol. water and alcohol (L. a. A.).

AMIDO-NAPHTHALENE SULPHONIC ACID

v. Naphthylamine sulphonic acid. AMIDO.(8)-NAPHTHO.HYDR3@UINONE

C10H2NO, i.e. C10H2(NH2)(OH)... Amido-di-oxy-naphthalone. The hydrochloride, B'HCl, is got by roducing nitro-(β)-naplitholydroquinone with tin and HCl (Groves, C. J. 45, 300). It is rapidly oxidised by air.

AMIDO-NAPHTHOIC ACID C,H,NO, i.e. C₁₀H₈(NH₂).CO₂H.

Amido-(a) naphthoio aoid.

[212]]. From nitro-(a)-naphthoic acid, [230°], by reducing with FeSO, and NH2. Colourless needles; may be sublimed; sol. alcohol, v. sl. sol. ether (Ekstrand. B. 18, 78).

Amido-(8) naphthoic scid. [211°]. Formed by reducing nitro-(8) naphthoic acid (269°), with FeSO, and NH,. Sisnder colourless needles (Ekstrand, B. 18, 1206).

[2I9']. From Slender Amido-(6)-naphthoio acid. nitro - (8) - naphthoic acid [289°]. needles. Salts .- * HA'HCl: small prisms, v. sol. water. - *HA'IINO3: largo thin lamine. -*(HA'), H.SO.: small prismatic needles (E.).

Amido-(3)-naphtheic acid [232°]. From nitro-(3)-naphthoic acid [293°]. Schall trimetric tables; sol. alcohol and boiling water. Salts .-CaA', 4aq: long violet needles. — HA'HNO,: largs needles. -(IIA'), H.SO,: needles (E.).

peri-Amido-naphtheic acid C10H.(NH.)CO.II

 $[\alpha_1 \alpha'_1].$

Preparation.-The orude product of the nitration of (a)-naphthoic acid is reduced with FeSO, The solution is boiled with HCl and, on cooling, the lactam of the peri-acid, [178°], crystallises out in yellow needles, whilst tho hydrochlorido of the isomeric acid remains in solution.

By diazotisation, treatment with cuprons oyanide, and saponification of the nitrile, naplithalic acid is formed (Bamberger a. Philip, B. 20, 242). This experiment determines the constitution of accomplithene (q. v.) which may be oxidised to naphthalic acid.

Lactam C₁₁H, NO

Formed by reducing nitro-(a)-naphthoic acid [215°]. Needles (by sublimation); sol. hot alcohol, sl. sol. water and ether. It is an indifferent substance insoluble in alkaline carbonates, but soluble in het aqueous NaOH with formation of the acid (Ekstrand, B. 18, **75**; I9, 1137; comparo Rakowsky, B. 5, 1020).

Acetyl derivative CullaAcNO. [125°]. Long hair-like needles (from alcohol).

AMIDO-NAPHTHOL C10H9NO.

(a)-Amido-(a)-naphthol Cioll (OH)(NH.) [1:1]. Formed by reducing nitro-(a)-naphthol [161°] (Liebermann, A. 183, 217) or p-sulpho-benzeneazo-(α)-naphthol (Liebermann, B. 14, 1796).

The free base is unstable; its salts produce

(a)-naphthoquinone when oxidised.

B'HCl: white needles; converted by bleaching powder into C₂₀H₁₀N₃Cl or C₂₀H₁₂N₃Cl which separates from HOAcAq in needles [85°], and explodes at 130° (Hirsch, B. 13, 1910).

(B)-Amido-(a)-naphthol C1011, (OH) (NH.) [1:2]. Formed by reducing nitro-(a)-naphthol [128°], or

nitroso-(*) naplithol (L.).

The free bass is unstable; in presence of alkalis, air Torms violet naphthoquinono-imide,

. Its salts give (β)-naphthoquinone

when oxidised. B'HCl: white lamine.-B'C,H2(NO:),OH.

(a) - Amido - (β) - naphthol C₁₀II (OH)(NH₂). Formed by reducing nitro-(\$)-naphthol [103] (Jacobsen, B.14,806; A.211,48) or nitroso-(B)-naphthol (Groves, C.J.45,296).

Priparation.—(β)-naphthol orange is heated with HClAq and SnCl, in slight oxcess over that

required by the equation:

HO.O., H., N., O.H., SO., Na + 2SnOl, + 6HOl = HO.O., H., NH, HCl + NH., C., H., SO., H+ 2SnCl. + NaCl.

Amido-(\$)-naphthol hydrochloride crystallises on cooling, and is freed from sulp'anilio acid washing with NaOHAq (Groves, C. J. 45, 291).

Properties.—Colourlsss scales, v. sl. sol. water, readily oxidised by air. Its ethereal solution fluoresces violet. Chromic mixture oxidises it to (B) naphthoquinone.

Salt .- B'HCl: white needles.

Benzoyl derivative.—C10H3(NHBz)OH. [245°]. Small colourless plates, soluble in alkalis. Formed by reduction of the benzoylderivative of (2)-nitro-(3)-naphthol, the benzoyl group wandering from the hydroxyl to the amidogen; this probably takes place by the intermediate formation of benzenyl-amidonaphthol C10 H3 C.C3 ll3 since this body

occurs in the reduction product.

Acetyl derivative. — C10 II. (NIIAc)OH. [225°]. Plates. Formed by reduction of ths acetyl derivative of (a)-nitro-(B)-naphthol, the same isomeric change taking place as in the preceding case (Böttcher, B. 16, 1935).

Di-amide-(a)-naphthel C10 H10 N2O i.e. C₀H₁(OH)(NH₂). From di-nitro-(α)-naphthol [138°] (Griess a. Martius, A. 134, 376). Its aqueous solution is turned red by Fe Cl, amido-

naphthoquinone imide (q. v.) being formed.
Salts. (Graebe a. Ludwig, A. 154, 307).—
B"H SnCl₁: monoclinic prisms, a:b:c= B"H SnCl₁: monoclinic prisms, a:b:c = 1.184:1:1.187, $\beta = 72^{\circ}33'$. - B"H SnCl₁4aq. -

B"H SO, 2aq.

Tri-amido (a) naphthel Cull N.O i.c. $C_{10}H_4(OH)(NH_2)_n$. From tri-mitro-(a)-naphthol (Dichl a. Merz, B. 11, 1665; Ekstrand, B. 11,

B"'H_SO aq: scales.—B"'3HCl aq: needles. AMIDO (B) NAPHTHOL SULPHONIC ACID. C10H,(OII)(NII,).SO2II. From nitroso-(B)-naph-

thol sulphonic acid (q. v.) with tin and HCl (Meldola, C. J. 39, 47). Long white needles, v. sol. water, turned brown by air. Gives phthalio acid with HNO, Aq. The acid obtained by reducing m-carboxy-benzene - azo - sulpho - (8) naphthol appears to be an isomeride (Griess, B. 14, 2032).

Di-Amido (a)-naphthel-sulphenic acid C₁₀H₄(OH)(NH₂)₂SO₂H [1:2:4:β']. Prepared by reducing naphthol yellow S. Lamine; gives

di-imido-(a)-napathol sulphonic acid when oxidised (Lanterbach, B. 14, 2029)

Amide (β) naphthel di-sulphonic acid C₁₀H₁(OH)(NH₂)(SO₂H) 3nq. From m-carboxybenzene azo sulpho (\$) naphthol by reduction. Laminæ, v. sol. water (Griess, B. 14, 2042).

AMIDO-NAPHTHOQUINONE C1011,NO2 i.e. C₁₀11₂(N11₂)O₂. Oxy - naphtho - quinone imide. Oximido-naphthal. Orange needles, formed by boiling amido-naphthoquinone imids with water (Martins a. Griess, A. 134, 377; Graebo a. Ludwig, A. 154, 307). Sl. sol. boiling water, v. sol. alcohol, insol. ether. Converted by boiling acids or alkalis into oxy-naphthoquinone. Aniline, whon heated with it in acetic acid solution, forms naphthoquinons di-anilide.

Its dihydride is described as AMIDO-NAPHTHO-

MYDROQUINONE.

AMIDO NAPHTHOQUINONE-IMIDE C₁₀H₈N₂O i.e. C₁₀H₃(NH₂)

Di-imido-naphthol. From di-amido-(a)-naphthol and Fc₂Cl₆Aq (Graebe a. Ludwig, A. 154, 307). Minute needles, v. sl. sol. cold water, v. sol. alcohol. Boiling alkalis and dilute acids at 120° convert it into oxy-naphtboquinons. Rsdut,ng agents producs di-amido-(a)-naphthol. An fine heatsd with its hydroebloride forms naph hoquinone di-anilide. Salt.-B'HCl: dark red monoclinic prisms with green lustrs $a:b:c=2.967:1:2.396 \ \beta = 74^{\circ} \ 28' - B'_{2}H_{2}PtCl_{0} - B'H_{2}CrO_{4}: S. \cdot 13 \ at \ 12^{\circ} - B'H_{2}SO_{4}$

Bromine-water added to an aqueous solution of amido-naphthoquinone imide produces two dsrivatives (Kronfeld, B. 17, 715):

1. C_bH₄Br₂O₂ [173°]: whits plates, so alcohol, benzene, HNO₃ (S.G. 1·4), and CHCl₃. 2. C₁₀H_aBr₃NO₃ i.e. CBr₃·CO.C₆H₄·C(NH).CO₂H [213°]: whits needles, sol. alcohol, benzene and HNO, (S.G. 1.4), insol. chloroform. It is split up by cold alkalis into phthalimide and bromo-When heated alone or better with H2SO, at 140° it gives CO2, Br2 and an acid C2H6Br2NO [237°]. This loses HBr when boiled for a long time with alkalis. When heated with H₂SO₄ it givos phthalic acid.

AMIDO - NAPHTHOQUINONE SULPHONIC ACID $C_{10}H_xN_xSO_4$. Minuts coppery needles, formed by oxidising di-amido-naphthol sulphonic

acid.

DI-AMIDO DINAPHTHYL C₁₀H₁₆N₂. Dinaphthyline. C₁₀H₆(NH₂).C₁₀H₆(NH₂). [273°]. Colourless plates (from benzene). Formed together with a smaller quantity of the isomeric naphthidine by warming (aa)-hydrazonaphthalone with two mols. of dilute HCl at 70°-80°; a clear solution is formed from which the naphthidine is ppd. as its sparingly soluble hydrochloride hy adding an excess of HCl, the easily soluble hydrochloride of dinaphthyline remaining in solution.

Reactions.-By diazotising and boiling with alcohol it is converted into $(\alpha\alpha)$ -dinaphthyl [154°]. By boiling with acids NH₃ is easily split

C₁₀H₆NH. off, giving imido-dinaphthyl |

CrO, it is oxidised to phthalic acid.
Salts. — B"H.Cl.*: easily soluble. B"H.Cl.PtCl,: sparingly soluble yellow plates

(Nietzki a. Goll, B. 18, 5254).

Di-amide-dinaphthyl (naphthidine). Probably [4:1] $C_{10}H_6(NH_2).C_{10}H_6(NH_2)$ [1:4]. [198°]. Silvery plates or colourless tables. Soluble in alcohol and benzone. Formed, together with its isomeride dinaphthyline, in small quantity by heating (aa)-hydrazonaphthaleno with HCl, and in large quantity by reduction of (ac)-azonaphthalene with SnCl, and HCl.

Preparation.—One pt. of azonaphthalone is dissolved in 45 pts. of hot acetic acid, and a solution of SnCl₂ (1 pt.) in 2 pts. of HCl and 2-3 pts. of water is added in sufficient quantity to decolouriss it; on adding an excess of HCl tho sparingly coluble hydrochloride of naphthidine is ppd.

Reactions. - Fe₂Cl₂, CrO₂, Cl₂, &c., produce a sarmine red colouration or pp. with solutions of

naphthidine salts. By heating with OrO, it is oxidised to (α)-naphthoquinone and finally to phthalio acid. Its diago-compound yields violet colouring-matters with the sulphonic acids of (8)-naphthol. By boiling the diazo-compound with alcohol it yields (aa)-dinaphthyl [1540].

Salts.—B"H,Cl2: sparingly soluble colourless silvery plates.—B"H Cl2PtCl4.—B"H2SO4: very sparingly soluble glistening platss.

Di-acetyl-a rivative C20H12(NHAc)2 [over 300°]; mearly insoluble in ordinary sol-

vents (Nietzki a. Goll, B. 18, 3254). Di - amido - $(\alpha\alpha)$ - dinaphthyl $C_{20}H_{12}(NH_2)_3$. Obtained hy reduction of di-nitro-dinaphthyl

with zine dust and HCl. It is readily oxidised .NH to the di-imido-compound C20H12 NH, so that it cannot be diazotised. B"H2Cl2: oasily soluble

colourless needles. The bichromate pps. in brown crystals.

C20H12(NHA0)2. Di-acetyl-derivative [Above 300°]. Colourless needles; insoluble in all solvents (Julius, B. 19, 2551).

Twenty-eight di-amido-(aa)-di-naphthyls are indicated by theory.

Tetra amide isc di naphthyl C₂₀H₁₀(NH₂)₄. [164°-167°]. From tetra nitro iso di naphthyl. Grsy powder, sl. sol. alcohol, m. sol. toluene (Staub a. Watson Smith, C. J. 47, 104).

AMIDO-NITRO- v. NITRO-AMIDO-.

AMIDO-NITRO-BROMO- v. BROMO-NITRO-AMIDO-

AMIDO - NITRO - CHLORO - v. CHLORO-NITRO-

AMIDO-NITRO-IODO- v. IODO-NITRO-AMIDO-. AMIDO-OCTOIC ACID C,II,NO, i.e. C.H. CH(NH2).CO2H. Amido - caprilic acid.

From heptoic aldehyde-ammonia (cnanthol-ammonia) and HCN (Erlenmeyer a. Sigel, A. 176, 341). Pearly white lamine, v. sl. sol. alcohol, ether, and cold water. Neutral. Volatilises before fusing. Salts. - HA'HCl. -IIA'HNO; slander needles. — (HA'), H2SO4. —

Nitrile C₆H₁₂.CH(NH₂).CN. [0°]. From HCN and cenanthol-ammonia (E. a. S.). Oil, miscible with alcohol and ether. Salts.—C,H_{1,N},HCl: satiny, six-sided plates.—(C,H_{1,8}N₂),H,PtCl₄.

Amide C₄H₁₃,CH(NH₂),CONH₂. From the

nitrile and conc. HCl. Salts .-

 $(C_aH_{18}N_2O)_2H_2PtCl_6$. $-C_aH_{18}N_2OHCl$. Converted by aqueous NaHCO₃ into an acid, $C_9H_{18}N_2O_3$ (?). AMIDO-OCTYL-BENZENE v. AMIDO-PHENYL-

AMIDO-OCTYL-TOLUENE v. AMIDO-TOLYL-OOTANE

AMIDO - CENANTHYLIC ACID v. AMIDO-HEPTOIC ACID.

AMIDO-OPIANIC ACID v. OPIANIC AOID.

TRI-AMIDO-ORCIN C,H,,N,O, i.e. CoMo(OH)2(NH2)3. From tri-nitro-orcin by sodium-amalgam or by Sn and HCl. Its solutions readily oxidise in air, forming amido-di-imido-orcin (q. v.) (Stenhouse, Pr. 21, 125). AMIDO-OXALOXYL-ACETIC ACID

CO.H.CH(NH.).CO.CO.H. Phenyl hydraside CO.H.CH(NH.).C(N.HPh).CO.H. Obtained by roduction of an alkaline solution of the diphenyl - hydrazide of di - oxy - tartario aoid C.(N.HPh),(CO.H), with sodium amalgam. White pp., very oxidisable, its alkaline solution quickly becoming reddish-violet on contact with the air. By warming with H2SO, it is converted into the anhydride C, H, N,O, which is probably oxy-amido-quinizine-carboxylic acid (q. v.). By further reduction of the alkaline solution with sodium-amalgam di-amido-succinio acid [125°] is formed (Tafel, B. 20, 214).

AMIDO-OXINDOLE v. OXINDOLE. AMIDO-OXY- v. OXY-AMILY-.

AMIDO-PHENANTHRENE v. IV FNANTHRENE. AMIDO-DI-PHENIC ACID C₁₁II₁₁NO₄ i.e. [1:2] CO., II. C., H., C., H., (NH.,). CO., H. [1:5:2]. hydrochloride, H.A"HCl, obtained from nitrodi-phonic acid, forms silvery lamine. distilled with lime it produces p-amido-fluorene, $\mathbf{C}_{13}\mathbf{H}_{1}\mathbf{N}\mathbf{H}_{2}$ [123] (Stasburger, B. 16, 2347).

o-Di-amido-di-phenic acid C,H,N,O, [3:2:1] (CO.JI)(NH.).C.H.3.C.H.3(NII.2)(CO.H) Formed, by intramolecular change, when o-hydrazo-benzoic acid is boiled with HCl (Griess, B. 7, 1609).

m-Di-amido-di-phenic acid C11II12N2O, 12 aq

i.e. [2:4:1] $(CO_2H)(NH_2).C_6H_2.C_6H_3(NH_2)(CO_2H)$ [1:4:2]. Benzilline di-carboxylic acid.

Formation.-1. From the corresponding dinitro-compound (Hummel, A. 193, 128; Struve, B. 10, 75; Schultz, B. 12, 235).—2. From m-hydrazo-benzoic acid by boiling with HCl (G.).

Properties .- Short needles (from water); v. sl. sol. water, alcohol, and ether. It gives benzidino and di-amido-fluorene when distilled with lime.

Salts. — Ag.A"aq. — H₂A"2HCl. H₂A"2HNO₃.—II₂A"IL₂PtCl₆ 2aq. AM1DO-PHENOL.

o-Amido-phenol C.H.NO i.e. C.H.(NII2)(OH) [1:2]. [170°]. S 1.7 at 0° (Körner).

Formation.-1. By reducing a-nitro-phenol (Hofmann, A. 103, 351; Fritzsche, A. 110, 166;

Schmitt a. Cook, K. 3, 62). Properties. - Scales; may be sublimed. Salts. B'HCl; S. 8 at 0°. - B', 11, SO.

Reactions.-1. Oxidised by KaFeCy, forming a dye (C₂₄H_nN₃O₂?) which sublines as pink needles (G. Fischer, J. pr. [2] 19, 319).—2. Nitrous acid forms o-diazo-phenol. - 3. Converted into oxy-quinoline by heating with glyeerin, o-nitro-phenol, and H.SO, (Skraup, B. 15, 893) .- -4. Lactic acid, o-nitro-phenol and H.SO, give oxy-methyl-quinoline (oxy-quinaldine) .- 5. CH. Cl. CO. H gives oxy-phenyl-amidoacetic acid, CO.H.CH..NII.C.H.OH (Vater, J. pr. [2] 29, 286).—6. Cl.CO. Et gives oxyphenyl - carbanic ether, HO.C. II. NII.CO. Et, [85°], which is converted by distillation into alcohol and oxy - methenyl - amido - phenol C₆H₄<N>C.OH, [138°] (Grönvick, Bl. [2] 25, 178). - 7. Polassic xanthate converts the hydrochloride of amido-phenol into sulpho-carbanil C_sH₄ < N CSH, called also thio-carbamido-

phenol and 'oxy-phenyl thio-carbimido' (Kalckhoff, B. 16, 1825; Zincke a. Hebebrand, A. 226, 60).-8. Amido phenol (30 g.) hoated with alcoholio quinone (43 g.) forms hydroquinone; and, on cooling, violet needles of a base C₂₄H₁₈N₁O₄ [250°] separato. This base is soluble in aniline, sparingly soluble in alcohol, benzene, and chloroform. In dilute acids it forms deep red solutions. Boiled with NaOH, ammonia comes off

and o-amido-phenol and other products are formed. The salts of the base have green metallic instre and are easily soluble in alcohol but sparingly soluble in water; much water decomposes them into acid and base.-B"2HCL —B"2HClPtCl₁,—B"₂H₂SO₂, Acetyl derivative C₂₁H₁₆Ac₂N₄O₄, [285°]. Oxidised in glacial aceticacid solution by HNO₃, this gives C₂₂H₁₅N₁O₄ [27] 280°]. Bonzoyl derivative C₂₁ d₄₀Bz₂N₄O₄. [265°]. Reactions. — Nitrous acid converts the base into C₂₁H₁₀N₆O₆, which forms small red needles, [above 290°]. HNO, oxidises it in acetic acid solution to a body which crystallises in yellow needles [260°] (Zincke a. Hebr brand, A. 226, 60).

Acetyl derivative CaH, (OH) (NHAe). [201°]. From its anhydro-derivative by boiling with dilute II.SO, (Ladenburg, B. 9, 1525). According to Morse (B. 11, 232) it can be formed by reducing o-nitro-phenol with Sn and IlOAc (cf. Zincke a. Hebebrand, A. 226, 69). Glittering white plates, v. sol. alcohol and hot water: sol.

KOHAq. Not affected by quinono.

Anhydro derivative $C_aH_4 < N > C.CH_3$.

Ethenyl-o-amido-phenol. [201°]. S.G. 2 1.1365. Formed by heating o-amido-phenol with Ac.O. or its acetyl derivative with P2O5 (I.).

Benzoyl derivative C.H.(NHBz)(OH). [167°]. From the anhydro-compound by boiling with aqueous acids, or from the di-henzoyl derivative by boiling with water and BaCO, (Hübner, A. 210, 387; Bottcher, B. 16, 629). Laminæ, sl. sol. cold water, v. sol. hot water, sol. alcohol. ether, and beuzene.

 $C_{\bullet}H_{\bullet} <_{N}^{O} > C.Ph.$ derivative Anhvdro

Benzenyl · o · amido · phenol. [103°]. (814°). Formed by heating o-amido phenol with BzCl, HOBz, or phthalic anhydride, and distilling the product (L.); or by reducing o-nitro-phenyl benzoate, with Sn and HCl. It is insol. water, v. sol. alcohol. Its salts are unstable, but B',H,PtCl, may be crystallised from alcohol.

Di-benzoyl derivative C₆H₄(NHBz)(OBz). [176°]. From o-amidophenol hydrochloride and BzCl (Hübner, A. 210, 387).

Formyl derivative. Only known in its Anhydro-derivativo C₆H₄<0>CH.

Methenyl-amido-phenol. [31°]. (183°). Formed by heating o-amido-phenol with formic acid.

Phthalyl derivative C_aII₄:(C₂O₂):N.C_aH₄.OII. Oxy Oxy-phenyl-phthalimide. [220°]. From o-amido-phenol and plithalio anliydrido at 220°. Yellowish prisms; converted by heat into CO2 and anhydro-benzoyl-o-amido. phenol, and by boiling Na, CO, Aq into phthaloxyl. amido-phonol, CO2H.C3H4CO.NH.C3H4OH, [223°] (Ladenburg, B. 9, 1528).

Methyl derivative CoH4(NH2).OMe. o-Anisidine (228°). From o-nitro-anisol by reduction (Brünck, Z. 1867, 205; Mühlhäuser, B. 13, 919, A. 207, 235; Herold, B. 15, 1684). Liquid. It acts npon quinone thus:

 $3C_6H_4O_2 + 2C_6H_4(OMe)NH_2 = 2C_6H_4(OH)_2 + C_6H_2O_2(C_6H_4(OMe)NH_2)_2$ The product forms reddish-violet needles (from glacial HOAo and benzene), [230°], and gives a fine blue solution in conc. H.SO. (Zincke, 4. \$26, 68). Salts.—B'HCl: needles (from alcohol); may be sublimed.—B'HB:.—B'H,50.,—B',H2PtCl.,—Chloroacetate (Vater, J. pr. [2] 29, 288). Acetyl derivative.—C,H.(NHAc).OMe. [34°] (204°). Pearly crystals. Sol. hot water. Benzoyl derivativo.—C,H4(NHBz).OMe [59°].

Ethyl derivative C₆H₄(NH₂).OEt. o-Amido-phenetol. (229°). A liquid preparation o-nitro-phenetol (M. Förster, from pr. 129, 344). Reactions—I. Bromine torms a diama a tri-bronno-derivative.—2. Cyanogen chloride passed into an ethereal solution forms ethoxy-phenyl cyanamide, [94°], o-amido-phenetol hydrochloride being ppd.:

 $2C_{c}H_{c}(NH_{c}).OEt + CNCl = C_{c}H_{c}(NH_{c})OEtIICl + C_{c}H_{c}(NH.CN).OEt$

C_aH₁(NH₂)ORtHCH + C_aH₄(NH₂CN).OEt (Berlinerblau, J. pr. [2] 30, 98). Chloroacetate.—C_aH₄(NH₂)(OEt)CH₂Cl.CO₂H (Vater, J. pr. [2] 29, 288).

Ethylene derivative C₂H₁(O.C₆H₁N11₂)₂. [128°]. By reducing the ethylene ether of nitro-phenol with Sn and HCl. Properties.— Trimetric plates (from alcohol or water). Feels greasy. Insoluble in cold water, soluble in alcohol, other, chloroform and benzene. Forms a bluish-black solution with H₂SO₄. Fe₂Cl₆ gives a sepia-brown colour. K₂Cr₂O₇ and HCl give a brownish-red colour (E. Wagner, J. pr. [2] 27, 201). Salte.— B'2HCl2aq. Silver-groy glittering flat needles grouped in tutts (from water). The acctate is extremely soluble. The sulphate is the least soluble salt, it crystallises in pearly plates but, like the oxalate, it readily becomes oxidised. Diacetyl derivative.—[226°]. Needles.

Oxethyl derivative.—
HO.C.H.,O.C.H.,NH., [90°]. Got by reducing BzO.C.II.,O.C.H.,NO., (v. o. NITRO-PHENOL) by Sn and HO! (Weddige, J. pr. 132, 252). Properties.—Colourless plates, slightly soluble in water, readily soluble in alcohol, ether and boiling benzene. A feeble base. Benzoyl derivative BzO.C.,H.,O.C.,H.,NH. [c. 100°]. Got by re-

ducing the benzoyl derivative of the exethyl ether of o.Nitro-phenel (q. v.).

Amido-ethyl derivative NIL,C,lL,O.C,L,NIL, Anhydro-benzoyl derivative

C₂H NII . C C II, [151°] v. o-Nitro-phenol.

m-Amido-phenol C_sII_s(NH₂)(OII) [1:3]. From m-nitro-phenol by reducing with Sn and HCl (Bantlin, B. II, 2106). The free base is extremely unstable. Its hydrochloride, B'IICl, is formed by reducing bromo-m-nitro-phenol (Pfaff, B. 16, 613).

Methyl derivative C.H.(NH₂)(OMe) m.Anisidine (251°). Salt.—BTICI (Pfaif). Ethyl derivative C.H.(NH₂)(OEt). (180°-

Ethyl derivative C₈H_{*}(NH_{**})(OEt). (180°-205°) at 100 mm. From the nitre compound by Sn and HCl (P. Wagner, J. pr. [2] 32, 71). A yellowish liquid, turning red in air. Salts.—B'*,H_SnCl_{*}: platoe.—B'HCl: silky grey needlee; insol. cther, eol. water and alcolol.—B*,H_SO₄I\[3aq.—*B'_4H_2C₂O₄: brownish platos.—B'*,HBr (Staedel, B. 16, 29). Acetyl derivative C₄II*₈(NHAc)(OEt). [97°]. Glittering white platos (from water).

Ethylene derivative C₂H₄(O.C_eH₄NH₂)₂. [135°]. Obtained by reducing the corresponding

nitro-compound. Short prisms (from alcohol). Sl. sol. hot water, sol. hot alcohol and benzene, sl. sol. ether. Forms crystalline salts.

p-Amido-phenic C.H.(NH₂)(OH) [I: 4].[I70°] (S. a. C.). [c. 184°] (L.). S. 1·1 at 0°. S. (alcohol) 4·5 at 0°.

Preparation.—1. From p-nitro-phenol by reducing with iron and acetic acid (Fritzsche, A. 110, 166) or Sn and HCl (Schmitt a. Cook, Kt. 3, 61).—2. If om amido-salicylio acid by distillation (Schmitt, B. 1, 67).—3. From nitrosophenol (Sueyer a. Caro, B. 7, 965).

Properties. Crystalline, but very unstable. May be sublimed (S. a. C.). In capillary tubes it turns brown at 140° and melts at 184° with decomposition (Lossen, A. 175, 296).

decomposition (Lossen, A. 175, 296).
Salts.—B'IICl. S. 71 at 0°. S. (alcohol)
10. Turns brown in air.—B'IIOAc. [183°].

S. II at 0°. S. (alcohol) 8.3 at 0°.

Reactions. 1. Bleaching powder solution added to a cold solution of p-amido-phenol hydrochloride forms quinone chloro-imide, but if added to a hot concentrated solution of the salt it forms a mixture of tri- and tetra-chloroquinones.—2. Bleaching powder added to a solution of p-amido-phenol in fuming HCl at 0° forms pure tri-chloro-quinone .- 3. Chlorine gas passed in oxeess into a solution of p-amidophenol in fuming HCl forms pure tri-chloro-amidophenol (Schmitta. Andresen, J. pr. 131, 435). The observation that chlorino converts p-amidophenol hydrochloride in aqueous solution into quinone-chloro-imide, while in presence of conc. HCl chloring converts p-amido phenol into trichloro-amido-phenol, is explained by the fact that quinone-chloro-imide is changed by cono. HCl into chloro-p-amido-phenols. In this decomposition the first reaction is:

$$C_6H_4 < _{NCl}^{O} + 4HCl = C_6H_4 < _{NH_2,HCl}^{OH} + 2Cl_2.$$

This reaction is similar to that between chloride of nitrogen and IICl:

 $NCl_3 + 4HCl = NH_4Cl + 3Cl_2$

Hence both ammonia and amidogen are protected by couc. HCl from the action of chlorine.

4. Acts on quinone as follows:

3C_aH₁O₂+2NH₂·C_aH₁OH = 2C_bH₁(OH)₂+C_aH₁OH₁C_aH₁OH)₂O₂. It ie beet to use the hydrochloride of p-amido-phenol in the aqueous solution; the product, which separates on cooling, is sparingly soluble in the usual menstrua, and does not melt below 290° (Zincke a. Hebebrand, A. 226, 70).—5. HCl₁NaNO₂, and K₂SO₂ produce p-diazo-phenol sulphite, HO.C_aH₁,N.SO₂H (Reisenegger, A. 221, 316).—6. Cl.CO₂Et forms p-oxy-phonyl-carbamic ether, HO.C_aH₁,NII.CO₂Et [120°].—7. A mixture of IfCl and potassium sulphosyamide produce, when the solution is evaporated, p-oxy-phenyl-thiourea, HO.C_aH₁,NH.CS.NH₂ [168°].—8. HCl and potassium sulphosyamide produce, when the solution is evaporated, p-oxy-phenyl-thiourea, HO.C_aH₁,NH.CS.NH₂ [214°].—9. GS₂ produces di-oxy-di-phenyl-thio-urea, CS(NH.C_aH₁,OH)₂.

Acetyl derivative C_eH₄(NHAo)(OH). [179°]. Large white prisms; obtained by reducing p-nitro phenol with tin and glacial acetic acid (Morse, B. 11, 232).

Benzoyl derivative C.H. (NHBz) (OH). [227°]. From p-amido-phenol and BzOl

(Hübner, A. 210, 878). Needles, insol. water, alcohol, and petroleum, sl. sol. hot alcohol.

Amido-phenyl benzoate C.H.(NH2)(OBz). [154°]. Obtained by reducing p-nitro-phonyl benzoate. Plates; sol. boiling alcohol, and water, v. e. sol. glacial acetic acid.

Di-acetyl derivative C.H. (NHAc)(OAc) [151°]. From p-amido-phenol and Ac2O.

Dibenzoyl derivative C.H. (NIBz) (OBz).
[231°]. From p-amido phenol and BzCl.

Methyt derivative C.W. (NH.) (OMe). Methyt derivative C.W.(NHL)(OMe). p.Anisidine. [56°]. (246°). From p.nitro-anisol (Brunk, Z. 1867, 205; Salkowski, B. 7, 1009). Formed also, together with CO₂, when anisoylhydroxylamine is distilled (Lossen, A. 175, 296). Tables (from water). B'HCl: long needles. B'2112PtCl8.

Ethyl derivative C₆H₄(NH₂).OEt. (253°). Obtained by re (Halloch, B. 14, 37). reducing p-nitro-phenetol

Ethylene derivative C2H4(O.C6H4NH2)2 [0. 170]. By reducing the ethylene ether of p-nitro phenol (q. v.). Properties. - Needles, which turn brown in the air. Crystallises from alcohol or from water. Very soluble in hot benzene, less soluble in CHCl, or ether. Forms a deep blue colour with K₂Cr₂O₇ and HCl. Fe₂Cl₆ gives a cherry red (E. Wagner, J. pr. [2] 27, 206). Salt .- B"2HCl: long thin needles grouped concentrically. The acctate is deliquescent. The sulphate is the least soluble salt. The exalate is but slightly soluble.

(a) Di-amide-phenol O₆Π₃(OH)(NH₂)₂ [1:2:4]. From (a) di-nitro-phenol (Gauhe, A. 147, 66; Stuckenberg, B. 10, 385; Post a. Stuckenberg, A. 205, 56). The free base is extremely unstable.

Salts.-B"2HCl. Precipitated by adding conc. HCl to its aqueous solution. Small prisms insoluble in cone. HCl, and in absolute alcohol. Dilute solutions are turned violet-red by Fe₂Cl₆ or bleaching powder (H. Köhler, J. pr. [2] 29, 270).—B"2H1.—B"H₂SO₄2aq: tables. Di-benzoyl derivative

C_eH_s(NHBz)₂OH (?). [187°]. From the hydrochloride and BzCl. Pale red leaflets; sol. alcohol, chloroform, and aniline, sl. sol. ether, insol. water. Forms a nitro derivative [c. 169°].

Tri-benzoyl derivative $C_6H_3(NHBz)_2(OBz)$ (?). [233°]. Rhombohedra, insol. alcohol, chloroform, and ether; sol. aniline.

(8) Di-amido-phenol $C_0H_2(NH_2)_2OH$ [2:6:1]. The free base is very unstable; its hydrochloride is got by reducing the corresponding di-nitrophenol (Post a. Stuckenberg, A. 205, 79).

Salts.-B"2HCl: thick pointed prisms, v. sol. water, sl. sol. alcohol. B"H.SO.: yellow needles.

DistRzoyl derivative C_eH₃(NHBz)₂OH. [209°-213°]. Minute crystals, sol. aleohol, sl. eol. benzene.

Tri-benzoyl derivative

 $\mathbf{C}_{\mathbf{s}}\mathbf{H}_{\mathbf{z}}(\mathrm{NHBz})(\mathrm{NBz}_{\mathbf{z}})(\mathrm{OH}).$ [184°]. Sol. warm Na₂CO₃Aq, insol. chloroform.

Tetra-benzoyl derivative

C.H.(NBz2)(NBz2)OH. [182°]. Leaflets; insol. water, sol. warm Na CO.Aq, alcohol, benzene, and ether.

Di-amido-phenol C₆H₂(OH)(NH₂)₂. [1:3:4]. Hydrochlorids B"2HOL—Formed heating ethoxy-p-amido-phenyl oarbamic ether, C.H. (OH)(NH.)NH.CO.Et with fuming H((H. Köhler, J. pr. [2] 29, 269). Oblong plate Very soluble in conc. HCl and in absolute alooho Dilute solutions are turned blood-red by Fe₂C or bleaching powder.

Tri-amido-phenol CaH2(NH2),OH. [2:4:6:1] Formation.-1. From pierie acid by reductio (Heintzel, Z. 1867, 338; B. 1, 111; Bamberger B. 66, 2400).—2. From picramide, Sn and HC (Hd.)p., A. 215, 350). The free base is unstable The salts give a blue colour with a large quantit of water containing air. In conc. solution Fe₂Cl gives deep blue glittering orystals of amido-di imido-phenol (Heintzel).

Salts. — B'''3HCl.—B'''23H2SO4.—B'''H3SnCl 1 a. B"HI H.SO. 2aq. B"HIH PO. 2aq. -B"H, FeCy. If the hydrochloride is boile with HCl di-amide-dioxy-benzene is go

(Salkowski, A. 174, 260).

Tri-acetyl derivative C, H, (NHAc), OH [263°]. From the hydrochloride of tri-amido phenol by hoating with NaOAc and Ac.O. White plates, soluble in acetic acid, hot alcohol, water aqueous acids and alkalis, very sparingly it benzene and acetone; by HNO2, CrO3 or Fe2Cl it is oxidised to the tetra-acetyl derivative o tetra-amido-di-oxy-diphonyl-quinone C₆H(NHAc)₂(OH).O

(Bamberger, B. 16, 2400). C₆H(NHAe)₂(OH).O

Reaction .- Tri-amido-phenol hydrochloride is converted by bromine water into 'bromo dichromazin' O18H8N3Br11O7. This body separates from alcohol in yellow needles with feeble violet dichroism. Boiling dilute H,SO converts it into 'bromo-dichroic acid C18H,Br11O11 and ammonia. Bromine converts bromodichromazin into hexa-bromo-acetone (Wedel a. Gruber, B. 10, 1137).

Tetra-amido-phenol

Ethyl-ether, hydroehoride CeH(NH2)4(OEt),2HCl. By reducing the produc of the action of IICl upon tri-nitro-ethoxy-phenyl urethane.—(1). $C_6H(NO_2)_3(OEt)(NH.CO_2Et) + HC$ = $CO_2 + EtCl + C_9H(NO_2)_2(OEt)NH_2$.

(2.) $C_6H(NO_2)_3(OEt)NIL_1 + 9H_2 = C_6H(NH_2)_4OEt + 6H_2O_4$

Properties .- Crystallises from dilute alcohol Insoluble in absolute alcohol, very soluble in water. Does not melt at 360°. Reduces solutions of Au and Pt. A feebly acid solution gives with Fe2Cl6 or bleaching-powder the following succession of colours: dark-green, violet-red, red dish-brown, yellowish-brown, yellow, colourless.
AM1D0-PHENOL SULPHONIC ACID

C_eH₇NSO₄ i.e. C_eH_e(NH₂)(OH)HSO₃. [2:1:4] S. 1 at 14°. Prepared by reduction of o-nitro phenol-sulphonic acid or by sulphonation o o-amido-phenol. Large colourless crystals like calc-spar. Does not form salts.

Anilide Coll_(NH2)(OH)(SO3.NHPh). [2050] Colourless needles; soluble in alcohol, acetic acid

and benzene insoluble in ethor.

Benzoyl derivative C.H. (NHBz)(OH)(SO.H). Salts .- NaA'4 aq: colourless needles, soluble in water a. alcohol.—BaA'; colourless spangles sparingly soluble.—CaA' 242 aq: sparingly soluble colourless scales (Post a. Holst, B. 43, 617; 4 205, 49).

p-Amido-phenel sulphonic said O,H,(OH)(NH,)8O,H [1:4:2]. S. .07 at 140. Formation.—1. From p-amido-phenol hydro-chloride and fuming H₂SO₄ (Post, B. 6, 897).— 2. From p-nitro-phenol culphonic acid (Post a. Holst, B. 13, 617).-3. Together with azoresorufin by heating a mixture of resorcin and nitro-benzene with H₂SO₄ (Brunner a. Krämer, B. 17, 1867) .- 4. From quinone chloro-imide and cone. Na₂SO₂ (Schmitt a. Bsnnewitz, J. pr. [2] 8, 7).

Properties. - White glistening needles; slasol. sold water, v. sl. sol. alcohol, insol. ether. Poes not combine with acids, but forms metallic alts, e.g. Ba(O.C_eH₃(NH₂)SO₃). Reduces cold ammoniacal AgNO₃. Turned violet by Fe₂Cl_e. Not ppd. by lead acetato.

Anilide C.H. (OH) (NH,)SO, NPhH. Small colourless crystals; v. sol. alcohol, acetic

acid, and benzene; insol. ether.
p-Amido-phenol di-sulphonic acid
C_H_2(OH)(NH_2)(SO_H)_2 [1:4:2:6] (?). From benzene-azo-phenol tri-sulphonio acid (q. v.) by ammonium sulphido (Wilsing, A. 215, 236; Limpricht, B. 15, 1293). White silky needles. Deliquescent; sl. sol. alcohol, insol. ether. Solution gives with Fe₂Cl₆ a deep violet colour. 1ts alkaline solutions show blue fluorescence for a short time. Salts.—KHA"aq: slightly sol. cold water.—NH,HA"aq.—PbA"aq.

 $C_{12}H_{11}N$ o-AMIDO-DIPHENYL C.H., C.H., NH₂ [1:2]. [45°]. From o-nitro-diphonyl with tin and glacial acetic acid (Hübner a. Lüddens, A. 209, 351).

Salts.-B'HCl: needles.-B'2H2PtCl64aq.:

orange leaflets.

p-Amido-diphenyl C.H., C.H., NH. [1:4]. Xenylamine; Martylamine. [49°]. (320°). Occurs in the high-boiling fractions in the preparation of aniline (Hofmann, Pr. 12, 389; G. Schultz, A. 174, 212; Osten, B. 7, 171). Prepared by reducing p-nitro-diphenyl with tin and HCl (Hübnor a. Östen, A. 209, 339). Colourless leaflets, sol. hot water, alcohol, and chioroform.

Salts.—B'HCl: leaflets.—B',H,PtCl,2aq; yellow loaflets, sl. sol. alcohol. - B'HNO₃: pearly leaflets. -B', II, SO₄: leaflets, v. sl. sol. water, sl. eol. alcohol.—B'2H2C2O4: long needles, sol. water and alcohol.

Acetyl derivative C.H.C.H.NAeH. [167°]. Long needles; v. sl. sol. water.

Benzoyl derivative C. H. NBzH. [230]. Leaflets; insol. water, v. sl. sol. alcohol.

Formyl derivative C.II. C.H. NH.CHO. [172°]. Prepared by heating p-amido-diphenyl with ethyl formato at 100°. Minute needles; sol. ether, sl. sol. alcohol, v. sl. sol. water (Zimmermann, B. 13, 1967).

p-Amide-diphenyl sulphenic acid $C_{12}H_s(NH_2)SO_3H$. [above 300°]. Formed by sulphonation of p-amido-diphenyl (Carnelley a. Sehlevelman, C. J. 49, 380). Insol. water.

Salts.--NaA'2aq: colourless needlos, m. sol.

water.—BaA'₂4aq; v. sl. sol. water. o-p-Di-amide-diphenyl C₁₂H C12H12N2 [1:2] NH₂.C₆H₄.C₆H₄.NH₂ [1:4]. Iso-benzidine. (β)- or (δ)-Di-amido-di-phenyl.
 (45°).
 (363°)
 (Schultz, A. 207, 348). Diphenyline.

Formation.—1. From o.nitro-p.amido-di-phenyl (Schultz, B. 9, 548; 14, 612).—2. From di-amido-diphenyl carboxylic acid (Strasser a.

Schultz, A. 210, 193).

Preparation.—An alcoholic solution of azo-

benzene (100 g.) is heated with a solution of SnCl, in cono. HCl; the liquid is evaporated to dryness, the residue diesolved in water and benzidine is ppd. as sulphate (100 g.) while o p-di-amido-diphenyl eulphate (30 g.) remains in solution (Sohmidt a. Schultz, B. 12, 482).

Properties.—Long needles, v. sl. sol. water. Salts.—B"HCl: laminæ.—B"2HCl: needles. -B" H_2SO_4 : prisms.— $B''_2H_2SO_4$.

Di-acetyl derivative C12H10Ac2N2. [2020].

m-m-Di-amidgediphsnyl [1:3] NH, C, H, C, H, NH, [1:3]. From the nitro compound (Brunner a. Witt, B. 20, 1028). Crys-

tals, sl. sol. water.—B"H2SO4.—B"H2PtCl4.
Di-acetyl derivative. [258°]. needles.

p-p-di-Amido-diphenyl. Benziline. [1:4] NH₂.C₆H₄.C₆H₄.NH₂ [1:4]. Xenyi $Xenylene \cdot di \cdot$

amine. [122°]. (above 360°).

Formation.—1. By the reduction of azobenzene or of azoxybenzene in alcoholic solution by SO₂ (Zinin, A. 85, 328).-2. From azobenzeno by SnCl. and HCl (v. sup.).—3. By heating azobenzene with fuming HCl (Zinin, A. 137, 376), HBr (Werigo, A. 165, 202), or H1 (Senziuk, Z. 1870, 267) in sealed tubes.— 4. By reducing nitro-benzene with sodium-amalgam in presence of acetic acid, the product being treated with H₂SO₄ (Werigo, A. 135, 176).— 5. From nitro-benzene, alcoholic NaOII, and zinc dust and subsequent treatment with acid (Aloxejeff, Z. 1867, 497).—6. From di-amido-diphenic acid by distilling with BaO (Schultz, A. 196, 29).—7. From p-amido p-nitro-diphenyl by Sn and HCl (Fittig, A. 124, 276).—8. From p-p-di-nitro-phenyl by tin and HCl (Schultz, A. 174, 227).-9. From hydrazo-benzene by treatment with mineral acids:

C₆H₅.NH.NH.C₆H₅ = NH₂.C₆H₄.C₆H₄.NH₂. Preparation.—V. o-p-di-amido-diphenyl.

Properties.—Silvery scales; may be sublimed. Sol. hot water, v. sl. sol. cold water, v. e. sol. alcohol and ether.

Salts.—B"H₂SO₄: small ecales, v. sl. sol. water and alcohol.—B"21ICl: lamina, v. sol. water and alcohol.—B"11Cl: long needles, sl. sol. water; ppd. when a large excess of water is added to the preceding salt.—B"2HNO,: foursided laming, sol. hot water. — B",H,C,O,: groups of silky needles, m. sol. water and alcohol.—B"C,H₀O₆: laminæ; sol. water.

Reactions.—1. Even very dilute solutions

give with polassic bichromate a deep blue pp. (Julius, M. 4, 193).—2. K_xFeCy_z gives a blue pp. 3. Chlorine-water gives a blue colour soon becoming red.-4. Exhaustive chlorination with SbCl, gives per-chloro-diplienyl and per-chlorobenzene (Merz a. Weith, B. 16, 2874).-5. If very dilute bromine-water be poured up 3.8 solution of benzidine in CS, the upper layer becomes blue, excess of bromine destroys this colour, the lower layer then turning red (Claus a. Risler, B. 14, 83).

Acetyl derivative NH2.C4H4.C4H4.NAcH. [199°]. Needles, sl. sol. water.

Diacetyl derivative

NHAc.C.H.C.H.NAcH. [3170]. Nearly insoluble in all solvents.

Di-formyl derivative C12Ha(NHaCOH),: crystalline powder, sublimable, insoluble in all ordinary solvents except nitrobenzens. Formed by heating hydrazobensene or benzidine with formic scid.

Di-bensoyl derivative CuHs(NHBz)2: solourless needles or pearly plates; insol. alcohol, ether, and aniline; sol. nitrobenzene. Formed by heating hydrazobenzene or benzidine

with BzCl (Stern, B. 17, 379).
Di-phthalyl derivative C., H1. N. O. [above 360°]: silky yellow needles, sol. hot nitro-benzene; insol. most other solvents. Formed by heating benzidine & hydrazobenzeno with phthalic anhydride (Buildrowski, B. 17, 1181).

Oxalyl derivative (C,H,NH),C,O,. insoluble powder, obtained by heating benzidine

oxalate at 200°.

Benzidine-v-sulphonic acid

H2N.CaH2.CsH2.NH.SOaH. Formed by heating an alcoholic solution of azobenzene with hydric ammonium sulphite (Spiegel, B. 18, I48I).

Gelatinous pp. It gives colourless crystalline calts.

The HSO, is readily split off with production of benzidine by dissolving the acid in strong H2SO.

Benzidine sulphenic acid (?)

C,H,(NH,).C,H,(NH,)(SO,H)(?). Hydrazo-ben-zene sulphonic acid C,H,NH.NH.C,H,SO,H (?). Hydrazo-ben-Ppd. by adding HCl to the product of the action of H.S on an ammoniacal solution of azo-benzene sulphonic acid (Griess, A. 154, 213).—Yellow needles or plates (from water).—BaA'₂: plates. The free acid is decomposed by solution in aqueons NH, into benzidino and H2SO4. The above azobenzene sulphonic acid is converted by potash fusion into p-oxy-azobenzene, and would therefore appear to be a p-sulphonic acid; in which case it is not clear how the conversion into a benzidine derivative could be effected. If, however, the acid is hydrazo-benzene sulphonic acid, we must assume that the benzidine transformation here takes place in alkaline solution, by displacement of SO₂H. In any case tho removal of SO₂H by ammenia is peculiar.

Benzidine sulphonic acid $\mathbf{C}_{12}\mathbf{H}_{11}\mathbf{N}_{2}(\mathbf{SO}_{3}\mathbf{H})$ $2\sqrt{2}$ aq. Obtained by heating (a) benzidine disulphonic acid with water at 210° (Limpricht, B. 11, 1048). Yellow needles (from alcohol); v. sol. water.—KA'4aq.—Ba\Lambda'_24aq.— PbA', 3aq.

Chloride C12H11N2(SO2CI) [above 210°].

Benzidine disulphonic acid $[4:8:I]C_bH_3(NH_2)(SO_3H).C_6H_3(SO_3H)(NH_2)[I:3:4]$

8. 08 at 22°. From azo-, or azoxy-, benzene disulphonic acid by reduction with SnCl., sodiumamalgam, or NaOH and zinc dust, followed by treatment with a mineral acid (Mährenholtz a. Gilbert, A. 30, 337; Brunnemann, A. 202, 314; Limpricht, B. 14, 1359). Monoclinic prisms (with 3aq). Dilute HCl at 230° gives benzidine and H.SO. Nitrous acid diazotises this acid.-Na₂A" δ]aq.— K_2 A" $1\frac{1}{2}$ aq.— $Ca\Lambda$ "4aq.— $Ba\Lambda$ "4aq.— $Pb\Lambda$ "4aq.

Benzidine di-sulphonic acid

C₁₂H₄(NH₁) (SO₂H)₂. From benzidine and fuming H₂SO₄ at 170°. Small white plates; v. sl. sol. water, insol. alcohol and ether (Griess, B. 14, 300). Salts. — BaA"5aq: white plates. — BaA"2aq: needles.—*Ag₂A": white crystalline ₽P.

Bensidine (a)-di-sulphonic scid C₁₂H₁₈N₂(SO₁H)₂. Hydrazo-benzene di-sutpnom acid (?). Prepared by reducing potassium azo benzene (a)-di-sulphonate with SnCl, (Limpricht B. 14, 1357). Tables (containing 2aq); sl. sol cold water, v. sl. sol. alcohol. — K_*A'' 8aq. — BaA''aq.— $\times Ag_*A''$: white pp.—PbA'': needles

sl. sol. cold, v. sol. hot, water.

Benzidine tetrasulphonic acid $C_{12}I_{\bullet}N_2(SO_3H)_{\bullet}$. Prepared by sulphonation of the freeeding with fuming H_*SO_{\bullet} — $Ba_2\Lambda''''I4aq$: large prisms, v. sol. hot water, sl. sol. alcohol.—
*K,A''' (Limpricht, B. 14, 1543).

Other sulphonic acids of benzidine

Benzidine heated with a large excess of fuming II SO, above 170° forms a mixture of di-, tri-, and tetra-, sulphonic acids, and di-amido-diphenylene sulphono sulphonic acids (Grioss, B. 18, Ref. 88).

Benzidine di-carboxylic acid v. di-Amido-di-

PHENIC ACID.

Benzidiae tetra-carboxylic anhydride

 $0 < {}^{CO}_{CO} > C_e H_2(N H_2).C_e H_2(N H_2) < {}^{CO}_{CO} > 0.$

[above 360°]. Formed by the action of an HCl solution of SnCl, on azo-benzene tetra-carboxylic acid (azo-phthalic acid). Light-yellow tasteless powder. Insol.water, alcohol, ether, or dilute acids.

With alkalis it gives anhydride salts: -C12II8N2(C2O3)(CO2K)25aq: large prisms. $C_{12}H_8N_2(C_2O_3)(CO_2Na)_2$ aq: small needles. $C_{12}H_8N_2(C_2O_3)(CO_2Ag)_2$: fine powder. $\begin{array}{l} O_{12}H_6N_2(C_2O_3)(CO_2)_2l^3b: \text{ amorphous powder,} \\ C_{12}H_6N_2(C_2O_3)(CO_2NH_4)(CO_2II): \text{ transparent} \end{array}$

prisms (Claus a. Hemmann, B. 16, 1759).

Di-amide-diphenyl C12H12N2. Iso-benzidine. [125°]. Occurs among the products obtained by passing aniline through a red-hot tube (Bernthsen, B. 19, 420). White iridescent plates, sl. sol. water. Its aqueous solution gives no colouration with potassic ferricyanide, and a greyish-brown pp. with chlorine water. The solid base is turned greenish-black by strong HNO,. The sulphate is sparingly soluble.

Tetra-amido-diphenyl [3:4:I] (NII₁)₂C₈II₃.C₆II₃(NH₂)₂ [1:3:4]. Obtained by reducing di-nitro-p-p-di-amido-diphenyl (Brunner a. Witt, B. 20, 1025). Silvery plates.

a AMIDO-PHENYL-ACETIC ACID C.H.NO. i.e. C.H. CH(NH.).CO.H. Phenyl-amido-acetio acid. [256°]. Formed by heating a bromophenyl-acctic acid with NH3Aq (S.G. 9) at 100° (Stockenius, B. 11, 2002); or by saponifying its nitrile, obtained by the action of alcoholic NH, on the cyanlydrin of benzoic aldehyde (Ticmann a. Friedlander, B. 14, 1967). White leaflets or prisms; may be sublimed. Sl. sol. cold water, in. sol. hot water. It forms unstable salts with bases, but more stable salts with acids, though these are decomposed by water. Distilled with lime, it gives benzylamine (Tiemann, B. 13, 383).

Salts. -B'HCl: trimetric prisms. -B'HNOs. $B'H_{\nu}SO_{\nu} - B'H_{\nu}C_{\nu}O_{\nu} - AgA'$: prisms, v. sl. sol. water.— BaA'₂: small white plates; v. sol. hot water.—MgA'₂; aq: plates, sl. sol. water.

Amide. The hydrochloride forms thick

prisms, sl. sol. alcohol.

Nitrice C.H.CH(NH.).CN. Yellow oil (v. sup.). m. Sulphonic acid

C.H. (SO.H).CH(NH2).CO.H. Minute needles; #1

sol. cold water, insol. ether (Plochl a. Los, B. 18, 1182).

o- Amido-phenyl-acetic acid. When o-nitrophenyl-acetic acid is reduced the product is not o-amido-phenyl-acetic acid bnt its anhydride, oxindol (q. v.) (Baeyer, B. 11, 583).

m-Amido-phenyl-acetic acid

 $\mathbf{C_6H_4}(NH_2).C\dot{H}_2.C\dot{O}_2H$ [1:3]. [148°]. Formed by reducing m-nitro-phenyl-acetic acid (Gabriel a. Bergmann, B. 16, 2065).

Nitrile C.H. (NH2). CH2. CN. m. Amido-benzyl cyanide. A liquid obtained by reducing m-nitro - phenyl - acetonitrile (Salkowski, 17, 506).

p-Amido-phenyl-acetic acid C_aH₄(NH₂).CH₂.CO₂H [1:4]. [[200°]. From pnitro - phenyl - acetic acid (Radziszewski, B. 2, 209; Bedson, C. J. 37, 92). White needles

(from water); v. sl. sol. cold water.

Nitrile C, H, (NH2). CH2. CN. n - Amido benzyl cyanide. [46]. (312°). V.D. 4.78 (for 4.56). Formation.—1. From p-nitro-benzyl cyanide (Szumpelik, B. 3, 474; Gabriel, B. 15, 831).— 2. As one of the products of the reduction of ap-di-nitro-cinnamie ether $(q.\ v.)$ by tin and HCl (Friedländer a. Mähly, $A.\ 229,\ 229)$. The yield is 15 p.c. of the substance used. Properties .-Satiny plates (from water). Sol. acids. Gives a di-bromo-derivative. HCl at 130° converts it into amido-phenyl-acetie acid. Salts .-B'_H_PtCl_a.—B'_H_SO. Acetyl derivative NHAc.C₀II. CHL.CN. [97°]. Slender needles; v. sol. alcohol and other. Di-acetyl derivativo NAc₂.C₈H₁.CH₂CN. [153°]. Glistening needles. Sol. boiling water, benzene, and CS₂; sl. sol. aleohol.

· Di-amido-phenyl-acetic acid C₆H₃(NH₂), CH₂GO₄H [4:3:1]. Formed by reducing (3, 4, 1)-nitro-amido-phenyl-acetic acid (Gabriel, B. 15, 1996). Short flat crystals (with aq). Sl. sol. hot alcohol, insol. ether, CS, chloroform, and benzene. Sol. acids and alkalis.

a.m.Di-amido-phenyl-acstic acid

[1:3] C₂II₄(NH₂).CH(NH₂).CO₂H. [214°]. Formed by reducing m-nitro-phenyl-α-amido-acetic acid with tin and HCl (Plochi a. Loë, B. 18, 1181). Flat silvery necdles.

Salt .- * CuA', : hluish-green crystalline pp.

o-AMIDO-PHENYL-ACETYLENE C,II,N i.e. C.H.(NH.).C:CH. Yellowish oil. Prepared by reduction of o-nitro-phenyl-acetylene with zinedust and NH3. It forms yellow pps. with ammoniacal AgNO, and Cu2Cl2.

B'HCl: soluble yellow crystals.

Reaction .- Converted by II, SO, (12 pts.) and H.O (4 pts.) into o-amido-acetophenone (Baeyer a. Bloem, B. 17, 964).

Acetyl derivative. [75°]. needles (Baeyer a. Landsberg, B. 15, 60).

Di-o-amido-di-phenyl-diacetylene [128°]. $\mathbf{C_{e}H_{4}(NH_{2}).C;C.C;\bar{C}.C_{e}\dot{H}_{4}(NH_{2}).}$ Prepared by the action of a solution of potassium ferricyanide on the cuprous compound of o-amidophenyl-acetylene. Long yellowish needles. Sol. alcohol, ether and acids, insol. water.

B"H.Cl. • colourless soluble crystals. Di-acetyl derivative. [231°]. Long needles (Baeyer a. Landsberg, B. 15, 60).

(B.2)-AMIDO-(A)-PHENYL-ACRIDINE

)C,H,(NH2). C10H14N, i.e. C4H4

Formed by heating phenyl-p-phenylene-diamine C,H,(NH,).NHC,H, with benzoic acid and ZnCl2 (Hess a. Bernthsen, B. 18, 692). Amorphous solid. Easily soluble in ordinary solvents. The solutions of the base are yellow, the benzene and ethereal solution having a splendid green It dyes silk a brownish yellow. fluorescence. The solutions of its salts are red.

Di-amido-phenyl-acridine v. Chrysaniline. AMIDO-PHENYL-ALANINE v. DI-AMIDO-PHENYL-PROPIONIC ACID.

AMIDO - PHENYL - AMIDO . U. DI-AMIDO.

PHENYL- OI PHENYL-DI-AMINO-

AMIDO-DI-PHENYL-AMINE C₁₂H₁₂N₂ i.e. NH₂-C₆H₁,NH.C₆H₅. [61°]. Prepared by the reduction of nitro-di-phenyl-amine or of phenylamido-benzene-azo-benzene, or its sulphonic acid

(Tropacolin O O.) (Nietzki a. Witt, B. 12, 1399).
Thin lamina. Gives quinone on oxidation.
Salt. B'₂H₂SO₄: silvery lamine, sl. sol. water. Acetyl derivative C12H11AcN2. [158°].

p.p. Di-amido-di-phenyl-amine NH_r.C_eH_,NH.C_eH_,NH_r. [158°]. Formed by reduction of anilino black. Prepared by roducing (a)-di-nitro-di-phenyl-amine (N. a. W.).
Acetyl derivative. [239°].

Di-amido-di-phenyl-amine. Prepared by redueing (β)-di-nitro-di-phenyl-amine. Liquid.

Salts.-B"H.Cl,; sl. sol. water. - B"H,PtCl, Acetyl derivative. [203°].

Tri amido tri phenyl amino (C,H,NH,),N. [230°]. Formed by the reduction of tri-nitrotri-phenyl-amine by SnCl2 (Heydrick, B. 18,

2157; 19, 759.) Salts.—B"3HCl: needles. Its solution exhibits the following colour reactions: blue, turning violet with Fe.Cl.; bluish-green with ppd. MnO.; blue with K.Cr.O.; red with chloranil in acctic acid (but if in this case the free base is used the colour is bluish-green). -B"23H2PtCla.

B"(C₆H₂(NO₂),OH)₃.

Tri-acetyl derivative N(C,H,NHAc)₃:

needles which do not melt below 240

AMIDO-PHENYL-BENZGLYCOCYAMINE v. AMIDO-DI-PHENYL QUANIDINE CARBOXYLIC ACID. p-AMIDO-PHENYL-iso-BUTANE C₁₀H₁₅N i.e.

C, II, C, H, NH, Butyl phenylamine. Amido. butyl-benzene. (230°). S.G. 25 937. From aniline hydrochloride (10 g.) and iso-butyl alcohol (8 g.) by heating for 6 hours at 230' (A. Studer, A. 211, 237; B. 14, 1172, 2186; Pahl, B. 17, 1232). Colourless oil; v. sl. sol. water, volatile with steam. Miscible with alcohol or ether. Nitrous acid converts it into butyl-phenol

Salts.-B'HCl.-B'HBr. -B'HL. Acetyl derivative [170°]: lamina Formyl derivative C₁₀H₁₃, NH.CHO [59°]: laminæ (Gasierowski a. Merz, B.18, 1009).

AMIDO - PHENYL - BUTINENE C, HIN CH probably CaH4(NH2).CII2.CII

(272° at 718 mm.). V.D. = 4.95 (for 5.02). Formed by reduction of m-nitro-a-methyl-cinnamic aldehyde in alcoholic solution with tin and HCl. Colourless glistening plates. Sublimes at 100°. Reduces ammoniacal AgNO,

hydrochloride, sulphate and nitrate are easily soluble in water. The hydrochloride forms colourless glistening plates. B'2H2Cl2PtCl, 2aq: slender needles.

Acetyl derivative C10H10NAc [140°]:

colourless concentrio prisms.

Benzylidene derivative C10H,N:CHPh [73°]: concentric light-yellow needles. Formed by heating the base with benzaldehyde (Miller a. Kinkelin, B. 19, 1249) o-AMIDO-PHENYL-CARRAMIC ETHER

C₈H₁₂N₂O₂ i.e. H₂N.C₆H₄NH.CO.Et. o-Amidophenyl-wrethane. [86°]. Formed by reducing o-nitro-phenyl-carbamic ether (Rudolph, B. 12, 1295). Long colourless needles; sol. water.

Salt .- B'HCl: large tables.

p-Amide-phenyl-carbamic ether. Amidoearbanilic acid. [74°]. Formed by reducing p-nitro-phenyl-carbamio ether (Hager, B. 17, 2626; Behrend, A. 233, 10). Needles (from dilnte alcohol); insol. water.

Salts.—B'HCl: long needles.—B'H.SO. B'H2C2O4: needles, sol. hot water, sl. sol. cold water.—B'2H2PtCls: brown pp. - (B'HCl)3SnCl. B'2SnCl2 aq. - (B'HCl) HgCl2.

Benzoyl derivative CoH. (NHB2).NH.CO.Et, [230°]: needles; sl. sol. alcohol, insol. water.

Di-p-amido-di-phenyl-carbamic ether (C.H.,NH.),N.CO,Et. Di-p-amido-di-phenyl-amine urethane. [101°]. Formed by reduction of di-p-nitro-di-phenyl-carbamic ether. Violet (C,H,NH),N.CO,Et. noedles (+aq). Soluble in water.

Di-benzoyl derivative

(C₆H₄.NHBz)₂N.CO₂Et [235°]: nearly colourless amorphous solid (Hager, B. 18, 2576)

DÎ-AMIDO-DÎ-PHENYL-CARBINOL C₁₈H₁₄N₂O i.e. C₈H₄(NH₂).CH(OH).C₆H₄NH₂. (β)-Di-amido-benzhydrol. [128°-129°]. From (3)-di-amido-benzophenone [149°] and sodium amalgam (W. Staedel, A. 218, 350). Glittering plates. Salts.—B"2HCl2aq.—B"H2SO, 2aq.

Acetyl-derivative. [220°], Di-amido-tri-phenyl-carbinol C19H18N2O i.e.

C_sH_sC(OH)(C_cH₄NH₂)₂ [below 100°].

Formation. — By the action of anilino in presence of H2SO4 upon the chloride C.H.CCI.C.H.NH.

Preparation.—From aniline hydrochleride (40 pts.), nitrobenzene (45 pts.), benzo-trichloride (40 pts.), and Fe at 180°. C₆H₃CCl₂ + 2C₆H₃NH₂ = C₆H₃CCl(C₆H₄NH₁), + 211Cl. The mass is extracted with dilute HCl (which leaves some blue colouring matters undissolved) and the nitro-benzene is distilled off by steam (Doebner, B. 15, 234; A. 217, 242).

Properties. - Small crystals (from dilute alcohol). Insol. in cold water; v. sol. alcohol or benzera On heating with MeI it gives

malachite green.

Salts .- Dilute acids dissolve it in the cold, forming nearly colourless solutions which on boiling (split off water and) change to deep reddishviolet. The salts dye violet, but the shades are not €ast. The coloured salts are probably of the

form C.H.C.C.H.NH2 This salt forms dark ∕\CI

blue needlos with coppery lustre.

Reaction .- Zino dust and HCl reduce it to di-amido-tri-phenyl-methane (q. v.).

Tri-amide-tri-phenyl-carbinels v. Rosannans. DI - AMIDO - TRI - PHENYL - CARBINOL CABBOXYLIC ANHYDRIDE O20H16N2O2 6.6. (O2H1NH2)2CO6H1.CO.O. [2650-2660]. Small

colourless needles. Is prepared by heating phenolphthalein with aquoous NH3. Gives a tetra-bromo-derivative [280°], and a tetra-acetyl-tetra-bromo-derivative [241°] (Baeyer a. Burkhardt, B. 11, 1297).

AMIDO-DIPHÉNYL CARBOXYLIC ACID v.

Amido-diphenio acid.

DI - AMIDO - DI - PHENYLENE KETONE OXIDE (so called) $C_{19}H_{16}N_{2}O_{2}$ i.e. $C_{13}H_{0}(NH_{2})_{2}O_{2}$. Lactone of oxy-di-amido-diphenylcarboxylic acid. From the nitro compound by Sn and HCl (A. G. Perkin, C. J. 43, 190). Orange needles (from xylene). Vory slightly soluble in boiling water. Ruby prisms (from diluto alcohol).

Salts. - Forms two hydrochlorides. -

(B"HCl), PtCl, -B"(2HCl)PtCl,

AMIDO-PHENYLENE OXIDE CaH, NO i.e. $C_nH_u(NH_u)O$ (?). Di-amido-di-phenylene dioxide. From nitro-phenylene oxide by alcoholic ammonium sulphide (Märker, A. 124, 251). Yellow needles, sl. sol. water, v. sol. hot alcohol.-B',H,PtCl,

DI-AMIDO-DIPHENYLENE-QUINOXALINE

 $N.C.O_{i}H_{i}$ N.C.C₆II, $C_6H_2(NH_2)_2$ Formed by the action

of phenanthraquinone upon tetra-amido-benzene [1:2:4:5]. Orange-yellow neodles. Nearly insol. acetio acid. Weak base. Dissolves in conc. H2SO4 with a greenish-blue colour, passing through violet into red on dilution (Nietzki a. Hagenbach, B. 20, 338)

AMIDO-PHENYLENE UREA C.H.N.O i.e. $OC < NH > C_e H_s(NH_z)$ [1:2:4]. Formed by reducing di-nitro-phenyl-urothano with tin and HCl (Hager, B. 17, 2631).

Salts. — B"H,SnCl,: long needles. — B"C,H,(NO₂),0H: greenish-yellow needles. o-AMIDO-PHENYL-ETHANE C,H,1N i.e.

 $C_sH_1(NII_2).CH_2.CH_1$ [1:2]. o-Ethyl-phenylamine. o - Amido - ethyl - benzene. (211°). S.G. 22 983. From o-nitro-phenyl-ethane, tin, and HCl (Beilstein a. Kuhlberg, A. 156, 206). Liquid at -10°. Salt .- B'HNO3

Acetyl derivative C₈H₁₀AcN. (305°).

Benzoyl derivative C, H10BzN. [147°]: small glittering plates (Paucksch, B. 17, 2800).

o-Amido-phenyl-ethane sulphonic acid C_aH₃Et(NH₂).SO₃H. Formed by sulphonation of the acetyl derivative. White needles (P.).

p-Amido-phenyl-ethane C.H.(NH.).CH.2.CH. [1:4]. p-Ethyl-phenyl-amine. Phenethylamine. [-5°]. (214°). S.G. 22 975. From p-nitrophenyl-ethane by reduction (B. a. K.) or from aniline by heating with cthyl alcohol and ZnCl₂ (Benz, B. 15, 1647). Formed also when ethyl-aniline hydrochlorido is heated at 800° (Hofmann, B. 7, 526). Colourless oil; volatile with steam. Salts.-*B'HNO; small needles or prisms, sl. sol. cold, v. sol. hot, water .-B',H,SO,: large white plates, sl. sol. cold water, m. sol. dilute H,SO,.—B'HCl.—B',H,PtCl,.

Acetyl derivative C.H.(NHAc).C.H.

[95°]. (816°).

[151°]: long nsedles (P.).

w-Amido-phenyl-ethane C.H. CH2.CH2.NH2.

[198°]. Phenylethyl-amine.

Formation.—1. By dry distillation of a smido-phenyl-propionic acid (q.v.) (Schulze s. Barbieri, J. pr. [2] 27, 346; Erlenmoyer a. Lipp, A. 219, 202).—2. By action of zino and HGl upon the cyanhydrin of benzoio aldehyde, or upon amygdalin (Fileti, B. 12, 297, 1700).—3. 'y action of bromine on an alkaline solution of phenyl-propionamide (Hofmann, B. 18 2740).

Preparation. - By reducing an alcoholic solution of bonzyl cyanide with zinc and HCl (Bernthsen, A. 184, 290), di-phenylothyl-amine (C_sH_s.CH_s.CH_s),NH, and tri-phenylethyl-amine (C_sH_sCH_sCH_s),N, being also formed (Spica, G.

1875, 124; 1879, 566).

Properties. - Liquid; sl. sol. water. Absorbs CO, from the air, being converted into a solid carbonato [105°], out of which, on heating, another carbonate, [88°], sublimes. Oxidised to benzoio aoid by chromio mixture.

Salts.—B'HCl, [217°]: trimotrio tablets (from cold alcohol) or satiny plates (from alcohol-ether): v. sol. alcohol or water, insol. ether.—B'2H2PtCl; more soluble in hot water than in hot alcohol.

Di-amide-di-phsnyl-sthans v. DI-AMIDO-DI-

ω-AMIDO-TRI-PHENYL-ETHANE C20H18N i.e. CPh₂.CH₂.NH₂. [116°]. From tri-phenyl-acetonitrile by reduction with zinc and HCl. Crystals; v. sol. ether, sl. sol. cold alcohol. The hydrochloride forms nesdles, [247°], v. sl. sol. water, v. sol. alcohol (Elbs, B. 17, 700).

AMIDO PHENYL-ETHYLENE AMIDO-STYRENE.

Di-p-amide-di-phenyl-sthylans C14H14N2 i.e. C₂H₂(C₆H₄,NH₂)₂. Di-amido-stilbene. [227°]. Formation.—1. By reduction of di-p-nitro-

di-phenyl-cthyleno with tin and HCl.-2. By reduction with SnCl, of the brownish-red product of condensation (azoxy-di-phenyl-ethylene?) obtained by the action of sodium methylate or alcoholic NaOH upon p.nitro-toluene.

Reactions.-By nitrous acid it is converted into a tetrazo-compound which by combination with the sulphonic acids of amines and phenols yields a series of colouring-matters which dye cotton from a soap bath. Thus (a)-naphtholsulphonic acid gives a bluish-violet, (3) naplithol. (R) di-sulphonic acid a blue, (a) naphthylaminesulphonic acid a red, and salicylic acid a yellow, colouring-matter.

Di-acetyl derivative C2H2(C6H4.NHA0)2: [312°] (Bender a. Schultz, B. 19, 3234).

D1-p-amide-di-phsnyl-sthylens-di-oarbexylic C,H,(NH2).C.CO

| > 0. [280°]. Formed anhydrids

C₆H₄(NH₂).C.CO by reduction of the nitro compound (Reimer, B. 14, 1802). Small plates. Insol. most solvents.

Di-p-amido-di-phenyl-sthylsne - di sulphonic acid C₂H₂(C₆H₃(NH₂)SO₃H)₂. Di-amido-stilbene-di-sulphonic-acid. Obtained by reduction with zinc dust of the brown product (azoxy- or azodi-phenyl-ethylene-di-sulphonic acid?) which is formed by boiling p-nitro-tolucne-o-sulphonic acid C₆H₂Me(NH₂)SO₅H [1:4:2:] with aqueous NaOH. Microscopic needles. Nearly insol. Vor. I.

Bensoyl derivative C.H. (NHBz)C.H. water. Its salts are easily soluble. By nitrous acid it is converted into a tetrazo-compound which by combination with amines or phenols gives colouring-mattere which have the property of dyeing cotton from a soap bath (Bender a. Schultz, B. 19, 3234).

o-AMIDO-PHENYL-GLYOXYLIC ACID

ISATIO ACID.

m-Amide-phanyl-glyaxylle soid C,H,NO, i.e C.H.(NH.)CO.CO.H [1:3]. Colourless prisms or needles. [270°-280° with decomposition]. Prepared by reduction of m-nitro-phenyl-glyoxylio acid with alkaline FeSO,

Salts .- A'Ag: sparingly soluble crystalline powder.—C,H,NO,HCl: soluble flat prisms

(Claisen a. Thompson, B. 12, 1946).

p-AMIDO-3-DI-PHENYL-GUANIDINE m-GARBOXYLIC ACID C₁,H₁,N₁O₂ i.e. [1:4] NH₂,C₃H₄,NH₄,C(NH),NH₄,CO₂H₄.

Amido-phenyl-benzglycocyamine. Prepared by heating cyano-carbimido-amido-benzoio acid (v. p. 157) with p-phenylene-diamine (Griess, B. 16, 338). Small prisms. Salt.—B"H₂Cl₂. 16, 338).

p-amide-s-di-phenyl-guanidins o-carboxylic acid. Anhydride H2N.C8H4.NH.C NH.C8H4 Amido-phenyl-benzglycocyamidine. Formed by boiling di-cyano-amido-benzoyl (v. p. 155) with an aqueous solution of p-phonylene-diamine (Griess, B. 18, 2421). Very small white needles: v. sol. hot water, m. sol. alcohol.

AMIDO-PHENYL-HYDROXIDE v. AMIDO.

m-AMIDO-PHENYL-HYDRAZINE CaH, Na i.o. C_uH₄(NH₂).NH.NH₂ Formed by saponification of the examic acid C_cH₄(NH.C₂O₂.OH).NH.NH₂ which is obtained by reduction of m-diazophenyl-examic acid C_eH₄(NH.C₂O₂.OH).N₂Cl with SnCl₂ (Griess, B. 18, 964). V. sol. aloohol and ether, sl. sol. water. Very oxidisable.

Amido-phenyl-hydrazins sulphonic acid $C_6H_3(NH_2)(N_2H_3)(SO_3H)$ [3:1:6]. Formed by reduction of nitro-phenyl-hydrazino sulphonic acid with NH,HS or SnCl₂ (Limpricht, B. 18, 2194). Very soluble in water. Salts:—A'H,HCl casily soluble fine white needles .- A'HH, SO .: microscopic needles.— *A'H,HNO; prisms. AMIDO-DI-PHENYL-KETONE v. AM

BENZOPHENCHE.

o-AM1DO-PHENYL-MERCAPTAN

C₆H₇NS i.c. C₆H₄(NH₂)(SH). Amido phenyl.

sulphydrate. [26°]. (234°).

Formation.—1. By fusing benzenyl-amidophenyl-mercaptan (q. v.) with potash (Hofmann, B. 12, 2363). -2. Anhydro-oxalyl-amido-phenylmercaptan (easily prepared from acctanilide and sulphur) is fused with potash (3 pts.). The yield is nearly theoretica! (Hofmann, B. 13, 1230). Colourless needles, very easily oxidised. It forms products of condensation with acids, aldehydes, and nitriles; thus acctic acid, acetyl chloride, acetonitrile, and aldehyde each produes ethenyl-amido-phenyl mercaptan (q. 8.) $C_{\bullet}H_{\bullet} <_{S}^{N} > C_{\bullet}CH_{\bullet}$

p-Amido-diphanyl-mercaptan

[1:4] H.N.C.H.C.H.SH [1:4]. Prepared by reducing p-nitro diphenyl sulphochloride with tin and HCl (Gabriel a. Damberger, B. 13, 1410).

Salt. B'HCl: small glittering prisms.

AMIDO-PHENYL-METHANE v. Tolumbia m-Amido-di-phenyl-methane O, H, N i.e. C.H. CH. C.H. NH. [46°]. Formed by reducing m-nitro-di-phenyl-methane (Beeker, B. 15, 2092).

Acetyl derivative. [91°]: pearly plates. p-Amido-di-phenyl-msthane. [35°]. Formed by reducing p-nitro-di-phenyl methane with tin and HCl (Basler, B. 16, 2718). The sulphate is sl. sol. cold water.

di-amide-di-phanyl-mathana C15H10(NH2)2 [85]. Formed by reducing (a)-di-nitro-di-phenylmethano (Doer, B. 5, 795). Pearly plates; sl. sol. water (Practorius, A. 194, 348). sulphate is v. sl. sol. water.

Tstra-amide-di-phsnyl-methane C13H2(NH2) .. [161°]. By reduction of the nitro compound [172°] (Staedel, A. 218, 341). White needles (from benzene). M. sol. water, sl. sol. bonzene.

Acctyl derivative C13H4(NHAc)4. Crystalline powder. V. sl. sol. water; m. sol. alcohol. Amide-tri-phenyl-methane C₁₉H₁₂N i.e.

CHPh. C.H. NH. [84°]. Prepared by heating aniline hydrochloride with di-phenyl-carbinol and ZnCl, at 180° (Fischer a. Roser, B. 13, 674; Is a weak A. 206, 155). Prisms or plates. Is a weak hase. The benzene compound (C₁₉H₁₇NC₈H₆) forms long colourless needles. Salts. B'HCl: needles, sl. sol. water.—B'H_PtCl₄.
Di-amide-tri-phsnyl-methane C₁₄H₁₈N₂ i.e.

C.H., CH(C.H.NH₂)₂. [139°].
Formation.—1. From benzylidene chloride, aniline, and zinc dust (Böttinger, B. 12, 976).-2. From di-amido-tri-phenyl-carbinol by reducing with zinc dust (Doebner, A. 217, 246; B. 15, 236) .- 3. By heating aniline hydrochloride with benzoic aldehyde and fuming HCl (Mazzara, G. 14, 510).

Preparation .- A mixture of benzaldchyde (10 pts.), aniline sulphate (28 pts.), ZnCl2 (20 pts.) and a little water, is heated on a water bath for several hours, the fused mass is boiled with dilute H,SO, diluted, filtered, and the base precipitated with NH, j yield 80 p.c. of the

theoretical (Fischer, B. 15, 676).

Properties.— Colourless crystals (from other) [139°]. Prisms containing benzene of crystalhisation (from benzene) [106°]; at 120° the benzene goes off. V. sl. sol. water, sol. alcohol or ether.

Additional References .- C. Böttinger, B. 11, 276, 840; 13, 958; O. Fischer, A. 206, 147, 153;

B. 13, 665.

Tri-amide-tri-phenyl-msthane C19H16Ns i.e. CH(C₈H₄.NH₂[1:4])₈. Para-leucaniline. [148°].

Formation.—1. By reducing tri-nitro-tri-phenyl-methane with zine dust and glacial acetic acid (O. a. E. Fischer, A. 194, 272).-2. Burn uoing para-rosaniline (Hofmann, Pr. 12, 9) .- 3. By reducing nitro-di-amido-tri-phenylmethane, prepared from aniline hydrochloride, p-nitro-benzoic aldeliyde and ZnCl2 (Fischer a.

Greiff, B. 13, 670; Fischer, B. 15, 678).

Properties.—Colourless plates. Readily converted by exidation into para-Resanitine (q. v.).

Salts.-B"H,Cl,aq: short prisms, sl. sol. alcohol, ether, and HClAq.-The sulphate forms needles, v. sol. water, sl. sol. alcohol, incol. ether.—The oxalate forms prisms, v. sol. water .- The platinoohloride forms sparingly soluble short needles.

Tri-acetyl derivative [177°].

tables; when oxidised by K, Or, O, and sootie acid it gives tetra-acetyl-para-rosaniline.

Tri-benzoyl derivative [1497]. Colourless needlss; sol. alcohol, v. sl. sol. watsr, ether,

and benzene (Renouf, B. 16, 1301).

m-p-p-Tri-amide-tri-phenyl-methane
[1:3] H.N.O.H., CH(C.H., NH,[1:4]). Pseudo-leucanitine. [150°]. Obtained by reducing mnitro-di-p-amido-tri-phenyl-methano (Fischer, B. 13, 673). Colourless crystals; sol. alcohol, sl. sol. ether or benzoline. Crystallises with benzene in white needles of $C_{19}H_{19}N_3C_4H_4$ [145°]. on oxidation, a violot colouring matter.

Salts .- B2"3H2PtCla: yollow erystalline pp.,

v. sol. water, m. sol. alcohol.

o-p-p-tri-amido-tri-phenyl-methans [165°]. [1:2] H₂N.C₆H₄.CH(C₆H₄NH₂[1:4])₂. Formed by reduction of the o-nitro-di-p-amidotri-phenyl-methane obtained by heating o-nitrobenzoic aldehyde with aniline sulphate and ZnCl. Small crystals. On oxidation it gives a brown colouring matter.

Salts.-B"H,Cl,: colourless easily soluble needles. The sulphato forms small quadratic tables, v. sol. water, sl. sol. alcohol. The oxalate forms small soluble needles (Renouf,

B. 16, 1304).

AMIDO-PHENYL METHYL KETONE v. AMIDO-ACETOPHENONE.

p-AMIDO-DI-PHENYL-METHYL-PYRAZOL-CARBOXYLIC ACID C,,H,,N,O, i.e.

∴NPh

CaHa(NH2).C.C(CO2H):CMe [251°]. Formed by reduction of p-nitro-di-phenyl-methyl-pyrazolcarboxylic acid with SnCl2 (Knorr a. Jödicke, B. 18, 2259). Crystalline powder. Sol. alcohol, ether, acids, and alkalis, insol. water. It evolves CO, at its melting-point.

m-AMIDO-(Py.3)-PHENYL-(Py.2)-METHYL C11:CMe QUINCLINE

C, H, N, i.e. C,H N : C.C.H.(NH.)

[115]. Obtained by reduction of m-nitro-phenylmethyl-quinoline with tin and HCl (Miller a. Kinkelin, B. 19, 533). Prisms. Very soluble in alcohol and benzene, tolerably in ether. Has no dyeing power, although it is isomeric with flavaniline. By further reduction with tin and HCl it yields a tetrahydride.

Salts.-The mono-acid salts are yellow, the di-acid colourless .- B"H,Cl, 2aq: easily soluble glistening prisms, B"H,Cl,PtCl, 2aq: orange tables.—B"H Cl,PtCl,: concentric yellow plates.

m-Amide - (Py. 3) - phenyl - (Py. 2) - msthyl-CH₂.CHMe tstrahydro-quineline C,H,(NH,)

Formed by reduction of amido-phenyl-msthylquinoline with tin and HCl. Dracetyl derivative O16H16N2Ac. [178°].

Thin colourless prisms; sol. hot alcohol p-Amide - (Py. 3) - phanyl - (Py. 1) - methyl-

quineline v. FLAVANILINE. AMIDO - PHENYL - HYDRO - QUINOLINE v.

AMIDO-PHENYL-QUINOLINE. p-AMIDO-PHENYL-OCTANE C, Han i.e. H.N.O.H., C,H., Capryl-phenyl-amine. Phen-capryl-amine. (291° corr.). Formed by hoszing a mixture of aniline, capryl alcohol, and ZnOl, at 2000 280°. Or by heating aniline hydrochloride and

Fluid at -20°. Colourisss oil.
Salts.—B'H2SO,: v. sol. hot water, v. sl. sol. $cold. - B'_2H_2O_2O_4$: small platss, v. sol. alcohol and hot water, sl. sol. cold water.

Benzoyl derivative C14H21.NHBz. [109°]. Slendsr falted nasdles, v. sol. alcohol and ether when hot, sl. sol. when cold.

o-Amido-phenyl-octans. From the nitro-compound (Ahrens, B. 19, 2725).--B'₂H₂SnCl₆. p-Amide w-phenyl-n-ectane. [19.5°]. (311° cor.). From n-octyl alcohol, aniline, and ZnCl. (B.).—B'HCl.—B'₂H₂SO₄.—B'₂H₂C₂O₄. Formyl derivative. [56°].

Acetyl derivative. [93°]. Benzoyl derivative. [117°].

AMIDO-OPIANIC ACID C10H11NO C₆H(OMe)₂(NH₂)(CHO)(CO₂H) [6:5:3:2:1]. Di-From methoxy-amido-aldehydo-benzoic acid. nitroso-opianic acid, SnCl2, and HCl. Crystalline. Balt.-HA'HCl: needles, decomposed by water.

Reactions .- 1. Baryta-water give a blue-violet colour.—2. Fe Cl, gives a green colour in solutions of NH, A'.—3. Hot Ac, O gives granules of C24H24N2O11 [233°] (Kleemann, B. 20, 876).

DI-AMIDO-DI-PHENYL OXIDE C,2H12N2O i.e. (C_aH₁.NH₂)₂O. [185°]. From the nitro compound (Hoffmeister, A. 159, 208). The sulphate forms slender needles.

AMIDO-PHENYL-PENTANE v. AMIDO-AMYL-

BENZENE.

TRI - AMIDO - TRI - PHENYL - PHOSPHINE **OXIDE** C₁₆H₁₈N₃PO i.e. OP(C₆H₄.NH₂)₃. [259°]. Obtained by reduction of tri-nitro-tri-phenylphosphine oxide (Michaelis a. Soden, B. 17, 923). White prisms. Soluble in hot water, hot alcohol, and acctone, sparingly in cold water, cold alcohol, and other. Its salts are very soluble in water. Tri-acetyl derivative

OP(C,H,NHAc),aq. [1882], colourless crystals. Tribensoyl derivative OP(C,H,NHBz),

[o. 180°], crystalline powder.

DI - AMIDO - DIPHENYL - PHTHALIDE $C = (C_6H_4NH_2)_2$ O20H16N2O2 i.e. CaH4 \c≤ŏ

Lactone of di-amido-tri-phenyl-carbinol carbo-xylic acid. [180°]. Tables. Prepared by roduction of dinitro-diphenyl-phthalide. By the action of HNO₂ it gives phonol-phthalein (Baeyer, B. 12, 642; A. 202, 66).

AMIDO PHENYL PIPERIDINE C,1H18N2 i.e. C,NH10 C,H.NH2. [40°]. Formed by reduction of the corresponding nitro-compound [105°] with SnCl, and HCl.—B"H,Cl,aq: large colourless orystals (Lellmann, B. 20, 681)

p-AMIDO-8-PHENYL-PROPANE C, H, N.O. H, N.O. H, CH, CH, CH, Mido-propyl-benzene.

Propyl-phenyl-amine. Phenpropylamine. (225°). From aniline, ZnCl, and propyl alcohol at 270° Liquid, volatile with steam; v. sl. sol. water. Salts: B'HCl: lamina, [204].—B'_LlptCl_—B'HB. [218]—B'HB. [218]—B'HL—B'_LlptCl_—A'HB. [218]—B'HL—B'_LlptCl_—B'HL—B'_LlptCl_—B'HL—B'LlptCl_—B'HL—B'LlptCl_—B'HL—B'LlptCl_

Bensoyl derivative C.H. BzN. [115°]. p-Amido-a-phenyl-propane H.N.C.H., CH(CH), CH,. Amido - isopropylbensons (217°). Similarly prepared from iso-

capryl alcohol at 200°-290° (Bsran, B. 18, 189). propyl alcohol (L.). Liquid, al. sol. water. Fluid at -20°. Colouriss oil. Salts: B'₂H₂SO₄: sl. sol. cold water. -B'₂H₂O₄o₄. Benzoyl derivative [1150]: lamine.

See also CUMIDINE and PHENYL-PROPYL-AMINE.

o - AMIDO - PHENYL - PROPIOLIC ACID

CoH, NO, i.e. CoH4(NH2).C:C.CO2H.

Preparation.—An ammoniacal solution of o-nitro-phenyl-propiolic acid is slowly added to a cold solution of FeSO, (11 pts.) saturated with NH3; after 1 or 2 hours' standing the mixture is filtered and the amido-acid ppd. from the filtrate by adding HCl in slight excess; yield: 65 p.c. of the nitro-acid used (Richter, B. 16, 679).

Properties. - Microscopic needles. Soluble in alcohol, sparingly in other, nearly insoluble in water, benzene, chloroform, and ligroine. Dissolves in aqueous acids. Decomposes on heating to about 125° and on boiling with water, in the latter case forming o-amido-acetophonone and CO.. By boiling with NaOH and then adding HCl a splendid red colour is produced.

Salt .- A'Ag*: insoluble pp.

Ethyl ether A'Et-[55°] needles (Baeyer a.

Bloem, B. 15, 2147).

a-AMIDO-a-PHENYL-PROPIONIC ACID C₀H₁₁NO₂ i.e. CH₂.C(C₀H₃)(NH₂).CO₂H. Anidohydro-atropic acid. From the nitrile by treatment with HCl (Tiemann a. Köhler, B. 14, 1981). Feather-like, satiny, needles. Sublimes about 260°. V. e. sol. water, insol. alcohol and ether. Converted by nitrous acid into atrolactic acid.

Nitrile CH3.CPh.(NH2).CN. Yellow oil. B-Amide-α-phenyl-prepienic acid

CH2(NH2).CH(C6H5).CO2H. [169.5°]. A product of action of cone. NH_aAq on β-bromo-α-phonyl-propionic acid (Fittig a. Wurster, A. 195, 158; Merling, A. 209, 11). Plates (from water). Sl. sol. cold water.

o.Amido-a-phenyl-propionic acid. Anhydride or lactam C₆H₄ NH CO.[119°]. Atroxindol. Formed, instead of the acid, by reducing C₀H₄(NO₂).CHMe.CO₂H. Needles (from dilute HCl). Sl. sol. cold water, forming a neutral solution; sol. alcohol and ether. When quite pure it has a pleasant smell. Slightly volatile with steam. It dissolves in alkalis but is reppd. by

CO, (Trinius, A. 227, 274).
p-Amido-a-phenyl-prepienic acid CH, CH(C,H,NH,),CO,H. [128°]. From nitro-hydro-atropic acid, Sn and HCl (Trinius, A. 227,267). Salt.—HA'HCl; needles; v. sol. water. α-Amide-β-phenyl-propionic acid C₂H₁₁NO₂ i.e. C₆H₂.CH₂.CH(NH₂)CO₂H.

Amido-hydro-cinnamic acid. Occurrence. - In the radicles of germinating lupin seeds, together with other amida acids. Forms about 1 p.c. of the dry seed. The mixture is heated with cupric hydrate and filtered, the acid is isolated from the residue by treatment with H2S and subsequent evaporation (Schulze a. Barbieri, J. pr. [2] 27, 342; B. 14, 1785).

Formation.—From its nitrile by HCl (Erler meyer a. Lipp, A. 219, 194). The acid formed in this way is perhaps not identical with that in lupin seeds. An amido-phenyl-propionic acid identical with that in lupin seeds occurs among ths products of the decomposition of proteids by HCl. It melts at [275°-280°] and is optically active, while the acid from phsnyl-acetic aldehyde melts at [263°-265°] and is inactive (Schulze |

a, Nägeli, H. 11, 201).

Properties.-Glittoring, anhydrous, (from hot saturated solutions); groups of slender needles, containing aq (from dilute solutions) (S. a. B.). Satiny plates (from alcohol); short anhydrous prisms or stars (from water) (E. a. L.). Sweet taste. Neutral reaction. M. sol. water, v. sl. sol. alcohol, insol. ether. Small quantities may be sublimed. Gives no colour with Millon's reagent.

Salts.—CuA'2: insol. water (S. a. B.). CuA', 2aq: rosettes of small blue prisms (E. a. L.). -AgA'. - HA'HCl: prisms or stars; v. sol. alcohol or water, insol. conc. HCl.

(HA'HCl)₂PtCl₄.—HA'HNO₃.—(IIA')₂II₂SO₄.

Reactions .- 1. With K.Cr.O, and H.SO, gives off odour of benzoic aldehydo, and ultimately forms benzoio acid.-2. When heated it cakes together and at about 270° it melts giving off CO_2 , H_2O , and exo-amido-phenyl-ethane (q. v.); the residue may be crystallised from alcohol, it melts at [280°] and has the formula C₈H₉NO. From the behaviour of the analogous amidopropionic acid, we may suppose this body to be phenyl-lactimide. - 3. By putrefaction it gives phenyl acetic acid.

Nitrile CaHa.CH2.CH(NH2).CN. From tho compound of HCN with phenyl-acetic aldehyde by heating with alcoholic NH, at 100° (Erlenmeyer a. Lipp, A. 219, 189). Small crystals.

Salts. B'HCl: trimetric prisms; v. sol. alcohol, insol. ether, v. e. sol. water.

Anhydride or lactam C.H.NO or $O_{14}H_{18}N_2O_2$ i.e. Ph.CH₂.CH $<_{CO}^{NH}>$ or

Ph.CH₂.CH<^{NH.CO}_{CO.NH}>CH.CH₂Ph.

Phenyl-lactimide. [291°]. A by-product in the conversion of the acid into amido-phenylethane by the action of heat. Very stender silky needles (from alcohol) forming an electric powder. May be sublimed as woolly needles. V. sl. sol. water, IICl, or KOHAq; insol. ether; al. sol. glacial acetic acid.

β-Amido-β-phenyl-propicnic acid

C.H., CH(NH2).CIL, CO.II.

β-Amido-hydro-cinnamic acid. [121°]. From β-bromo-β-phenyl-propionic acid and cone. aqueous NII, at 0° (Posen, A. 195, 144; 200, 97). Large monoclinic crystals (from water); m. sol. cold water, v. sol. alcohol, v. sl. sol. ether. Boiling IIClAq splits it up into NII, and cinnamic acid. Salt .- B'HCl; prisms, v. sol. water.

Anhydride or lactam

 $C_{e}H_{s}.CH < CH_{o}NH$ >CO. Phenyl-lactimide. [147°].

Formed, instead of a sulphate, when the acid is added to H.SO, (1 vol.) diluted with water (1 vol.). Needles, insol. cold water, m. sol. hot water, alcohol, or other. Is not reconverted into the amido acid by prolonged boiling with water.

o-Amido-β-phenyl-propionic acid Anhydride or lactam C, H, NO i.e.

 $C_{\bullet}H_{\bullet} < NH \cdot CO > \text{ or } C_{\bullet}H_{\bullet} < NCOH_{\bullet} > \cdot H_{\bullet}CH_{\bullet}CH_{\bullet}CH_{\bullet} > \cdot H_{\bullet}dro-carbostyril. Di-hydro \cdot (Py. 3) - oxy - ox$ quinoline. [160°]. Formed, instead of the amide acid, when o-nitro-B-phenyl-propionio acid is reduced by tin and HCl (Glaser a. Buchanan, Z.

1869, 194). Prisms; v. sl. sol. water v. sol. alcohol, ether, and warm conc. HClAq. May be distilled. PCl, at 140° converts it into di-ohloroquinoline.

Ethyl derivative C.H.N(OEt). [1992]. Formed by reducing the ethyl derivative of carbostyril with sodium amalgam (Friedländer a. Ostermayer, B. 15, 335). Silvery plates.

m-Amido-S-phenyl-propionio acid C.H. (NH.).CH., CH., CO.H. m - Amido - hydro - cinnamic acid. [85°]. Formed by reduction of m-nitro-\$-phenyl-propionic acid with tin and HCl (Gabriel, B. 15, 846). Colourless crystsls. V. sol. water, alcohol, and ether. Salt.-A'HHCl: colourless needles or scales.

p-Amido-\$-phenyl-propionic aoid

C₄H₄(NH₂).CH₂.CH₂.CO₂H. p-Amido-hydro-cinnamic acid. [131°] (Glaser a. Buchanan, Z. 1869, 195). Prepared by reduction of p-nitro-phenyl-propionic acid with FeSO₄ and NH₂. Salts.—B'HCl.—B'₂H₂SO₄.

Acetyl derivative C₆H₄(NHAc)C₂H₄,CO₂H. [143°]. Long colourless needles or short prisms. Sol. alcohol and ether, insol. CS2 (Gabriel, B. 15, 843).

α-β-di-amido-β-phenyl-propionio acid. Anhydride or lactam

C₆H₃.CH.CH(NH₂).CO NII -

a-Benzoyl derivative C.H., CH.CH(NHBz).CO [187°]. Formed by NII -

heating benzoyl-imido-cinnamic acid,

C,H,.CH.CH.CO,H

, with strong aqueous NH3. ŇΒz Glistening needles or prisms; sol. hot aloohol

and acctic acid, sl. sol. ether, insol. water, dilute acids and alkalis. By boiling with HCl it loses NH3 giving the benzoyl derivative of a-amidoeinnamic acid (Plöchl, B. 17, 1616).

a-p-di-amido-β-phenyl-propionic acid C₀H₁₂N₂O₂aq i.e. C₀H₁(NH₂)CH₂. CH(NH₂)CO₂Haq [245°-250°]. p-Amido-phenyl-alanine. Got by reducing p-nitro-a-amido-phenyl-propionio acid (Erlenmeyer a. Lipp, A. 219, 219), or by reducing a-p-di-nitro-einnamic ether and saponifying tho product (Friedländer a. Mühly, B. 16, 852; A. 229, 226). Silky needles (from water), sl. sol. alcohol, insol. ether. Neutral; has a sweet tastc. Reduces salts of gold and silver. Does not give Hoffmann's mercury reaction (A. 87, 124). Gives off no NII, when boiled with KOH. Converted by nitrous acid into tyrosine.

Salts. - IIA'2IICl. - 11A'H2PtCl - CuA'2; small violet-blue crystals, sl. sol. water .-HA'ILSO,: small needles.

(4,2,1) - di - amido - β - phsnyl - propionic soid (4:2:1), C_aH₃(NH₂)₂.CIL₂.CH₂.CO₂H.

Anhydride or lactam C.H. N.O i.s. $H_2N.\dot{C}_0H_3 < \begin{array}{c} CH_2.CH_2 \\ NH.CO \end{array} > .$ Amido - hydro -

carbostyril. (Py. 3)-oxy-(B. 3)-amido-di-hydro-quinoline. [211°]. Formed by reducing (4,2,1)di-nitro-B-phenyl-propionio acid (Gabriel a. Zimmermann, B. 12, 602). Needles or prisms.

Not affected by boiling alkalis. Salt.—BHCl. (£:3:1)-Di-amido-β-phenyl-prepionio acid. [4:3:1] C_aH₁(NH_a)₂.CH₂.CH₂.CO₂H. Di-amido-hydro-cinnamic acid. [144°, dry]. Formed by reduction of m-nitro-p-amido-phenyl-propionic soid with tin and HCl (Gabriel, B. 15, 2291). Thick orystals containing aq. Sol. alcohol and acetic acid, v. sl. sol. ether, chloroform, benzene, and CS2. Dissolves in aqueous acids and alkalis.

m-AMIDO-(Py. 3)-PHENYL-QUINOLINE CH:CH

C18H12N2 i.e. C6H4 [120°]. N : C.C.H.(NH2)

Formed by reduction of m-nitro-phenyl-quinoline with tin and HCl (Miller a. Kinkelin, B. 18, 1904). Long glistening needles. Distils at a high temperature undecomposed. Sol. ether, benzene, and hot water, v. sl. sol. cold water.

Salts .- B"H2Cl2: easily soluble colourless needles.—B"H₂Cl₂PtCl₄: yellow crystalline powder.—B"₄H₂Cl₂PtCl₄: long fine needles.— B"H2SO 2aq: thick colourless prisms.

(a) - Amido - (Py. 1) - phenyl - quinoline

C₁₈H₁₂N₂ i.e. C₈H₄C(C₈H₄·NH₂):CH N=CH [150°].

Obtained by reduction of the corresponding nitrocompound [187°] with SnCl₂. Colourless glistening plates (from alcohol). V. e. eol. alcohol, benzene, and chloroform, v. sl. sol. ether with a bluish-violet fluorescence. Volatilises undecomposed. Its mono-acid salts have a dccp yellow colour and dye wool yellow, the di-acid salts are colourless.- *B"HI: soluble yellow needles.-*B"2H2Cl2PtCl1: yellow prismatio needles. Tho ehromate is a sparingly soluble brown pp. (Koenigs a. Nef, B. 20, 627).

(β) - Amido - (Py. 1) - phenyl - quinoline C(C₆H, NH₂):CH $C_{15}H_{12}N_2$ i.e. C_5H_4 [198°].

Obtained by reduction of the corresponding nitrocompound [118°] with SnCl2. Four-sided prisms. Sl. sol. alcohol and benzene, v. sol. chloroform, v. sl. sol. ether. The ethereal solution has a bluish-violet fluorescenco. It volatilises undecomposed. Its mono-acid salts have a yellow oolour and dye wool yellow, the di-acid salts are colourless.— *B"H,Cl,PtCl,: yellow prisms, sol. HCl, nearly insol. water (Keenigs a. Nef, B. 20, 628).

Amido-phenyl-quinoline. [136.5°]. V.D. 7.67 (for 7.62). Obtained by heating quinoline hydroohlorido with aniliuo (Jellinck, M. 7, 351). Yellowish white needles; insol, cold water, sol, benzone, alcohol, and chloroform.

Salts.-B"2HCl.-B"H,PtCl. Methylo-iodide. B"Mel. [220°] m-Amido-(Py. 3)-phenyl-hydrequinoline

CH_CH2 Thick syrup. NH.CII.C₈H₄(NH₂)

Formed by reduction of m-amido-phenyl-quinoline or of m-nitro-phenyl-hydroquinolino with tin and HCl (Miller a. Kinkelin, B. 18, 1907).-B"H2Cl2: monoclinio tables.

(Py. 1:2)-AMIDO-PHENYL-ISOQUINOLINE C(NH₂):CPh C1.H12N2 i.e. C4H4 [o. 100°]. CH:N

Formed by reduction of (Py. 4:1:2)-chloro-nitrophenyl-isoquiaoline by heating with H and P (Gabriel, B. 19, 834). Yellowish plates or needles. Easily coluble in ordinary solvents, moderately in other and ligroine. Dissolves

readily in acids. Salts .- B'HI: ysllow orystals.—B'₂H₂Cl₂PtCl₄: long orange red needles.
—B'HCl*: flat needles.

DI-AMIDO-DI-PHENYL SULPHIDE

12H12N2S i.e. (CaH4NH2)2S. Thioaniline. [105°]. Mol. w. 216. Obtained by heating anilino (6 pts.) with sulphur (1 pt.) at 160°, with gradual addition of PbO (Merz a. Weith, B. 4, 384); or from di-phenyl sulphide by nitration and reduction (Krafft, P. 7, 384). A small quantity is got by the action of S_cCl₂ on aniline (Schmidt, B. 11, 1168). Long thin needles (from water). V. sl. sol. cold water, sl. sol. hot water, v. sol. alcohol, ether, and hot benzene. Not attacked by hot cono. IICl, by hot alcoholic KOH, or by sodium-amalgam. Its solutione give a blue colour when warmed with Fo₂Cl₄.

Conc. H₂SO₄ dissolves it with violet colour. Salts.—B"H₂Cl₂2aq: prisms, v. sol. water, sl. sol. alcohol or cold conc. HCl. B"HCl2aq. B"H2PtCl4. — B"H2SO4aq. — B"2H2SO4aq. — B"H2C2O4.

Diacetyl derivative (C,H,NAcH),S.

[215°]. Needles.

Di-o-amido-di-phenyl di-sulphide C12H12N2S4 i.c. (C₆H₄NH₋)₂S_x, [93⁵]. Prepared by oxidising o-amido-phenyl mercaptan with Fc₂Cl₆ (Hofmann, B. 12, 2363). Plates; insol. water, sol. alcohol. Readily reduced back to the mercaptan. The hydrochloride forms lamina, sl. sol. HClAq.

Di-p-amide-di-phenyl di-salphide

(C₀H₄NH₂)₇S₂. [79°]. Formed by saponifying its acetyl derivative with dilute H₂SO₄. Long greenish needles (from water); v. sl. sol. water, v. sol. alcohol.

Salts .- B"H2SO, 2aq: small needles.

Di-acetyl derivative (C.H.NAcII)282. [c. 217°]. Formed, together with the discetyl derivative of di-amido-di-phenyl tri-sulphide, by heating acetanilide with S₂Cl₂ at 100° (Schmidt, B. 11, 1171).

Di-amide - di - phenyl tri - snlphide. acetyl derivative (C_cH₄NAcII)₂S₃. [214°]. Prepared as described above; forms lamine (from glacial acetic acid).

AMIDO-DI-PHENYL SULPHONE

C₁₂H₁₁NSO₂ i.e. C₆H₅.SO₂.C₆H₄.NH₂. Amido-sulphobenzide. From nitro-di-phonyl sulphone by alcoholic ammonium sulphide (Gericke, A. 100, 209). Minute prisms, sl. sol. cold water.

Salts .- B'IICl: [c. 90]; reddish four-sided

prisms.-B',11,PtCl,

Di-amide-di-phenyl sulphone C12H12N2SO2 i. (C₈H₄NH₂)₂SO_. [168°] (Schmid a. Nölting, B. 9, 80). Obtained in the same way from di-nitrodi-phenyl sulphone. Four-sided prisms, sl. sol. cold water.

Salts .- B"H2Cl2: long prisms .- B"H2Ttois. Di-amido-di-phenyl-sulphone di-carboxylio acid C_{1,}H₁₂N₂SO₆ t.c. SO₂(C₆H₂(NH₂)CO₂H)₂, [above 350°]. Obtained from p-amido-benzoic acid and furning IL₂SO₄ at 180° (Michael a. Norton, B. 10, 580). Roso-red tufts of orystells (from water), sl. sol. alcohol.

Salt .- Ag2A": small white lamine.

AMIDO-DI-PHENYL SULPHONIC ACIDS o. AMIDO-DIPHENYL.

AMIDO-PHENYL SULPHYDRATE v. AMIDO-PHENYL MERCAPTAN.

AMIDO-DIPHENYL DI-SULPHYDRATE C₁₃H₁₁NS₂ i.e. C₁₂H₁(NH₂)(SH)₃. (153°]. Prepared by reducing nitro-diphenyl di-sulpho-chloride with tin and HCl (Gabriel a. Dambergis,

B. 13, 1411). Long needles.

p-AMIDO-DIPHENYL. p-THIO-GLYCOLLIC ACID C_{1.}H₁₃NSO₂ i.e. H₂N.C_cH_{1.}C₆H_{1.}S.CH₂CO₂H₁ [Over200°]. Formed by action of a chloro-acetate npon p-amidodiphenyl p-sulphydrate (Gabriel a. Dambergis, B. 13, 1410). Plates; sl. stl. water. o-AMIDO-s-DI-PHENYL-THIO-UREA

C13H13N3S i.e. C0H3NH.CS.NH.C6H1.NH2. From o-phenylene-diamine and phenyl thio-carbimide in benzene (Lellmann a. Würthner, A. 228, 212). Glittering prisms, v. sol. alcohol and glacial acetie acid, sl. sol. benzeno, insol. ether. In a capillary tube it cakes together at I4I°; at 185° aniline distils out of it; but at 250° it is still solid: o-phenyleno-thio-urea being left: PhHN.CS.NH. C_6H_1 NH₂=

CS CNH C.H. + PhNH2.

m-Amido-di-phenyl-thio-nrea. [148°]. From m-phenylene-diamine and phenyl thio-carbi-mide in benzene (L. a. W.). Amorphous yellow powder or colourless prisms (from alcohol). V. sol. glacial acetic acid, m. sol. alcohol, insol. ether and benzenc. May be melted without decomposition. Decomposed by long boiling with alcoliol, as follows : 2CS(NPhH)(NH.C.H.NH.) = (PhHN.CS.NH), C, H, + C, H, (NH2), Tho o and p isomerides behave similarly.

p-Amido-s-di-phenyl-thio-urea. From phenylene diamine and PhNCS dissolved in benzene (L. a. W.). Reddish prisms (from alcohol). Sol. glacial acetic acid, insol. ether or benzenc. Begins to decompose at 163°, forming p-phenyleno-thio-urea and aniline.

AMIDO-p-PHENYL-TOLUENE C₁₂H₁₃N i.e. C₁₃H₁₁, NH₂. [93°-97°]. Amido-tolyl-phenyl. Obtained from p-phenyl-toluene, C₅H₃, C₅H₄, CH₈, by nitration and reduction (Carnelley, C. J. 29, 21).

Salt. B'HCl. [c. 283°]

DI AMIDO-PHENYL-TOLYL-KETONE $0_{i_1}H_{i_1}N_{i_2}O$ i.e. $H_iN.C_iH_i.CO.C_iH_i(CH_i).NH_i.$ [about 220°]. Colourloss needles. Formed togother with oxy-amido-phenyl-tolyl-ketone and di-oxy-benzophenone by heating commercial rosaniline with water at 270°.

Di-benzoyl derivative C, H, O(NHBz). [226°], colourless needles (Liebermann, B. 16,

1927).

DI-AMIDO-PHENYL-TOLYL-METHANE v.

DI-AMIDO-BENZYL-TOLUENE.

Di-amido-phenyl-di-tolyl-methane $O_1H_{22}N_2$ i.e. $C_6H_6CH(C,H_6NH_2)_2$ [185°]. Formed by heating a mixture of p-toluidine, p-toluidine in anochloride, and benzou, aldehyde for several hours at 120° (Ullmann, B. 18, 2091). It forms a compound with benzene, crystallising in glistening needles.

Tri-amido-di-phenyl-tolyl-methane G₀H₂₁N₃ i.e. (H,N.C,H),CH.C,H₈NH₂. Leucaniline. [100°]. Obtained by reducing rosaniline (q. v.) (Hofmann, Pr. 12, 9; Rosenstiehl a. Gerber, A. Ch. [6] 2, 341). Small crystals (from boiling water). Sl. sol. hot water, or other, v. sol. Converted into rosanilino by oxidaalcohol. tion. Salts. — B"'H₂Cl₂ aq. — B"'₂3H₂PtCl₂.— B"'3HNO₂.

Tri-acetyl derivative [168*]. Needles. Gives tetra-acetyl-rosaniline when oxidised with K, Cr. O, and acetic acid (Renouf, B. 16, 1808).

Tri-p-amido-phenyl-di-tolyl-methane
C₁₁H₂₂N₃ i.e. (H₂N.O.H₂), CH.O.H₄.NH₄. Prepared
by reduction of nitro-di-amido-phenyl-di-tolylmethane with zino dust and HCl (Fischer, B. 15, Small colourless prisms or long nesdles. On oxidation it gives a rosaniline which dyes a bluer shade than ordinary rosaniline.

o-AMIDO-s-DI-PHENYL-UREAC, 3H13N3O i.e. NH2.C6H4.NH.CO.NHC6H3. From phenyl cyanate and o-phenylene diamine in benzone solution (Lellmann a. Würthner, A. 228, 220). Slender silky needles (from alcohol). V. sol. glacial acetic acid, m. sol. alcohol, v. sl. sol. benzene, insol. ether. In capillary tubes it cakes together and partially melts at 182°, splitting up into anilino and phenylene-urea, [305°

m-Amido-s-di-phenyl-urea. From m-phenylene diamine and PhNCO in benzeno (L. a. W.). Grey needles (from dilute alcohol). V. sol. alcohol, and glacial acetic acid, sl. sol. cther and benzene. In capillary tubes it decomposes at 185° into aniline and m-phenylene-urea,

[above 300].

p-Amido-s-di-phenyl-nrea. From p-phenylenediamine and PhNCO in benzene (L.a. W.). Slondor white needles (from alcohol). Sol. glacial acetic acid, v. sl. sol. benzeno, insol. ether. Decomposes about 210°-220° into aniline and p-phenylenc-urea [above 320°].

Di-amido-di-phenyl-nrea C13H14N,O CO(NH.C.H.NH.)2. From tetra-nitro-di-phenyluren, (C,H3(NO.)2.NH)2CO by reducing with tin and HCl (Fleischer a. Nemes, B. Io, 1296). Lamina (from alcohol); sl. sol. cold water .-B',H.PtCl.

AMIDO-PHENYL-URETHANE v. AMIDO-

PHENYL-CARBAMIC ETHER.

o-AMIDO-PHENYL-VALERIC ACID [I:2] C.H.(NH.).CH..CH..CH..CH..CH..CO.H. [629]. White needles. Formed by boiling an alcoholio solution of eso-di-bromo-amido-phenyl-valerio acid with sodium-amalgam. It could not be converted into an inner-anhydrido even by dehydrating-agents. Acetyl derivative: [1510] (Diehl a. Einhorn, B. 20, 385).

AMIDO-PHOSPHENYLIC ACID v. PHOS-PHINES

AMIDO-PHTHALIC ACID C.H,NO. i.s. $C_eH_3(NH_2)(CO_1H)_2$ [1:2:3].

Salt.-II.A"IISnCl, 2aq: noedles, got from nitro-plithalic acid by tin and HCl. The hydrochlorido gives off CO2 on evaporation, becoming m-amido-benzoic acid (Miller, A. 208, 245).

Ethyl ether Et.A". Oil; got by reducing

ethyl con-nitro-phthalate.

Amide - phthalic acid O.H. (NH2)(CO.H) [1:3:4]. Its hydrochlorido splits up, like that of the preceding acid, into CO, and m-amido-

benzoic aoid (M.).

Ether Et.A" [95°] (M.). Got by reducing u-nitro-phthalio ether (M.; Koenigs, B. 10, 125). Monoclinic prisms (from alcohol). Etheresi solutions show faint blue fluoresoence.

Adetyl derivative [1229]. Minute

Amido - iso - phthalic acid C.H.NO. 2aq. [above 300°]. S. 104 at 15°; 92 at 99°. Formed by reducing nitro-iso-phthalic acid, [249°] (Storrs a. Fittig, 4, 158, 285; Beyer, J. pr. [2] 25, 491). Prisms (from alcohol) or plates (from water). Solntions give a deep reddish-

brown colour with Fe,Cl,.

Salts: K2A"; gives no pps. with salts of Ca or Ba, but pps. with salts of Zn, Cd, Cu, Ag, 1'b, and Hg.—Na,A",—MgA"4; aq: S. 20 at 15°.—CaA" 3; aq: S. 74 at 15°.—SrA" aq: S. 86 at 15°.—BaA"1; aq: S. 543 at 15°.—ZnA".—CdA".—AgHA".—H,A"HClaq.— (H,A"HBr.—H,A"HNO, 1;aq.

Methyl ether Me₂A" [176°]; solidifies

at 164°.

Ethyl ether Et.A" [118°]; solidifies at 113°. Prepared by treating a mixture of nitro-iso-phthalic ether (50 g.), alcohol (300 g.), and cono. HCl (500 g.), with zino dust at 0°. Tufts of thin plates (from alcohol) or slender needles arranged in crosses (from water). V. sl. sol. water. Solutions fluoresce violet-red.

Amido-tere-phthalic acid C₃H₃NO₄ i.e. C₄H₄(NH₂)(CO₂H)₂[2:1:4]. Obtained by reducing nitro-terephthalic acid with tin and HCl (Warren de la Ruo a. Hugo Müller, Pr. 11, 112). Thin lemon-yellow prisms; v. sl. sol. cold water, alcohol, ether, or chloroform. Decomposed by heat without previous fusion. Its solution fluoresces blue.

Di-methyl ether Me2A" [126]. Salts: Me, A"HCl: white needles, saponified by water. --(Me₂A"HCl)₂PtCl₄ (Ahrens, B. 19, 1636)

Di-amido-terephthalic acid C.H.N.O. i.e.

C₆H₂(NH₂)₂(CO₂H)₂ [3:6:1:4). Ether Et₂A" [168°]. Formed by the action of bromine upon di-amido-di-hydro-terephthalic ether (di-imide of succino-succinic ether) dissolved in strong H₂SO₄. Glistening orange needles. Sparingly soluble in alcohol and ether with a yellow fluorescence. By diazotisation and treatment with Cu.Cl, it is converted into di-chloro-terephthalic ether, which is reduced by The sodium-amalgam to terephthalio ether. sulphate forms very sparingly soluble colourless needles (Baoyor, B. 19, 430).

AMIDO - PHTHALIDE C,H,NO, $C_4H_3(NH_2) < CO \atop CH > O [4:\frac{1}{2}]$. [178°]. Formed by roducing nitro-phthalide [141°] (Hoenig, B. 18, 3448). Short prisms; sol. chloroform, sl. sol. alcohol, ether, and benzene, v. sl. sol. cold water. Salts: B'HOl: needles.—B'2H.PtCl.

D1 - AMIDO - ISO - PHTHALOPHENONE C20H16N2O2. Two isomeric compounds of this formula are obtained by reducing the two di - nitro - phthalophenones that are got by nitrating iso-phthalophonone C₆H₄(CO.C₆H₅)₂ [1:3] (Ador, Bl. [2] 33, 56).

AMIDO-PODOCARPIC ACID v. Ponocarpio

DI-AMIDO-PROPANE v. TRIMETRYLENE-DIAMINE and PROPYLENE-DIAMINE.

(3:4:1)-AMIDO-PROPENYL-BENZOIC ACID C10H11NO2i.e. C6H3(NH2)(C3H5)CO2H[3:4:1][94°]. Fornation:—1. By reduction of nitro-propenyl-benzoic acid with FeSO, and NH.—2. By boiling amido-oxypropyl-benzoic acid with HCl (Widman, B. 16, 2572). Long with readles. Facility of the control of the white needles. Easily soluble in alcohol, ether, and benzene, sparingly in water and ligroine.

Tolerably marked basic properties.

Salts: A'H,HCI; long colourless easily soluble prisms. — (A'H,HCl),2PtCl,; easily soluble yellow needles.—A'H, AcOH: colourless

prisms, [c. 160°].

Acetyl derivative C_aH_a(NHÃc)(C_aH_a)CO₂H—[212°], long white needles, sl. sol. hot water. By the action of nitrous acid it is converted into methyl-cinnoline carbox ylic acid $\&O_2H.C_6H_2 < \stackrel{CMe:CH}{N} >$, diazo-propenyl-benzoio acid,

 $CO_2H.C_0H_2$ < $N:N.OH^2$, probably being the in-

termediate product (Widman, B. 17, 722).
Amido-propenyl-benzoic acid

C₆H₃(NH₂)(C₃H₃)CO₂H [2:4:1]. [165°]. Formed by

Heating anido-oxypropyl-benzoic acid with dilute HCl (Widman, B. 19, 272). Yellow plates.

Acetyl derivative: [122]; white prisms.

a-AMIDO-PROPIONAMIDE C.H.N.O i.e.
CH.3CH(NH.).CO.NH₂. [above 250°]. Occurs in urine (Baumstark, A. 173, 342). Small columns, sl. sol. cold water, m. sol. hot water, insol. ether insol. alcohol (difference from urca). Converted by nitrous acid into sarco-lactic acid, and by baryta-water at 150° into CO2, NH4, and ethylamine.

a-AMIDO-PROPIONIC ACID v. ALANINE.

β-Amido-propionic acid C₃H₇NO₂ CH_(NH_).CH_,CO_H. [180°]. Mol. w. 89.

Formation .- 1. Together with \$\beta\$-imido-propionic acid, by the action of NII, upon 8-iodopropionic acid (Heintz, A. 156, 36; Mulder, B. 9, 1903).-2. From cyano-acetic acid by reduction with Zn and H2SO4 (Engel, B. 8, 1597).

Properties.—Prisms; v. e. sol. water, sl. sol. alcohol. Sweet taste. Split up by distillation

into NH, and acrylic acid.

Salt.—CuA', 5aq: dark-bluo prisms.
(a)-AMIDO-PROPIONITRILE C,H,N, CH, CH(NH,).CN. A mixture of aldehydeammonia and prussio acid (30 p.c. solntion) is acidified with H.SO,Aq (1:3) (Erlenmeyer a. Passavant, A. 200, 121). Liquid; quickly changes to imido-propionitrile, giving off NH2. -B'HCl.--B',H,PtCl.

AMIDO-PROPYL-ALCOHOL v. OXY-PROPYL-AMINE.

AMIDO-ISOPROPYL-BENZOIC

AMIDO-CUMINIC ACIN. AMIDO-n-PROPYL-CINNAMIC ACID

C₁₂H₁₅NO₂i.e. C₂H₃(C₂H₃)(NH₂).C₂H₂.CO₂H[4:2:1] [155°]. Formed by reduction of nitro-n-propylcinnamic acid with FcSO, and NH, (Widman, B. 19, 277). Glistening yellow needles. Easily soluble in hot alcohol. By heating with dilute HCl for a long time it is converted into npropyl-carbo-styri. [162°].

a-AMIDO - p - PROPYL - PHENYL - ACETIC ACID C₁₁H₁,NO₂ i.e. C₂H₁,C₄H₁,CH(NH₂).CO₂H₂. [197°]. Prepared by saponifying the product of the action of HCN npon cumin-hydramide (Plochl, B. 14, 1316). Sl. sol. cold water, fisol.

alcohol and ether.

AMIDO-PYRENE v. PYRENE.

AMIDO-PYROCATECHIN *C.H.NO. *C.H.(NH.)(OH)2. E pound by Sn and HCl. By redneing the nitro-com-

Salt: B'HCl: dark needles. Sodie car-

bonate liberates the free base which, however, is rapidly exidised by air forming a violet solution (Benedikt, J. pr. [2] 18, 457; B. 11, 363).

Methylene derivative C,H,NO

derivative C,II,NO2 $H_2N.C_eH_3 < O > CH_2$. Obtained by reducing methylene - nitro - pyrocatechin or nitropiperonylic acid (Hesso, A. 199, 34I). Brownish oil. Salt: B'HCl.

AMIDO-PYROGALLOL C.H.NO. i.e.

C.H.(NH2)(OH), Amido-pyrogallic acid. From the nitro-compound. Its alkalino solution turns blue in air.

Salt.-B'HCl: needles (Barth, M. 1, 884). AMIDO-PYRO-MECONIC ACID C.H. NO. i.e. C3H3(NH2)O3. From nitro-pyro-meconic acid, tin and IICl (Ost, J. pr. [2] 19, 194). Needles (from water). Fe₂Cl₈ gives a bluo colour, changed to red by excess. B'HCl aq.

Di-amido-pyro-mellitic ether

C₄(NH₂)₂(CO₂Et)₄ [134°]. From the nitro compound (Net, A. 237, 24). Diacetyl derivative [149°]

(a)-AMIDO-PYRRYL METHYL KETONE C_sH_sN₂O i.e. C₄H₃(NII₂)N.CO.CH₃. Formed by reduction of (a)-nitro-pyrryl methyl ketone with tin and HCl (Ciamician a. Silber, B. 18, 1460).-B'₂H₂PtCl_a: long yellow needles.
(B.4)-AMIDO-QUINOLINEC,NH_a.NII₂.[67°].

Preparation .- I. By reducing nitro-quinoline, [89°] (Koenigs, B. 12, 451).—2. By heating oxy-quinoline with zine-chloride ammonia (Bedall a. Fischer, B. 14, 2573). Plates. Discolves in acids. CrO, gives a blood-red colour.

(B. 2)-Amido-quinoline C, H, N, [114]. Pre. pared by reduction of nitro-quinolino from pnitraniline (La Coste, B. 16, 670). Colourless plates or flat needles (containing 2aq). Sublimable. V. sol. alcohol and other, less in water and ligroin. Salts: B"HCl: large colourless prisms.—B",II,Cl,PtCl, 2 aq.: crystalline pp.

Picrate B"(C, II2(NO2),OH)2: needles. (B. 3)-Amido-quinoline

CII:CH

O4H3(NH2) [110°]. Prepared by $N = \dot{C}\Pi$

heating (B. 3)-oxy-quinoline with ammoniacal ZnCl₂ (Ricmerschmied, B. 16, 725). Yellow plates. Sublimable. Sol. alcohol, ether, and hot water; sl. sol. cold water. The picrate forms long red needles, v. sl. sol. ether.

(a)-Di-amido-quinoline C,H,N, C.H. (NH2)2N. [156° uncorr.]. Formed by reduction of (a)-di-nitro-quinoline [183°] with SnCl. (Claus a. Kramer, B. 18, 1247). Thick SnCl₂ (Claus a. Kramer, B. 18, 1247). Thick yellowish needles.—B"H.Cl₂PtCl₄: red needles.

(B)-L1 - amido - quinoline. [163° uncorr.]. Forme by reduction of (β) -di-nitro-quinoline [132] with SnCl. (C. a. K.). Small yellow needles or plates. Is not sublimable or volatile with steam. V. sol. water and alcohol, sl. sol. ether, benzene, and ligroin.-B2H2Cl2PtCl4:

yellow crystalline powder.
DI-AMIDO-QUINONEC, H2(NH.), O. [6:2:4:I]. Diacetyl derivative C.H. (NHAc), O.: [265°-270°]. Formed by oxidation of tetra-acotyl-di-amido-hydroquinone C_cH₂(NHAc)₂(OAc)₂ or tri-acetyl-tri-amido-phonol C_cH₂(NHAc)₃OH (from picric acid). By heating with SnCl, dissolved in conc. HCl it yields di-amido-hydroquinone (Nietzki a. Preusser, B. 19, 2247; 20, 797).

DI-AMIDO-QUINONE-IMIDE #. AMIDO-DI-IMIDO-PHENOL

(B. 2)-AMIDO-QUINOXALINE C.H.N. i.s. N:CH CaH3(NH2) [159°]. Formed by conden-

N:CH sation of glyoxal with (I:2:4)-tri-amido-benzone (Hinsberg, B. 19, 1254). Yellow neodles or large crystals. Sublimable. V. sol. water, alcohol, and chloroform, m. sol. ether and bonzene. The ethercal and chloroform solutions have a yellowish green fluorescence.
The aqueous solution gives yellow pps. with
AgNO, and HgCl₂. Its solution in HCl is deep violet.

Salts.-B'HCl: brown plates with green reflection.-B'_H_SO_.-B'_H_Cl_P'tCl_.

AMIDO-RESORGIN C_H_NO_ i.e. [I:2:4] CoHa(NHa)(OH)2. Formed by reducing nitroresorcin with tin and HCl (Weselsky, A. 164, 6). -B'HCl 2aq: gives brown colour with Fo2Cl. The free base is unstable.

Ethers: CoH3(NH2)(OEt)2: [32°]; (251°). From benzene azo-di-cthyl-resorcin (Will a. Pu-

kall, B. 20, I124).—C₀H₂(NII.₂)(OII)(OEt). [I48°]. Amido-resorcin. Behyt ether C₆II.₃(NII.₂)(OEt).₂ [I:2:6]. [124°]. From benzeneo-azo-di-ethyl-resorein (Pukall, B. 20, 1148).

Di-amido-resorcin C.H.N.O. i.e. C.H.(NH.).(OH). [1:3:4:6]. The hydrochloride is obtained by reducing dinitroso-resorcin (Fitz, B. 8, 633) or benzene-disazo-resorcin (Liebermann a. Kostanecki, B. 17, 881). It gives a blue colour with Fe Cl. The free base is unstable. If the hydrochloride is suspended in chloroform, a little aqueous NaOH added, and then a large quantity of water, a beautiful blue

colour is produced.—B"ILSO, I aq.

Di-amido-resercin. Formed by reduction of di-nitro-resorein with tin and HCl (Typko, B. 16, 555). The hydrochloride (B'H,Cl2) forms easily soluble flat needles. Fe.Cl, produces a ppn. of steel-blue prisms of di-imido resorcin. AMIDO-SALICYLIC v. OXY-AMIDO-BENZOIO.

DI - AMIDO - STILBENE v. DI-AMIDO-DI -PHENYL-ETHYLENE.

AMIDO-STRYCHNINE C21H21N3O2 i.c. C₂H₂₁(NH₂)N₂O₂. [275°]. (c. 280°) at 5 mm. From nitro-strychnine and SuCl. (Loebisch a. Schoop, M. 6, 848). Cubes (from alcohol). Insol. water, sl. sol. benzoline, m. sol. alcohol, v. e. sol, ether and chloroform. Its salts are very much more solublo than those of strychnine; they turn reddish-violet in moist air. They give tho general reactions for alkaloids. Give no colour with conc. H2SO, and K2Cr2O,. A dilute acid solution is turned blue by aqueous K2Cr2O, or by Fe Cl. Salts .- B"211Cl: prissus .- B"H, PtCl.

Acctyl derivative C2, II2 (NAcH) N2O2 aq

[205°] (L. a. S. M. 7, 77). Di - amido - strychnine C21 II 20 (NII 2) 2 N2 O2 [263°]. From di-nitro-stryelinine, tin, and HCl (Hanriot, C. R. 96, 586; Bl. [2] 41, 236). Prisms (from chloroform); v. sl. sol. water and ether. m. sol. alcohol, v. sol. chloroform. Gives no colour with conc. H2SO, and K,Cr2O, A dilute acid solution is turned violet-blue hy oxidising agents such as K2Cr2O, Aq or NaOCl

p-AMIDO-STYRENE C. H., N i.e. C.H., (NH.,).CH:CH., [76°-81°]. A body of this composition is formed by reducing p-nitrocinnamic soid with tin and HCl (Bender, B. 14, 2359), and by heating p-amido-cinnamic acid (Bernthsen a. Bonder, B. 15, 1982).—B'₂H₂PtCl₃.

o-AMIDO-STYRYL-ACRYLIC ACID C₁₁H₁₁NO₂

i.e. C.H. (NH2).CH:CH.CH:CH.CO.H. o-Amidoeinnamenyl-acrylic acid. [177°]. Formed by reduction of o-nitro-styryl-acrylic acid with forrous sulphate and ammonia. Yellow needles. V. sol. chloroform, ether, alcohol, and acctic acid, sl. sol. CS2 and hot water, v. sl. sol. cold water. Its ethereal solution has a green fluores-It forms salts with acids and with The hydrochloride is easily soluble, bases. the sulphate sparingly soluble. The salts with bases are deep yellow.

Acetyl derivative

C.H. (NHAe).C.H. CO,H: [253°]. Small white tables, sol. hot alcohol, sl. sol. cold alcohol and ether, insol. water (Diehl a. Einhorn, B. 18, 2332).

o-AMIDO-STYRYL-PROPIONIC ACID C₃H₄(NH₂).CH:CH.CH₂.CH₂.CO₂H. o - Amido cinnamyl-acetic acid. [59° hydrated]. Crystals (+H2O). Easily soluble in ordinary solvents. Formed by reduction of o-amido-styryl-acrylic acid with sodium amalgam (Diehl a. Einhorn, B. 20, 378).

AMIDO-SUCCINAMIC ACID v. ASPARAGINE. AMIDO-SUCCINIC ACID v. ASPARTIO AOID. Di-amido-succinio acid C₄H₈N₂O₄ i.e. CO₂H.CH(NII₂).CH(NH₂).CO₂H. [125°].

Formation .- 1. From di-bromo-succinic acid and NH₃ (Lchrfeld, B. 14, 1817).-2. By reducing the di-phonyllydrazide of di-oxy-tartaric acid, CO₂H.C(N₂HPh).C(N₂HPh).CO₂H, in alkaline solution with sodium amalgam. The yield is 35 p.c. of the theoretical (Tafel, B. 20, 247).

Properties .- Prisms; v. sl. sol. water, alcohol, ether, acctone, acctic acid, chloroform, aniline,

phenol, and CS. Sol. aqueous acids and alkalis. Di-amido-succinic acid C₄H_aN₂O₄. [151° uncorr.]. White needles or prisms. Sol. water, alcohol, and ether. The acid is isomeric with the preceding. The ether is formed by the action of NH, on di-chloro-succinic ether.

Diethyl-ether A"Etz. [122° uncorr.] Colourless needles or trimetrio prisms. Sol. alcohol and other, v. sl. sol. water.

Salts.-A"Ag, and A"Pb: insol. pps.-

A"Cu: green pp.
Di-amide C.H., N. (CO.NH.) 2. [160° uncorr.].

Long slender needles. Insol. water and ether (Claus a. Helpenstein, B. 14, 624; 15, 1850). AMIDO - SUCCINURIC ACID v. URAMIDO-

SUCCINIO AOID. AMIDO - SULPHOBENZIDE v. AMIOO-DI-PHENYL SULPHONE

AMIDO-SULPHO-BENZOIC ACID C,II,NSO, i.e. C₆H₃(NH₂)(SO₃H)(CO₂H) [1:3:5]. nitro-m-sulpho-benzoic acid and aqueous ammonium sulphide (Limpricht a. Uslar, A. 106, 29). Needles, v. sol. hot water, m. sol. alcohol, v. sl. sol. ether. Blackened by heat. Combines with bases but not with acids.

(a)-amido-sulpho-benzoio acid C,H,NSO, aq i.e. $C_{6}H_{3}(NH_{2})(SO_{3}H)(CO_{2}H)$ [1:x:5]. Obtained, togother with the following acid, by sulphonation of m-amido-benzoio acid (Griess, J. pr. [2] 5, 244). Rour-sided lamina, m. sol. hot water.-Salt:-BaA"2aq: v. sl. sol. water.

(B)-amido-snlpho-benzoio acid O.H.NSO. lamines; v. al. sol. hot water. -- Salt: BaA" Saq: m. sol. water.

Amido-sulpho-benzoio acid

C_tH₄(NH₂)(SO₄H)(CO₂H) [1:3:6]. Rhombic plates, sol. hot water. Dilute solutions show blue fluorescence (Hart, Am. 1, 363).

Amido-sulpho-benzoio acid $C_2H_3(NH_2)(SO_3H)(CO_2H)$ [1:2:4].

Imide C_eH₃(NH₂)<SO₂>NH. [285°]. From the amide of p-nitro-toluene sulphonic acid by oxidation and reduction (Noyes, Am. 8, 167). Colourless crystals, v. sl. sol. water. Its solution shows dark blue fluorescence.

AMIDO-SULPHO-BENZOLIC ACID. An old name for amido-benzene sulphonio acid v. Amino-Benzene,

AMIDO-SULPHO-PHENOLIC ACID. An old name for amido-phonol sulphonic soid v. Amido-

a-AMIDO-p-SULPHO - PHENYL - PROPIONIC ACID C. II, NSO, i.e.

SÖ₃H.C₈H₂.CH₂.CH₂.CH(NH₂).CO₂H. From α-amido-phenyl propionic acid (20 g.), conc. H_2SO_4 (30 g.) and Nordhausen acid (25 g.) (Erlenmeyer a. Lipp, A. 219, 209). Groups of short prisms (from water). M. sol. water, v. sl. sol. alcohol, insol. ether. Does not combine with HCl. Fused with KOH gives p-oxy-benzoic acid. -Salts: BaA', 4aq: flat prisius.
AMIDO-TEREPHTHALIC ACID v. Amido-

PHTHALIC ACID.

m-AMIDO-THIO-BENZAMIDE C.H.N.S i.e. C, H4(NH2).CS.NH2. Obtained by boiling m-nitrobenzonitrile with aqueous ammonium sulphide (Hofmann, Pr. 10, 598; B. 1, 197). Needles (from water). Weak base. Decomposed by heat into H.S and amido-benzonitrile. Alcoholio solution of iodine converts it into C14H12N4S, crystallising from water in slender needles Forms [129°]. a platino - chloride

C₁₄H₁₂N₄SH₄PtCl₅ (Wanstrat, B. 6, 332).

p - Amido - thio - benzamide (170°). From
p-nitro-benzonitrile and cone. H₂SO₄ (Engler, A.
149, 299). Crystals; m. sol. alcohol.

AMIDO - THIO - CRESOL v. AMIDO-TOLYL

AMIDO-THIOPHENE C.SH.2(NII2). pared by reducing nitro-thiophene with tin and alcoholie HCl (Stadler, B. 18, 1490, 2316). Yellow oil. Very unstable; being changed in 12 hours into a brittle resin. The hydrochloride reacts with diazo salts forming stable azo compounds.

Salts.—B'HCl.—B'.H.SnCl,. a-AM(DO-THIENYL-ACETIC ACID

C_eH₇SNO₂ i.e. C₄SH₂.CH(NH₂).CO₂H. Formed by reducing the oxim of thionyl-glyoxylio acid C,SH, C(NOH) CO,H with tin and HCl (Bradley, B. 19, 2115). Phites or grains; decomposite 235°-240°. Saits.—The acid gives pps. with salts of Cu, Hg, Bi, and Zn, but no pps. with salts of Fe, Mg, Mn, Ur, Ni, Ba, Ca, Sn, or Pb.-CuA', aq. -HA'HCl.

AMIDO - THIOPHENOL v. AMIDO-PRENTA MEROAPTAN.

C10H15NO AMIDO - THYMOL C_sH₂Pr(NH₂)(CH₃)(OH) [1:3?:4:6]. Nitrosothymol, prepared from sodium-thymol, KNO₂ and H₂SO₄ (Schiff, B. 8, 1500), is reduced by Sn and HCl to the well-crystallised tin salt of i.e. C.H.(NH.)(SO.H)(CO.H) [1:2:5]. Six-aided p-amido-thymol. This is dissolved in water and decemposed by H.S (Andresen, J. pr. 181, 169).—Salt: B'HCl: decomposee at 210°-215°.

Reactions.—1. Bleaching powder solution converts it into thymo quinone-ohloro-imide (q.v.).-2. A solution of bromine in NaOH oxidises it to thymo-quinono .- 3. Bromine water has the same effect.

Amido-thymol sulphonic aoid C10H14(SO3H)NO is among the products of the action of cone. NaHSO3Aq upon thymoquinone-chloro-imide

(A.). Needles or prisms.
Di-amide-thymequinene C10H14N2O2 C.PrMe(NH2)2O2 or Oxy-amido-thymo-quinon-

imide C_PrMe(NH_2)(OH) Formed by

heating phenylamido-oxy-thymoquinone with alcoholio NH, at 100° (Anschütz a. Leather, C.J. 49, 725). Dark blue crystals, insol. water, ether, benzene, ohloroform, and CS2; v. sl. sol. alcohol; sol. glacial HOAo (crystallising with HOAc); v. col. HClAq, forming a red solution.

AMIDO-TOLUENE v. Toluidine and Benzyl-

Di-amido-tolueue v. Tolylene-diamine and AMIDO-BENZYLAMINE.

TRI - AMIDO - TOLUENE

C_sH₂Me(NH₂), [1:3:4:5].

p-Acetyl derivative C.H.,Me(NHAc)(NH.), [1:4:3:5] [o. 264°]; pearly rods (containing aq); sol. acetio acid and hot alcohol, insol. water, ether, and benzene. Formed by reduction of acctyl-di-nitro-p-toluidine (1 pt.) with tin (3 pts.) and conc. HCl (8 pts.). The hydrochloride and conc. HCl (8 pts.). (B'HCl aq) forms white concentric easily eoluble needles (Niementowski, B. 19, 716).

Benzoyl derivative C.H₂Me(NHBz)(NH₂)₂ [1:4:3:5] [c. 185°]. Formed by reducing benzoyl-di-nitro-p-teluidine (Hübner, A. 208, 318). 1nsol. water, sol. alcohol and ether. Salts.—B"211Cl.—B"H.SO.

Tri-amide toluene C₈H₂(CH₃)(NH₂)₃ [1:2:4:7]. Very oxidisable crystalline solid. Tri-acid base. Prepared by reduction of nitro tolylene m-diamine. B"(HCl), and B",(H,SO), are white crystallino solids (Ruhemann, B. 14, 2657).

AMIDO-TOLUÈNE SULPHINIC ACIDS

C,H,NSO,

o Amido-tolusne sulphiuic acid

O.H.Me(NH.)SO.H[1:2:4]. o.Toluidine sulphinic acid. S. 148. From o.amido.tolucne thiosulphonic acid and sodium amalgam (Paysan, A. 221, 361). Rectangular tables, sl. sol. water or alcohol, insol. ether or benzene. At 160° it decomposes without melting.

Reactions.-1. With yellow ammonic sulph. ide forms amido-toluene thiosulphonio acid.-2. KMnO, forms amido - tolueno sulphonio acid _ 3 Boiling HCl forms the isomeric toluene sulphamine. _ 4. Nitrous acid forms a diazo compound which when warmed with alcohol forms the ethyl derivative of cresol sulphonic acid.

Salts.—KA'.—BaA', 2aq.—AgA'.
Toluene sulphamine C,H,NSO, [175°]. Got by heating o amido toluene sulphinic acid with HCl and ppn. by NH₂. Needles in stars (from alcohol).—B'HCl: groups of slender needles.

p-Amido-telusne sulphinic acid C,H,NSO, i.o. C₀H₂(NH₂)Me.SO₂H [1:4:5]. From C.H. (NH2)Mc(SO2SH) by boiling with HCl but since much then changes to toluene sulph-

amine it is better to reduce it with sodium amaigam (Heffter, A. 221, 847). Hard prisms. Does not melt below 240°. Insol. alcohol. al. sol. cold water, v. sol. hot water.

Reactions.-1. Warmed with a solution of sulphur in ammonic sulphide it changes to the thiosulphonio acid, C,H,(NH,)Me.(SO,SH).-2. Bromine converts it into amido-toluene sulphonic acid .- 3. Not reduced by Sn and HCl.

Salts.-KA'.-BaA', raq.

Tolueue sulphamine (isomerio with the above). [132°]. Got by heating p-amido-toluene sulphinic acid with conc. HCl. It is a base. Microscopic prisms (got by adding NH₃ to its solution in HCl). V. sol. alcohol and ether, but ecparates from them in a resinous form; sl. sol. water. Dissolved by treatment with water and sodium amalgam (not NaOH alone) forming eodic amido toluene sulphinate.

Salts .- B'HCl: sl. sol. HCl, v. sol. water or alcohol. - B'2H2SO4 - B'HBr: changes readily into amido toluene sulphonic acid. -B'HNO3: warmed with HNO, forms amido - toluene sulphonic acid.

Di-amido-toluene sulphinic acid C, H, N, SO, C.II.Mc(NH₂).SO.H. Tolylene di-amins hinic acid. S. 047 at 20°. From sulphinic acid. S. 047 at 20°. From C.H.Me(NII.), SO.SH by boiling with HCl (Porl, B. 18, 70). Silky needles (containing aq). V. sl. sol. water, insol. alcohol, ether, and glacial

HOAo.—PbA', 2aq: minute needles.

AMIDO-TOLUENE SULPHONIC ACIDS C,H,NSO, (Limpricht, B. 18, 2172). o-Amide-toluene sulphonio acid

C,H,Me(NH,)SO,Haq[1:2:5]. o.Toluidine sul phonic acid. S. 2.76 at 12' (II. Hasse).

Preparation .- 1. By heating the acid sulphato of o-toluidine at 220° 230°, or in a metal dish till solid (Nevile a Winther, C. J. 37, 626; B. 13, 1941; Gerver, A. 169, 374; Pagel, A. 176, 292). - 2. By reducing the corresponding nitro acid (Foth, A. 230, 306).

Salts. -- KA'aq: tables and prisms. -- NaA'4aq: tables. -BaA'27aq: tables and prisms.

-AgA': prisms.

Reactions .- 1. Bromine water forms first CII., C.H.Br(NII.)SO,H [1:3:2:5] then di-bromotoluidine C, II. (CH,)(NH,)Br,[1:2:3:5][46°] is ppd. 2. Fused with alkalis or heated with water or aqueous HCl to 190° it forms o-toluidine .- 3. Nitrous acid and alcohol give m-toluene sulphonio acid.-4. With o-toluidine at 235° it forms a red dye.

o Amido-tolueue sulphouic acid

C_sH_sMe(NH₂)(SO₃H) [1:2:3]. Obtained by reducing the corresponding nitro acid (Pechmann, A. 173, 215). Minute needles; sl. sol. cold water. Gives o toluidine when fused with KOH.

o. Amide-tolusue sulphenio acid C₂H₃Me(NH₂)(SO₃H)aq[1:2:4]. From the nitro acid (Bek, Z. 1869, 211; Beilstein a. Kuhlberg, A. 155, 21; Weckwarth, A. 172, 193; Hayduck, A. 172, 204; 174, 343; Herzfield, B. 17, 904). Long needles or four-sided prisms. S. 974 at 11° insol. alcohol. The aqueous solution is turned violet by Fe₂Cl_s. Potash-fusion gives o-amidobenzoic acid. Bromine gives di-bromo-toluidine sulphonic soid. Salts .- NaA' 4aq .- KA'aq .-BaA'221aq.-PbA'

A mide. — CaHaMe(NH2)SO2NH2 S. 22 at 23°. From C.H.Me(NO.)SO.NH.

m-Amido-tolnene sulphonic acid

C.H.Me(NH.)SO.H [1:3:2]. m-Toluidine sul-phonic acid [275°]. By sulphonation of m-toluidine (Lorenz, A. 172, 185). Tables or plates; sl. sol. water. Bromine-water produces tri-bromo-toluidine.

Salts. -BaA'29aq. -PbA'232aq. m-Amido-toluene sulphonic acid

C₆H₃Me(NH₂)SO₃Haq. S. (dry) ·14 at 19°. From bromo-tolucue sulphonic acid C₆H₃MeBrSO₃H [1:2:4] by nitration and reduction (Hayduck, A. 174, 350). Minute needles.

p-Amido-toluene e.ro-sulphonic acid

C.H.(NH.).CH.,SO.H [1:4]. p-Amido-benzyl-sul-phonic acid. S. 097at 10°. Formed by reducing the nitro acid by NH, and H.S (Mohr, A. 221, 219). Prisms, insol. alcohol, sl. sol. cold water.

Salts.-KA'2 aq. - BaA' 8aq.

The diazo derivative, C.H. < CH. SO. is converted into C.H. (OEt).CH2.SO3H by heating with alcohol under 1100 mm. pressure.

p-Amido-toluone sulphonic acid

C₃H₃Me(NH₂)SO₃llaq [1:4:2]. p-Toluidine sulphonic acid. S. 45 at 20°. A product of sulphonation of p-toluidine (Sell, A. 126, 155; Malycheff, Z. 1869, 212); formed also by resistant the telescopic product of the control of ducing p nitro-toluene sulphonic acid (Beilstein a. Knliberg, A. 172, 230). Rhombohedra (containing aq). Reduces warm ammoniacal AgNO₃. Its aqueous solution is turned red by $\mathbf{Fe_2Cl_s}$ (Herzfeld, B. 17, 904).

Salts.-KA'.-BaA', aq.-PbA', Amide. - C.H.Me(NH.)SO,NH, From C, H, Me(NO,).SO, NH, [186°] by reducing with NII, and H.S (Heffter, A. 221, 209). Salt: C_aH₃Me(NH₃Cl)SO₂NII₂; converted by conc. IICl and nitrous acid into C_aH₃MeCl.SO₂NH₂ [138°]. p-Amido-toluene sulphonic acid

C.H.Me(NII.)SO.H [1:4:3]. S. 10.

Preparation .- 1. By sulphonating p-toluidine at 180°; the preceding acid is also formed, especially if the operation is protracted (Pechmann, 4. 173, 195). -2. By heating p-toluidino acid sulphate at 220°-240° (Nevilo a. Winther, C. J. 37, 632).

Properties. -Yellowish crystals. Lcss soluble

in cold water than the o-compound.

Reactions. - 1. Bromine forms much di-bromo-toluidine, C_uH₂(CH₃)(NH₂)BrBr [1:4:3:5], [73°] and also a bromo-toluidine sulphonic acid. 2. Water at 180° forms p-toluidine and H.SO .. -

8. Potash-fusion gives p-oxy-henzoic acid. -

4. Nitrous ether givos m-toluene sulphonic acid.
Salts.—Bah', 3aq.—Pbh', 2aq.—Agh'.—The
K salt is insol. in cold KOHAq (difference from preceding acid; Schneider, Am. 8, 274).

Amido-tolnene-o-sulphonio acid C.H.Me(NH.)(SO.H) [1:x:2]. S. 34 at 22°. From the (1, 4, 2) acid by nitration, removal of NH, and reduction (Pagel, A. 176, 305).— BaA', 2, aq.—PbA', aq. Amido tolnene sulphonic acid. Obtained by

reducing the product of successive sulphonation and nitration of toluene (Hayduck, A. 177, 57).— Minute oryetale (containing aq) .- BaA'z.

o-Amido-tolnene di-sulphonic acid $C_sH_sNS_sO_s$ i.e. $C_sH_sMe(NH_s)(SO_sH)_s$ [1:2:3:5].

[128], NH, and H, S (Paysan, A. 221, 210). Four-sided columns.

Salt.—C, H, Me(NH, Cl). SO, NH. [240°].

C, H, Me(NH, NH, SO, H [1:2:5] and finning H, SO, by heating an hour at 160° (Nevile a. Winther, C. J. 41. 421). Needles, grouped in stars; sol. water and alcohol.

Salts (H. Hasse, A. 230, 287). -BaA"3aq. -

Reactions.—1. By conversion into the diazo compound and subsequently boiling with HNO, it is converted into di-nitro-o-cresol C.H.Mc(OII)(NO₂)₂ [1:2:3:5].—2. At about 240° it splits np into SO, and C.H.Mc(NH.)(SO₂H) [1:2:5].—3. By Cl.SO₂H at 230° it is changed into an isomeric acid with a salt K.A" Gaq.

o-Amido-toluene disulphonic aoid C.H.Me(NII.)(SO,H) [1:2:4] and CISO,H at 170° (Saworowicz, B. 18, 2181). Minute prisms. At 300° it decomposee into SO, and C.H.Mc(NH.)SO,H[1:2:4]. Salte.-BaA"2aq.-CaA" 2aq.

m-Amido-tolnene disulphonic acid

 $C_uH_uMe(NH_u)(SO_uH)_u$ [1:3:2:x]. By sulphonation of m-toluidine (Lorenz, A. 172, 188). Easily splits up into SO₃ and the mono-sulphonic acid.

Salts.—Ball, A". (?12½) aq.—PbA" 2aq. p-Amido-toluene-disulphonic acid

C₆II₂Me(NII₂)(SO₃II)₂ [1:4:2:3]. From p-toluidine and fuming H₂SO₄ at 200° (Pechmann, A. 173, 217). Nodules; v. o. sol. water and alcohol. Salt. - BaA" 3aq: laminæ.

n-Amido-toluene-disulphonio acid

 $C_0H_2Me(NH_2)(SO_3H)_22_2aq[1:4:2:x]$. Formed from C.H.Mc(NH.)(SO.H) [1:4:2] by CISO.H at 150° or funning II.SO. at 180° (L. Richter, A. 230, 331). Long silky needles, v. sol. water, sol. alcohol. At 290 it splits up into SO. and

C₂H₂Me(NH₂)(SO₃H) [1:4:2]. Salts. — BaA"aq. — - BaH, A", 11 aq. -

BaH2A"2 2aq. - K2A"2aq. - PbA" 12aq.

p. Amido-toluone-disniphonio acid C.H.Me(NII.)(SO.11).2aq [1:4:3:x]. Formed from C.H.Mc(NH.)(SO₃H) [1:4:3] and H.SO₄ or ClSO₃H (L. Richter, A. 230, 314). Mass of minute needles (from water). With water at 140 (or dry at 200°) it splits up into SO, and C.H.Mc(NH.)(SO.H) [1:4:3]. This acid is perhaps identical with that of Pechmann.

Salts.-BaA"3aq.-BaH,A", 3aq.-K,A"2aq.

-PbA".--PbA"2aq.

Diazo derivative *C.H.Me(N.SO.)"SO.H. V. sol. water, insol. alcohol. KA'. - BaA', - PhA', Hydrazine derivative.—From the diazo

acid by SnCl,.
(C, H, Me(N, H,)(SO, H)SO,), Ba 2 aq. Reduces HgO, ammoniacal AgNO, Fe Cl, and Folling solution.

Amido-toluene di-snlphonic acid C_oH₂(NH₂)Me(SO₂H)₂(?2)aq. From p-bromo-toluene disulphonic acid by nitration, and reduction of the resulting nitro-toluene disulphonic acid-(Kornatzki, A. 221, 198).

Di-amido-tolnene exo-snlphonic acid C.H., N.SO, i.e. C.H., (NH.), CH., SO, H. Di-amida-benzyl-sulphonic-acid. Formed by reducing Formed by reducing C.H.,(NO.),CH.SO,H with NH, and H.S (Mohr, A. 221, 228). Silky needlee.

Di-amido-telnene sulphonic acid C,H,Me(NH,),SO,H [1:2:4:5]. Formed from C,H,Me(NO,)(NH,)SO,H and SnCl, (Foth, A. 280, 809). Small brownish prisms, rhombohedra (from water). Salts.—HA'HCl aq: prisms, decomposed by boiling water. — HA'HBr aq.— BaA', 5 aq.—KA' aq. o-AMIDO - TOLUENE - THIO - SULPHONIC

ACID C,H,NS,O, i.e. C,H,Me(NH,)SO,SH [1:2:4]. From C,H,(NO,)MeSO,Cl and ammonio sulphide (Limpricht a. Paysan, A. 221, 360). Four-sided prisms. Decomposes without melting at 115°. Sl. sol. oold water, insol. stoohol. Warmed with HCl forms S and tolucne sulphamine. Salt .-AgA'.

p-Amido-tolnene thio-sulphonic acid C_cH_s(NH_s)Me.SO_sSH [1:4:5]. Forme Formed from C.H.(NO.)MeSO Cl [44°] and ammonic sulphide (Limpricht a. Heffter, A. 221, 315). Hard yellowish prisms (from water). Decomposed at 120° without melting. Insol. alcohol or ether, sl. sol. water. Decomposed by HCl with deposition of S and formation of C, H, (NH2)MeSO2H.

Salts: BaA', 2aq. -AgA'. Di-amido-toluene thiosulphonio acid

C₁H₁₀N₂S₂O₂ i.c. C₆H₂(CH₃)(NI1₂)₂SO₂SH. [152°]. Formation.—1. By reduction of di-nitrotoluene-sulphonic chloride with NH,11S.-2. By reduction of di-nitro-toluene sulphinic acid with NH,11S. Small silky prisms. V. sl. sol. water, iusol. alcohol and ether.

Salts.—A'Ag: white insol. pp. -A'Na: largo tables - A'2Pb: easily soluble (Perl, B. 18, 67).

AMIDO-TOLUIC ACIDS C.H., NO. Amido-

toluylic acids.

(a)-amido-o-toluic acid CallaMe(NHa)COaH [1:4:2]. [196°]. Formed by reducing (a)-nitro-o-toluic acid (Jacobsen, B. 17, 164). Small prisms, v. sol. hot alcohol, and hot water, sl. sol. cold water. Converted by nitrous acid into oxy-toluic acid [172°].

(\$)-amido-o-toluic acid C.H.Me(NH2)CO2H [1:6:2] [191°]. Formed by reducing (β)-nitroo-toluic-acid (Jacobsen, B. 16, 1959; 17, 164). Small needles, v. sol. cold water. Converted by nitrous acid into oxy-toluic acid [183°].

(γ)-amido-o-toluio aoid C_eH₂Me(NH₂)CO₂H [1:5:2]. [153°] (Hoonig, B. 18, 3449); [c.

165°] (J.).

Formation.—1. By reducing (γ) -nitro-o-toluic acid.—2. By heating nitro-phthalide [141°] with HI and P at 205°. Colourless needles; may be sublimed, but at 200° it splits off CO. forming m-toluidine. V. sol. hot alcohol, m. sol. hot water and ether, sl. sol. chloroform, benzene, and cold water. Nitrous acid produces oxy-toluic acid [179°].

Salts: HA'HCl: slender needles .- * CuA'2.

-HA'H₂PO₄: plates.
Amido-m-tolnio acid C₆H₂Me(NH₂)CO₂H

[I:4:3] or [1:4:5].

ation .- 1. The hydrophloride is obtained by warming methyl-isatoic acid with HClAq (Panaotović, J. pr. [2] 33, 61).-2. The same acid is got from m-toluio acid by nitration and reduction (Jacobsen, B. 14, 2354; compare Panastović, loc. cit.).

Properties.—Trimetric, thread-like rods (from water). Sl. sol. water, v. sol. alcohol and ethor. Salt: HA'HCl. [207°]. Colourless trimetrio prisms; m. sol. water and alcohol, sl. sol. ether.

Methyl ether C₆H₃Me(NH₂)CO₂Me. [62°] From methyl-isatoio acid and MeOH at 1800 Flender columns; sl. sol. water.

Amids C.H.Me(NH.) CO.NH. [178°]. From mathyl-isatoic acid and NH.Aq. Small columns (from watsr); v. sol. aloohol.

Anilide C.H.Me(NH.)CONPhH. [240°].

Pearly tablets (from alcohol); v. sl. sol water.

Phenyl-hydrazide C.H., Me(NH.). CO.N., H., Ph. [1980]. From methyl-isatoio acid and phenylhydrazine. Pearly crystals (from aloohol); v. sl. sol. water. Forms a violet solution with cono. H.SO.

(β)-Amido-toluic acid C₆H₃(CH₃)(NH₂)(CO₂H) [1:2:3]. [132°]. Obtained by nitration and reduction of m-toluio acid (Jacobsen, B. 14, 2354; compare Panaotović, J. pr. [2] 33, 61).

Small flat prisms; m. sol. water.

Amido-m-toluio acid. Benzoyl derivative C,H,(NHBz)MeCO,H [5 or 6:1:3]. Formed by oxidation of benzoyl iso-cymidino (Kelbo a. Warth, A. 221, 168). Small yellowish nesdlss (from alcohol).

DI-AMIDO-DITOLYL C,HisNg.

Di-amido-ditolyl

[4:3:1] $(NH_2)MeC_6H_3.C_6H_2Me(NH_2)$ [1:3:4]. o-

Tolidine. [112°].

Formation.—1. By passing Cl₂O into an ethereal solution of o-hydrazotoluene (Petrieff, B. 6, 557).—2. By heating o-hydrazo-toluene (Petrieff, B. 6, 557).—3. By heating o-hydrazotoluene with HCl (Schultz, B. 17, 467).-4. By warming an alcoholic solution of o-azo-toluene with SnCl, and HCl (S.).

Pearly plates; v. sol. alcohol and ether, sl. sol. water. Converted by diazo reaction into mditolyl. Converted by boiling its diazo-perbromide with alcohol, into a di-bromo-ditolyl, which oxidises to bromo-m-toluic acid [1:2:4], [205°]. Salts.-The sulphate and hydrochlorido are sparingly soluble in water.

Acetyl derivative C12HaMex(NAcH)x

[315° cor.]

Di-amido-ditolyl. m-Tolidine.

The sulphate, B"H2SO4, separates slowly when a few drops of H,SO, are added to an alcoholic solution of m-hydrazotoluene (Goldschmidt, B. 11, 1626). The free base has a low

melting point. Gives a bluo colour with Fe₂Cl₈.
'Di-amido-ditolyl.' [107°]. Formed by the action of SO2 or of SnCl2 and HCl upon an alcoholic solution of p-azo-toluene (Melms, B. 3, 554; Schultz, B. 17, 472). Silvery plates, Gives a blue colour with Fe₂Cl_v. Fischer (B. 25, 1019) has shown this body to be tolylenetolyl-diamine.

Di-amido-u-ditolyl

[4:3:1]. (NH.)MeC.H.C.H.Me(NH.) [1:2:4]?

o.m.Tolidinc. Formed by the action of SnCl, and HCl on an alcoholic solution of o-m-azo-toluene. By diazotisation in alcoholic solution it is converted into a ditolyl of boiling point 270° which on oxidation gives isophthalic acid.

Salts.-B"H2Cl2: easily soluble silky needles. -B'H₂SO₄*: very sparingly soluble plates (Sohultz, B. 17, 471).

AMIDO-TOLYL-BENZAMIDINE C₁₄H₁₅N₁ i.a. NH₂,C₂H₃Me.NH.C(NH).C₆H₅ [212°]. From benzonitrile and (1:2,4)-tolylene-diamine hydro-chlorids (Bernthsen a Trompetter, B. d.1, 1758),

White needles.—B'HCl: prismatic tables.

AMIDO-TOLYI-ISO-BUTANE CHI, N i.e. C,H,Me(C,H,)NH, [1:5:2]. (248°). From o-

toluidine and isobutyl alcohol (Effront, B. 17, 2320). S B'₂H₂C₂O₄. Salts.—B'HCl.—B'HBr.—B',H,SO,

Acetyl derivative [162°]. Plates.

Bensoyl derivative [168°]. Nesdles.

Amido-tolyl-isobntane C_cH₃Me(C,H_c)NH₂ [1:3:2]. (244°). From o-toluidine, isobutyl alcohol, and ZnCl2 (Erhardt, B. 17, 419; Effront, B. 17, 2340). Formyl derivative [105°].

Acetyl derivative [142°].

o-AMIDO-TOLYL-ETHANE $C_9H_{13}N$ O.H. Ms(NH2)Et (230°). Mcthyl-ethyl-phcnylamine. Amido-ethyl-toluenc. From o-toluidine, alcohol, and ZnCl₂ at 270° (Benz, B. 15, 1650). Salts.-B',II,SO, -B',H,C,O.

Acetyl derivative [106°]. (314°).

DI-AMIDO-DI-TOLYL-ETHYLENE DIA-MINE v. DI-TOLYLENE-ETHYLENE-TETRA-AMINE.

AMIDO - TOLYL MERCAPTANS C,11, NS. Amido-thio-cresols. Amido-tolyl sulphydrate. Prepared by reducing the chlorides of the corresponding nitro-toluene sulphonio acids (Hess, B. 14, 488).

Amido-o-tolyl mercaptan CaHaMe(NHa).SH $[42^{\circ}].$ Sol. alcohol, other, alkalis, [1:4:2]. and acids; oxidised by air.

Salt.-B'HCl: prisms or tables.

Acetyl derivative [195°]: slender needles; insol. HČl.

Amido-m-tolyl msrcaptan CaHaMe(NH2)SH [1:4:3]. Oil; oxidised in air gives with HgCl, a white crystalline pp.; with Pb(OAc)2 a yellow amorphous pp. Gives anhydro compounds with formic acid, acetic anhydride, and benzoyl chloride.

Amido-p-tolyl mercaptan C_eH₂Me(NH₂)SH [1:2:4]. Oil. Oxidised by air to the disulphide. Salt .- B'HCl : short needles.

Acctyl derivative [240°].

Amido-tolyl mercaptan Cell, Me(NH2)SII [1:2:?] From o-nitro-toluene sulphochloride [36°]. Oil.

Salt .- B'HClaq: six sided tables. Gives with Fe2Cla a pp. of the disulphide. gives glistening plates, and alkaline lead acctate gives a yellow pp.

AMIDO-TOLYL METHYL KETONE C,H,,NO [1:2:5] $C_0H_3(CH_3)(NH_2).CO.CH_3$ [102°]. (280°-284°). Prepared by heating a mixture of o-toluidine (1 pt.), ZnCl, (2 pts.), and acetio anhydride (3 or 4 pts.), for 8 or 9 hours (Klingel, B. 18, 2696). Flat white needles. V. sol. alcohol, other, and hot water, v. sl. sol. benzenc and petroleum-ether. Salts .- B'HCl: flat white soluble prisms. -B'2H2Cl2PtCl4: yellow nesdles, v. sol. alcohol, sl. sol. hot water, insoluble ether. -B'2H2SO4: white needles.

Acetyl derivative C.H.Mc(CO.Me)(NHAc), [144°]; white crystals; v. soi. alcohol and warm water.

AMIDO-TOLYL-(aa)-DIMETHYL-PYRROL C.NH.Me. C.H. NH2. [73°]. (322°). Obtained by heating its dicarboxylic acid (v. infra).

m. AMIDO . TOLYL . (aa) . DI.METHYL. PYR.

ROL (ββ)-DICARBOXYLIC ACID C13H18N2O4 i.e. C.NMe₂(C,H₅NH₂)(CO₂H)₂. From m-tolylene diamine and diacetyl-succinic ether (Knorr, A. 236, 313). Yellow plates (containing 2aq). At 203° it gives CO₂ and m-amido-tolyl-(αα)-dimethyl-pyrrol. Ether Et₂A" [134°].

AMIDO-TOLYL-OCTANE C18H28N f.c. C₆H₃Me(NH₂)C₃H₃... [325°]. From *n*-octyl alcohol, o-tolnidine, and ZnCl₂ at 280° (Beran, B. 18, 145). Salts.—B'HCl.—B'₂H₃SO₄.—B'₂H₄C₂O₄. Acetyl derivative [81°].

DI-AMIDO-DI-TOLYL-OXAMIDE v. Oxalyl-DI-TOLYLENE-TETRA-AMINE.

AMIDO-TOLYL SULPHYDRATE v. AMIDO-TOLYL MERCAPTAN.

DI-AMIDO-DI-TOLYL SULPHIDE

 $C_{14}H_{16}N_2S$ i.e. $(C_6H_3MeNH_2)_2S$ [1:4:2]. Thiotoluidine. [103 3]. Prepared by heating p-toluidine, sulpliur, and litharge together at 150° (Merz a. Weith, B. 4, 393). Laminæ (from aloohol); sl. sol. water.

Salts.—Decomposed by water.—B" 2HCl sl. sol. conc. HClAq.—B"H,l'tCla.—B"H,SO, —B"H,SO, 2aq. — B"H,Br., — B"H,Iz. — B"(CaH,(NO.),3OH), [179°]; silky yellow needles (from benzene); v. sl. sol. ether and cold water.

Diacetyl derivative [211°] Dibensoyl derivative [1860] (Truhlar, B.

20, 664).

Di-amido-di-p-tolyl snlpbide v-carboxylio sr S(C₆H₂Me,NH.CO₂Et)₂. Thio-p-tolyl ethsr Thio-p-tolyl urethane [113°]. From the preceding and Crystals; v. sol. alcohol, ether, and ClCO₂Et. benzene (T.).

AMIDO-TOLYL-UREA C,H,N,O Formed, in small NH2.CO.NH.C6H3Me.NH2. quantity, by the action of tolylens diamine sulphate on potassium oyanate (Strauss, A. 118, 159). V. sol. alcohol.

Di-amido-di-p-tolyl-ursa CO(NH.C, He.NH2) a. Formsd by reducing the corresponding nitrocompound (A. G. Perkin, C. J. 37, 700). Minute satiny needles; sl. sol. alcohol .- B" 2HCl.

AMIDO-TYROSINE C,H12N2O3 i.e. C₀H₃(OH)(NH₂).C₂H₃(NH₂).CO₂H. From nitrotyrosine (Beyer, Bl. 1867, ii. 369). Crystalline powder; v. sol. water, sl. sol. alcohol.

Salts .-- B"H_Cl2aq .-- B"H2SO4 .-- B"2H2SO4.

(B"H₂SO₄)₂ZnSO₄.

AMIDO-URAMIDO-BENZOIC ACIDS

C₈H₆N₃O₃ i.e. NH₂.CO.NH.C₆H₃(NH₂).CO₂H. Prepared by reducing the two nitro-uramidobenzoic acids (Griess, B. 5, 195).

(a)-Acid. Plates; sl. sol. water, v. sl. sol.

alcohol. Salts .- HA'HCl .- AgA'.

(β)-Acid. Plates; ni. sol. hot water. Forms no hydrochloride. Boiling aqueous baryta or HCl forms NH, and amido-carboxamido. benzoic acid, of which the barium salt, Ba(C_aH₃N₂O₃), 4aq crystallises in needles.
(α) - AMIDO - UVITIC ACID C_aH_pNO₄ i.e.

C₆H₂(NH₂)(CH₃)(CO₂H)₂ [2:1:3:5] (?). Colourless solid. Sl. sol. water. Prepared by reduction of (a)-nitro-uvitic acid (Böttinger, B. 13, 1933).

(β)-Amido-uvitio acid [o. 255°] (1966).

B. 9, 807).

AMIDO-VALERIC ACIDS C.H., NO.

a-Amido-n-valerio acid

CH, CH, CH, CH(NH2).CO2H.

Formation .- 1. From n-butyric aldehydeammonia and aqueous HCN (Lipp, A. 211, 359).
2. From its benzoyl derivative which occurs among the products of exidation of benzoylconiine (Baum, B. 19, 506).-3. From bromovalsric acid and NH, Aq at 130° (Juslin, Bl. [2] 87, 3).

Properties. White glistening plates; v. sol. water, sl. sol. alcohol, insol. ether; may be sublimed. Has a sweet taste. Is optically inactive. Nentral to litmus.

Salts. - HA'HCl: needles or groups of prisms; only deliquescent in very moist air. -HA'HNO, - (HA'HCl), PtCl, - CuA'2: small blue plates, sl. sol. water, insol. alcohol. AgA': orystalline pp., sl. sol. water.

y-Amido-valeric acid OH, CH(NH2).CH, CO,H. [193° uncorr.]. Obtained by reduction of the phenyl-hydrazide of \$\beta-aceto-propionic acid-

CH, C(N2HPh). CH2. CH2. CO2H in alcoholic solution with sodium amalgam and acctio acid (Tafel, B.19,2415; 20,249). White plates. V. sol. water, nearly insol. alcohol, insol. benzeno and ether. On heating it splits off H2O and the anhydride distils. The hydrochloride forms glistenin plates, easily soluble in alcohol.

Anhydride Oxy-CH.CH.CH.CH..CO.

methyl-pyrrol di-hydride. (248° i.V.) at 743 mm. colonriess liquid which solidifies in a freezing mixture. V. sol. water, alcohol, ether, and benzene. Its nitrosamine is a yellow oil.

a-Amide-ise-valeric acid (CH₃)₂CH.CII(NII₂).CO₂II.

Formation.-1. From NH, and a brome isovalerio acid (Cahours, A. Suppl. 2, 83; Fittig a. Clark, A. 139, 199; Schmidt a. Sachtleben, A. 193, 105) or a-chloro-iso-valeric acid (Schlebnsoh, 4. 141, 326).—2. From its nitrile (Lipp, A. 205, 18; B. 13, 905).

Properties. - Colourless laminæ, composed of minute monoclinic prisms; v. sol. water, v. sl. sol. cold alcohol or other. Neutral. May be sublimed.

Salts. - HA'HCl: large tables, not deliquescent in moist air. - HA'HNO'3. - CuA'2: scales, sl. sol. hot water. - AgA': spherical groups of crystals, v. sl. sol. water.

Amide Me_CH.CH(NH2).CO.NH2. The hy. droohloride, got by action of fuming HCl on the nitrile, forms monoclinic plates, v. sol. water.

--(B'HCl)₂PtCl₄: prisms.

Nitrile Me,CH.CH(NH,).CN. From isobutyrio aldehyde - ammonia and HCN (L.). Yellow oil; m. sol. water, v. sol. alcohol and ether; gradually changes to imido-di-valero. nitrile, NH(CHPr.CN) ., giving off NII, Salts .-B'HCl: insol. ether. - (B'HCl) PtCl.

B-Amide-iso-valeric acid (CH₂)₂C(NII₂).CH₂.CO₂Haq [c. 215°].

Formation .- 1. Among the products of oxidasion of the sulphate of diacetonamine (Heintz, 198, 51).—2. By reduction of β-nitro-isovalerio soid (Bredt, B. 15, 2321).

roperties. - Crystalline powder; bogins to anblime at 180°. V. e. sol. water, sl. sol. alcohol,

insol. ether.

Salts.-HA'HClaq: needles, [o. 1200].-(HA'HCl) PtCl .- CuA', 2aq : large crystals .-

AgA'. * (AgA')₂AgNO, aq.

Amido-valeric acid C.H.(NH2)O2. Found in the radicles of the sprouting lupin seeds. It occurs along with amido-phenyl-propionio acid, from which it may be separated by virtue of the reater solubility of its copper salt (Schulze a. Barbieri, J. pr. [2] 27, 352).

Properties. — Gilttering plates, resembling leueine (from alcohol). When heated, a woolly substance sublimes out of it. Gives no pp. with ouprio hydrate or acetate (difference from leucine). Salt.—HA'HCl: doliquescent prisms.

Constitution. — Probably identical Lipp's a amido-n-valeric aoid (A. 211, 354).

Amide-valeric acid. Occurs in the pancreas of oxen (Gorup-Besanez, A. 98, 15). Prisms; v. sl. sol. alcohol (difference from leuoine). The hydrochloride forms slender deliquescent needles. An amido-valerio acid was found by Schützenberger (A. Ch. [5] 16, 283) among the products of the decomposition of albumen by baryta-water.

AMIDO-VERATRIC ACID v. di-methyl-di-

OXY-AMIDO-BENZOIC ACID.

AMIDOXIMS. Oxy-amidines. Oxy-imidgamides. The oxims of amides, the general formula being R.C(NOH).NH2.

Formation. - 1. By action of hydroxylamine upon nitriles: R.CN + $N(OH)H_2 = R.C(NOH).NH_2$. 2. By heating the thio amides with alcoholio solution of hydroxylamine (Tiemann, B. 19, 1668): $R.CS.NH_2 + H_2.NOH = R.C(NOH).NH_2 + H_2S.$

Properties. - The amidoxims combine with acids; they also contain hydrogen displaceable by metals. The acid salts of the alkaline metals, RC(NOK).NH,,RC(NOH).NH, crystallise well. The stability is increased by the presence of electro-negative substituents; thus, nitrobenz-amidoxim can be reduced to amide-benzamidoxim without destruction of the amidoxim group

 \hat{E} thers.—R.C(NOEt).NH $_2$. These are formed by the action of iodide of cthyl (or of otheralkyls) upon the alkalino salts, such as R.C(NOK).NII2.

They are bases.

Reactions .- 1. Split up by treatment with acids or alkalis into NH, hydroxylamine, and the corresponding acid: R.C(NOH). $NH_2 + 2H_2O =$ R.CO.OH + H.NOH + NH,. This reaction takes place most readily with methenyl amidoxim (isuretine) and ethenyl-amidoxim, while benzenylamidoxim requires long boiling with HCl before it is decomposed. In this saponification of amidoxims the amide seems to be first formed: $R.C(NOH).NH_2 + H_2O = R.CO.NH_2 + H_2NOH.$ 2. The hydrochlorides are converted by sodium nitrite into amides: R.C(NOII).NH2HČl + NaNO. R.CO.NH, + NaCl + N.O + H.O. - 3. They combine with phenyl cyanate forming bodies called uramidoxims; e.g.: Ph.C(NOH).NH₂ + PhNCO = Ph.C(NOH).NH.CO.NIIPh.—4. Chlorides of acid radicles, R'.CO.Cl, form alkoyl derivatives, R.C(N.O.CO.R').NH₂, which can split off water forming azoxims R.C $\langle N \rangle$ CR. azoxims, although of high boiling point, are extremely volatile in the vapour of other liquids, even ether.—5. Dibasic organic anhydrides give rise to carboxylio acids of azoxims, e.g.: R'.C(NOH).NH₂ + R"<CO>O=

R'.C N.O C.R".CO.H + H.O.-6. Chloroformic ether produces bodies of the composition R.C(NH₂):NO.CO₂Et.—7. Carbonyl chloride gives carbonyl-di-oxims, (R.C(NH₂):N.O)₂CO.— 8. Chloral forms orystalline addition-products. References .- Tiemann, B. 18, 1060, 2456;

19, 1475. The Amidoxims are described as FORMAMIDOXIM, ETHENYL-AMIDOXIM, HEXOAMID-OXIM, BENZAMIDOXIM, CINNAMIDOXIM, TOLUAMID-OXIM, BENZAMIDOXIM CARBOXYLIO AOID, NITRO-BENZamidoxim, &c.

AMIDO-XYLENE v. XYLIDINE.

Exo-Amido-xylens CH, C,H, CH, NH, [1:3?]. From CH, C,H, CII, Cl and alcoholio NH, (Pieper, A. 151, 120). B'HCl [185°].—B',H,PtCl, Oil. Salts .-

Exo. Amido. p. xylene OH, C, H, CH, NH, [1:4]. From CH, C, H, CS. NH, tin, and HCl (Paterno a. Spica, B. 8, 441).

di-Amido-xylene v. Xylylene diamine.

tri-Amido-xylens CaH, N. i.e. CaHMe2(NH2). [1:3:4:6:2]. Formed by reducing tri-nitro-m-xylene [177°] (Greving, B. 17, 2427). White noodles which may be sublimed.

AMIDO-XYLENE-SULPHONIC ACID

CaH, NSO.

Am do xylene snlphonio acid C.H.(NH.)Me.SO.H [4:1:3:6]. S. 276 at 0°; 735 at 100°. From (1,3,4)-xylidino and H.SO. or from nitro-m-xylene sulphonio acid (Jacobson a. Ledderboge, B. 16, 193). Salts.—NaA'aq and KA'aq form large trimetrio tables. -BaA', aq: miuute needles, v. sol. water.—BaA', 2aq (Sartig, A. 230, 334; Nölting a. Kolın, B. 19, 137). Diazo compound C₆U₂Me₂N₂SO₃: plates.

Amido-p-xylene snlphonic acid

 $\mathbf{C}_{\mathbf{s}}\mathbf{H}_{\mathbf{s}}\mathbf{M}\mathbf{e}_{2}(\mathbf{N}\mathbf{H}_{2})(\mathbf{SO}_{\mathbf{s}}\mathbf{H})$ [1:4:6:2]. From p-xylone snlphonic soid by nitration and reduction (Nölting a. Kohn, B. 19, 143). Needles (with aq): sl. sol. cold water. Its salts are easily soluble. Does not give xyloquinone on exidation.

Amido-p-xyleno sulphonic acid

C.H.Me.(NH.)SO.H [1:4:2:5]. From amido-p-xylene and fuming H.SO. or by heating its acid sulphate at 230°. Readily oxidised by CrO. to xyloquiuone. Salts.-NaA': plates, v. sol. water. Bah', 7aq (Nölting, B. 18, 2664; 19, 141).

Di-amido-xylens sulphonic acid C₂HMo₂(NH₂)₂SO₃H [1:3:6:?:4]. From nitroxylidine sulphonio acid and ammonium sulphide (Limpricht, B. 18, 2190; Sartig, A. 230, 313). Fawn-coloured prisms, sl. sol. water, insol. alcohol. For Clacolours the solution winc-red. Salts: BaA', 3 laq.—KA'aq.—PbA', .-HA'IlClaq. AMIDO.m-XYLENOL C,H,1NO i.e.

C₂H₂Me₂(NH₂)(OH) [1:3:x:y]. [161°]. Got by reducing nitro-xylenol (Pfaff, B. 16, 1137). White [161°]. glistening crystals. Salt: B'HCl: plates.
Amido-p-xylsnol C₂H₂(CH₃)₂(NH₂)(OH)

[1:4:3:6]. [242°]. Whito scales. Formod by reduction of nitroso-p-xylenol (phlorone-oxim) with tin and HCl (Goldschmidt a. Schmid, B. 18, 570; Sutkowski, B. 20, 979). CrO, oxidises it nearly quantitatively to phlorone.

Salt: B'HCl: white orystals.

AMINES. An amino is a body obtained by displacing hydrogen in ammonia by one or more alcohol radicles. They may be divided iuto mono-, di-, tri-, and tetra- amines according as it is considered that their molecule is derived from one, two, three, or four molecules of ammonia. Monamines are spoken of as primary, secondary, or tertiary, according as one, two, or three of the atoms of hydrogen in the molecule of ammonia is held to have been displaced by one or more alkyls. If part of the hydrogen has been displaced by an acid radiols (alkoyl) The reaction is completed by heat, and the

and part by an alcoholic radiole (alkyl) the product may be viewed either as an amide or as an amine, thus NMeAoH may be called msthylacetamids or acetyl-methylamine. In this dictionary the latter name will be used, such dorivatives being described under the amines from which they may be hold to bs derived. Fatty amines are amines in which the nitrogen is attached to carbon that does not form part of a ring; aromatic amines are bases in which the nitrogen is attached to carbon in a benzeno nucleus. In addition to these thore are amines, such as pyridino and quinoline, in which the nitrogen itself forms part of a ring, and also others in which the nitrogen is united to carbon in rings other than that peculiar to benzene.

Formation.—1. By the action of ammonia on the ethers of inorganic neids. The iodides, bromides, and ohlorides of fatty, but not of aromatic, alkyls, combine with ammonia and with the amines (Hofmann, T. 1850, i. 93; 1851, ii. 357):

 $NH_9 + EtI = NEtII_9I$ $NEtH_2 + EtI = NEt H_2I$ NEt,II + EtI = NEt,III $NEt_1 + EtI = NEt_1I$.

The fatty alkyl iodides also act upon the hydriodides of the amines, in presence of ammonia:

 $NH_3 + NEtH_1I + EtI = NEt_2H_2I + NH_4I$ NH3 + NEt, ILI + EtI = NEt, HI + NH, I NH, + NEt, HI + EtI = NEt, I + NH, I.

It is therefore impossible to prepare a purs base by this method; methyl iodide gives chiefly NMo.I, while ethyl iodide gives chiefly NEtH,I, but in the easo of primary iodides, whatever proportions are taken, the entire series of salts is formed. Isobutyl iodide does not form the quaternary iodide, secondary butyl iodido forms hardly any tri-butylamine, but only mono- and di-butylamine, while tertiary butyl iodide is split up by NH, into isobutylene and HI. NEt, at 100° splits up isopropyl iodide and tertiary butyl iodide forming NEt, III and olefine (Hofmann, B. 7, 513; Reboul, C. R. 93, 69). Secondary propyl, hexyl, and octyl iodides form only mono-amines when heated with ammonia (Jahn, M. 3, 165). In the action of alkyl chlorides upon aqueous NH,, the higher the molecular weight, the less primary amine is formed (Malbot, C. R. 104, 998).

Ammouium iodide can be separated by its insolubility in alcohol. The compounds NR, HI, NR,H,I, and NRH,I are decomposed by KOHAq with formation of KI and NR, NR, H, or NRH, respectively, while tetra-alkylated ammonium iodides are not affected. The following method may be employed in the preparation of fatty anines (Hofman, B. 3, 776). The alky jodine, lkI, is heated with alcoholic NII, at 100°; the product is filtered from NH₁I, evaporated, and distilled with potash. NR,I remains behind. The distillate, dried by means of solid KOH, is cooled and treated with oxalic ether which Is slowly added. The following reactions then ocour:

EtO.CO.CO.OEt + 2NRH2 = HRN.CO.CO.NRH + 2HOEs EtO.CO.CO.OEt + NR2H= EtO.CO.CO.NR₂ + HŌEt.

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tertiary base, NR, which does not react with oxalio ether, is distilled off. The residue is well cooled and the solid di-alkyl oxamide separated from the liquid di-alkyl-oxamio ether by pressnre. The latter is purified by washing with water. Boiling potash liberates the alkylamine from the di-alkyl oxamide and the di-alkyl-annine from the di-alkyl-oxamic ether.

When the halogen is situated in a benzene nuclcus ammonia cannot effect its displacement by amidogen unless other chlorous groups are also present in the nucleus. Thus o- and p-chloro-nitro-benzene (but not m-chloro-nitrobenzene) are converted into nitro-anilines by

alcoholic NH₃ at 100°. Primary monamines may be prepared by

acting with KOII on the alkyl ammonium sulphates (Morrison, Pr. E. 28, 693): $MeNH_{1}SO_{4} + 2KOH = K_{2}SO_{4} + MeNH_{2} + 2H_{2}O.$

2. By boiling alkyl cyanates with potash (Wurtz, C. R. 28, 223), thus: EtNCO + H₂O = EtNII, + CO. The primary bases prepared by this reaction may be contaminated with secondary and tertiary bases. This occurs when the potassic cyanate used to prepare the alkyl cyanates contains cyanide (Silva, C. R. 64, 299).

3. Similarly, from thiocarbinides and H₂SO₄: EtNCS + H₂O - EtNH₂ + COS.

4. By the reduction of nitro compounds: $RNO_2 + 3H_2 = RNH_2 + 2H_2O$. This reaction is chiefly used in the aromatic series, inasmuch as it is easy to prepare nitro derivatives of compounds containing a benzene nucleus.

The following reducing agents may be used:

(a.) Alcoholic ammonium sulphide. compound is dissolved in alcohol, saturated with NH, and H2S is then passed in. The solution is boiled, filtered from S, acidified, and evaporated; a salt of the base is then left: $C_6H_5NO_2 + 3H_2S =$ C.H.NH. + 2H.O + S. (Zinin, A. 44, 283). This method is especially useful in reducing nitro-azo compounds which would give hydrazo contpounds if reduced in acid solution.

Substances containing several nitroxyls nsually have only one of them reduced when

treated in this way.

(b.) Zinc dust may be used either alone, by mixing the substance with it and distilling, or it may be used in conjunction with water or aqueous potash: $Zn + 2KOII = K_2ZnO_2 + II_2$.

(c.) Ferrous sulphate and aqueous ammonia are used in reducing unsaturated and unstable

compounds.

(d.) Arsenious acid and NaOHAq.

(e.) An acid and a metal. For the acid, HClAq or HOAc is used; for the metal, zinc, tin, or iron, is taken. Tin and HClAq give, as a rule, the best results. A mixture of zine and in is as effective as pure tin, for the zinc pps. the tin as fast as it dissolves. The amount of acid used may sometimes bo very small; thus, in the preparation of aniline, the action seems to be:

4C,H₃NO₂+4H₂O + 9Fc = 4C₆H₃NH₂+3Fe₅O₄.
(f.) Stannous Chloride. In reducing with SnCl₂ and HCl the resulting SnCl₄ sometimes chlorinates the product; thus o nitro-tolnene gives chloro-o-toluidine. The nitroxyls of polynatro derivatives may be reduced one by one by adding to their cold alcoholic solution the calculated quantity of SnCl, dissolved in alcohol

saturated with HCl. In the case of dinitro-toluene C.H.Me(NO2)2 [1:2:4] the nitroxyl in the o position is first reduced, forming CaH, Me(NH2)(NO4) [1:2:4]; whilst alcoholie ammonium sulphide reduces the nitroxyl in the p position, forming C.H.Me(NO2)(NH2) [1:2:4] (Anschütz a. Heusler, B. 19, 2161)

(g.) Hydric iodide solution, alone, or with

addition of phosphorus.

5. By the reduction of nitriles (Mendius, A. 121, 229): $CH_3CN + 2H_2 = CH_3.CH_2.NH_2$

The reduction is effected by Zn and dilute H.SO, but it is slow, and a great deal of nitrile is saponified: CH, CN + 2H, O = CH, CO, NH, 6. Primary bases are instantly formed when

carbamines are treated with acids:

 $CH_3NC + 2H_2O = CH_3NH_2 + HCO_2H$.

7. By boiling bromo-amides with aqueous NaOH. If bromine and potash be simultaneously supplied to an amide, a potassium bromo-amide, X.CO.NKBr, is formed. If this compound be treated with silver carbonate, an alkyl cyanate is produced: X.CO.NKBr = KBr + X.N.CO.

When this cyanate is boiled with potash an alkylamine is formed (by Formation 2). Thetwo last stages may be performed simultaneously by boiling the potassium bromo-amide with

aqueous NaOH.

The operation is conducted as follows:

Bromine is mixed with its equivalent of amide, and a 10 p.c. solution of potash is added till the colonr of the bromino has nearly disappeared.

 $X.CO.NII_2 + Br_2 + 2KHO =$ X.CO.NKBr + KBr + 2H,O.

Three equivalents of potash dissolved so as to form a 30 p.c. solution are now heated to 70° in a rctort, and the first solution is added gradually through the tubulus. Finally the whole is distilled, and the base collected in a receiver containing hydric chloride. A mixture of ammonium chloride and the hydro-chloride of the base is thus got; they may be separated by alcohol, which does not dissolve the former (flofmann, B. 15, 765).

8. Amides can be converted into amines by heating with alcohols: thus acctamide and cthyl alcohol give ethylamine acctate CH₃.CO.NH₂+HOEt=CH₃.CO.ONEtH₃, while ethyl-acctaniide and cthyl alcohol give diethyl-

amine acetate (Baubigny, C. R. 95, 646). CH, CO.NEtH + HOEt = CH, CO.ONEt.H. Sodium alcoholates act similarly (Seifert, B.

18, 1355) X.NH.CO.Y + NaOR = X.NH.R + NaO.CO.Y.

9. From amido-acids by heating alone or with baryta : $C_6H_4(NH_2)CO_2H = C_6H_3NH_2 + CO_2$.

10. From alcohols or phenols by displacing hydroxyl by amidogen. Ethyl and methyl alcohols give a little ethyl- and methyl-amine when heated with NH,Cl at 300° (Weith, B. 8, 459). Similarly, phenols produce small quantities of amines when heated with NH,; this reaction takes place very readily in the naphthalene and anthracene series. Ortho- and para-, but not meta-, nitro-phenols are converted by aqueous ammonia into nitranilines (Merz a. Riz, B. 19, 1749). The reaction takes place more readily when the alcohols are heated at 260° with the compound ZnCl₂NH₂ or CaCl₂NH₄ (Merz a. Weith, B. 13, 1300; 14, 2343; Merz a. AMINES.

Gaslorowski, B. 17, 628; Mers a. Buoh, B. 17, 2634). Ammonia-zine-chloride converts phenol into aniline; aniline-zino-chloride acting upon phenol gives di-phenylamine. S-naphthol is converted by heating with NH, into naphthylamino, but by ammonia-zinc-ohloride into di-8naphthyl-amine. Fatty alcohols act differently upon aromatio bases in presence of ZnCl., the alkyl entoring the nucleus; thus aniline-zinc-chloride and alcohol produce amido-phenylethane:

 $C_6H_4NH_2 + HOEl = C_6H_4EtNH_2 + H_2O.$

11. By roduction of the phenyl-hydrazides of the aldehydes and kotones in alcoholic selution by sodium amalgam and acctio acid (Tafel, B. 19, 1924):

RR'C: N, HPh + 2H, = RR'CH.NH, + PhNH,..

12. By reduction of aldoxims and ketoxims in alcoholio solution by sodium amalgam and acetic acid (Goldschmidt, B. 19, 3232):

 $RR'C:NOH + 2H_2 = RR'CH.NH_2 + H_2O.$

13. From sulphonates by heating with sodamide (Jackson a. Wing, B. 19, 902):

 $R.SO_3K + NaNH_2 = R.NH_2 + NaKSO_3$

Properties .- Most amines are volatilo or can be distilled alone or with the aid of steam. Primary bases in which amidogen is not united to earbon in a benzeno nucleus turn red litmus paper blue and combine with carbonic acid; aniline and its homologues are neutral to litinus, and do not combine with carbonic acid. Ammonia pps. the amines from cold aqueous solutions of their salts; but at high temperatures the amines expel NH, from its salts. The relative saponifying power of amines has been studied by Ostwald (J. pr. [2] 35, 112). If a mixture of aromatio bases is dissolved in an excess of glacial acctic acid, and the solution is diluted with three times its volume of water and then boiled, the primary amines remain in solution while the acetates of secondary and tertiary amines are decomposed and the bases are found on the filter (Michael, B. 19, 1391). To determine whether a given base is primary, secondary, or tertiary, it is heated with methyl iodide until a quaternary iodide is formed; this iodide is known by its stability towards potash. The original base and the animonium iodide are both analysed. If the ammonium iodide differs in composition by containing CH,I more than the base, then the base was tertiary. If it differ by C.H.I, this shows that the original base was secondary, and had to exchange hydrogen for methyl before it could become tertiary. If the iodide contains C.H.I more than the base, then the latter was primary.

Reactions I, 2, 3, 4, 5, 6, 11, I2, I5, 20, 27, 28, may also be used to distinguish between primary, secondary, and tertiary bases.

When a quaternary ammonium base is distilled, if it contains ethyl it splits up thus: $NRR'R''C_2H_3(OH) = NRR'R'' + C_2H_4 + H_2O;$ (Hofmann, B. 14, 494).

Quaternary ammonium chlorides containing methyl split off MeCl on distillation:

NRR'R''MeCl = NRR'R'' + MeCl;

(Lossen, A. 181, 877).

Vol. L

Reactions 3, 5, 6, 12, 13, 26, and 28, serve to distinguish o-diamines from m- and p- diamines. Reactions. -1. If a primary base be boiled with alcoholic potash and chloroform the dis-

gusting odour of the corresponding carbamine will be noticed (Hofmann, B. 8, 767)

EtNH, + CHCl, + 8KHO = EtNC + 3KCl + 8H,O. 2. If a primary fatty base be dissolved in alcohol mixed with an equal volume of CS, and the liquid be boiled down to half its volume, a thiocarbamate will be formed:

 $2RNH_2 + CS_2 = RNH.CS.S.NRH_3$.

If the liquid be now boiled with a little aqueous mercurie or ferric chloride a pungent odour of an alkyl mustard wil (or thiocarbimide) will be perceived, thus: RNH.CS.SNRH₃ + $HgCl_2$ = $HgS + RNCS + NRH_3Cl + HCl$. In the aromatio series the product of the action of alcoholic CS is usually a thio-urea which requires to be treated with P2O3 in order to get the thiocarbimido (Hofmann, B. 3, 768; 8, 107; Weith, B. 8, 461). Mesidine and amido-penta-methyl-benzeno give thio-carbimides in addition to smaller quantities of the thie-ureas (Hofmann, B. 18, 1827).-3. Nitrous acid converts primary fatty amines into alcohols: $RNH_2 + HNO_2 = ROH + N_2 + H_2O$. It converts primary aromatio amines into diazecompounds: RNH. + HNO. = RN.OII + II.O. converts all secondary bases into nitrosamines, which are neutral substances, volatile with steam: $RR'NH + HNO_2 = RR'N.NO + H_2O$. cannot act upon tertiary fatty bases, except with elimination of an alkyl. It converts most tertiary aromatio bases into nitroso derivatives, which still possess basic properties:

 $C_6H_5NMc_2 + HNO_2 = C_6H_1(NO)NMe_2 + H_2O_4$ Aromatio nitrosamines are converted into p-nitroso derivatives under the influence of acids. $C_6H_5NMe(NO) = C_6H_4(NO).NMeH.$ means of the preceding reactions, nitrous acid may be used to separate secondary from primary and tertiary bases, for the nitrosamines do not combine with acids, and may therefore be extracted from the acid solution by ether, or by distilling with steam: and on reduction they give the secondary base. If the diazo compounds are hoiled with water phonels are formed: $RN_2Cl + H_2O = ROII + N_2 + IICl$, while if they are boiled with alcohol, the amidogen is usually displaced by hydrogen: RN_CI+C_H_O=RH+IICI+C_H_O. Frequently, however, boiling with alcohol displaces amidogen by ethoxyl: $C_{u}Me_{i}H.N_{2}Cl + HOEt = C_{u}Me_{i}H.OEt + HCl + N_{2}$ (Hofmann, B. 17, 1917). Amidogen may also be displaced by hydrogen by reducing the diazo compound to a hydrazino and boiling the latter with aqueous CuSO₄ (Haller, B. IS, 90). In order to displace amidogen by chlorine we may distil the platinochlorido of the diazo derivative; to displace amidogen by bromine we may boil the perbromide of the diazo derivative with alcolol; to displace it by iodine we may boil the diazo salt with aqueous HI or KI. These operations may be more conveniently performed by the method of Sandmeyer (B. 17, 1633, 2650). This method consists in boiling the diazo compounds with ouprous chloride, bromido, iodide, or cyanide.

Examples.—(a) 4 g. m.nitro-aniline, 7 g. HCl (S.G. 1.17), 100 g. water, and 20 g. of a 10 p.c. solution of cuprous obloride in HClAq are heated to near boiling and 2.5 g. sodie nitrite dissolved in 20 g. water are slowly added, the mixture being well shaken. 4 g. pure m-chloro-nitro-

benzene is obtained.

(b) 12.5 g. crystallised cupric sulphate, 36 g. KBr, 80 g water, 11 g. H2SO4 (S.G. 1.8), and 20 g. copper turnings are boiled until the dark colour has nearly dicappeared. Aniline (9.8 g.) is now added, and the boiling liquid treated as before with NaNO, (7 g.) dissolved in water (40 g.). Bromo-benzene passes over en subsequent dietillation.

(c.) 25 g. crystallised CuSO, 150 g. water and 28 g. KCN (96 p.c.) are dissolved in hot water. A solution of diazobeuzene chloride is run in, this is prepared from 7 g. NaNO₂ dissolved in 20 g. water added to a solution of 9.3 g. aniline in 20.6 g. HCl (S.G. 1.17) and 80 g. water. The yield of benzonitrile is 63 p.c. of

the theoretical.

In these reactions a double compound between the cuprous salt and the diazo salt is perhaps an intermediate body. Such a double compound has been isolated in the case of 8-naphthylamine, C₁₀H,N₂BrCu_Br₂ (Lellmann a. Remy, B. 19, 810). Substitution of amidogen by halogens may also be effected by gradually adding IINO, to a hot solution of the amine in IICl, HBr, or HI (Losanitsch, B. 18, 39).

Amidogen may be changed into SH by heating the diazotised base with warm alcoholic potassium sulphide. By oxidising the resulting mercaptan with KMnO, a sulphonic acid is got

(Klason, B. 20, 349).

Nitrous acid serves to distinguish o-, m-, and p. diamines (v. di-Azo-compounds).

4. Benzoyl chlorids acts on primary and secondary antines:

 $RNH_2 + BzCl = RNBzH + HCl$

RR'N11 + BzCl = RR'NBz + HCl(Hofmann, B. 5, 716; Hallmanu, B. 9, 846) Tertiary aromatic amines heated with it at 200° may exchange alkyl for benzoyl:

 $NPhEt_2 + BzCl = NPhEtBz + EtCl$

(Hess, B. 18, 685).

5. Acetyl chloride converts primary and secondary amines into acetyl derivatives.

The di-alkylated tertiary aromatic amines readily allow one of the alkyl groups to be replaced by acetyl when treated with acetyl bromide, the alkyl bromido formed converting another portion into quaternary ammonium bromido: 2XNR₂+AcBr=XNRAc+XNR₃Br.

The reaction sets in spontaneously, and is completed on gentle warming (Staedel, B. 19,

1947).

Primary aromatic amines may be converted into acctyl derivatives by boiling not only with AcCl or Ae,O but oven with glacial HOAc.

The alkoyl derivatives of c but not of m and p aromatic diaminee givo rise to anhydro compounds:

C.H. $\stackrel{\text{NH.CO.CH}_2}{\text{NH}_2} = \text{H}_2\text{O} + \text{C.H.}_4 \stackrel{\text{NH}}{\sim} \text{C.CH.}_3$ (Hübner, A. 208, 278).

6. Aldehydes form producte of condensation with amines. Enanthol is recommended by behiff (A. 159, 158) as a means of distinguishing between the different classes of amines. Primary amines require one equivalent of cenanthol:

 $C_7H_{14}O + PhNH_2 = PhN(C_7H_{14}) + H_2O_7$ while eccondary amines require only half as

much aldehyde:

 $C_1H_{14}O + 2Me_2NH = (Me_2N)_2(C_1H_{14}) + H_2O_2$ The base is dissolved in benzene and a standard

solution of cenantnot in pensens is run, in as long as it produces further separation of drops of water.

Tertiary aromatic amines can also condense with aldehydee: $2PhNEt_2 + [1:2]C_6H_4(NO_2)$. CHO = $NO_2.C_6H_4.CH(C_0H_4NEt_2)_2 + H_2O.$ Arematio amines heated with aldehydes and ZnCl, give tri-substituted methanes (Fischer, B. 15, 676).

In order to distinguish whether an arcmatic diamine is an ortho compound, Ladenburg (B. 11, 600) heats its hydrochloride with benzoic aldchyde; if the compound is ortho an aldehydine (q. v.) is formed and HCl is ovolved, while no HCl ie evolved in the case of m or poompounde.

 $R''(NH_2HCl)_2 + 2PhCHO =$ R"(N,C,H,Ph,)HCl + 2H,O + HCl.

7. Bromine and aqueous potash convert primary amines into di-bromamines: $MoNH_2 + 2Br_2 + 2KOH = MeNBr_2 + 2KBr + 2H_2O$. Secondary amines, containing one divalent alkyl,

behave similarly: $\mathbf{C_sH_{14}NH} + \mathbf{Br_2} + \mathbf{KOH} = \mathbf{C_sH_{14}NBr} + \mathbf{KBr} + \mathbf{H_2O}$ but secondary amines coutaining two monovalent alkyls are split up into an alkylene bromido and

a primary alkylamine (Hofmann, B. 16, 559). The di-brome-amines containing hexyl and its higher homologues are split up by aquecue NaOII iuto HBr and nitriles:

 $C, H_1, CH_2NBr_2 + 2NaOH =$

C, H, CN + 2NaBr + 2H, O.

Hence amides may be converted first into aminee and then into nitriles by treatment with bromine and NaOIIAq, the first reaction being:

 $\begin{array}{l} C_{7}H_{12}CII_{22}CO.NII_{2}+Br_{2}+4NaOH=\\ C_{7}H_{12}CII_{22}NH_{2}+2NaBr+Na_{2}CO_{3}+2H_{2}O \end{array}$ (Hofmann, B. 17, 1920).

8. Sulphuric oxide combines with primary and secondary fatty amines, forming small quantities of sulphamic acids:

 $NEt_{2}H+SO_{3}=NEt_{2}\,SO_{3}H.$ It also combines with tertiary fatty amines:

 $NEt_3 + SO_2 = Et_3N < \frac{SO_2}{O} >$ (Beilstein a. Wiegand, B. 16, 1261).

It combines with arematic amines forming sulphonic acids:

 $C_6H_5NH_2 + SO_3 = C_6H_4(SO_3H)NH_2$ Aromatic amines may also be sulphonated by

H₂SO, and by ClSO,II.

9. Sulphuryl chloride acts upon secondary fatty amines thus, forming tetra-alkyl eulpli-amides: $SO_2Cl_2 + 2HNEt_2 = SO_2(NEt_2)_2 + 2HCl$. With the hydrochlorides of these bases the reaction stops half way: SO₂Cl₂+ HCl.NHEt₂= Cl.SO₂.NEt₂+2HCl (R. Behrend, A. 222, 116).

10. Zinc ethide does not attack tertiary amines, but acts upon primary and secondary

amince in the following ways:

 $2RNH_2 + ZnEt_2 = R_2N_2H_2Zn + 2HEt_4$ $2RR'NH + ZnEt_2 = R_1R_2R_2 + 2HEt$ (Frankland, Pr. 8, 504; Gal, C. R. 96, 578).

11. Cyanic ethers unito with primary and secondary bases forming alkyl-ureas

RNH2+R'NCO=RNH.CO.NR'H RR''NH + R'NCO = RR''N.CO.NR'H.

Cyanic acid acte similarly:

RNH2HCl+KNCO=KHN.CO.NH2+KCl 12. Thio-carbimides unite with primary and secondary amines forming thio-ureas: RNH, + R'NCS = RNH.CS.NR'H.

When the solid product obtained by boiling an

aromatic di-amine with alcohol and oil of mustard (O.H.NCS) is gradually heated, then if the diamins were o it would solidify above its melting-point; if it were m it would melt without further change and would therefore solidify on oooling; if it were p it would melt and undorgo decomposition and on cooling would remain liquid (Lellmann, A. 221, 1; 228, 248; B. 19, 808). In all cases di-thio-ureas, R"(NH.OS.NIIC, II,)2, are first formed; those from m-diamines are not affected by heat, while the o and p dorivatives split up on melting thus:

$$R''(NII.OS.NHC_3H_5)_2 = R'' < \frac{NH}{NH} > CS + CS(NHC_3H_5)_2.$$

18. The di-sulphocyanides of the o-diamines are changed at 120°-130° into thio-ureas, $C_xH_y <_{NH}^{NH}>CS$, which are not dosulphurised by hot solution of PbO in NaOHAq. The m and p diamines give compounds of the form $C_xH_y(NH.CS.NH_z)$, which are desulphurised by this reagent (Lellmann, A. 228, 8, 248).

14. Small quantities of orthodiamines are readily detected by adding a few drops of a hot acetic acid solution of phenanthraquinone to an alcoholic solution of the substanco; if an orthodiamino is present a yellow crystalline pp. of the corresponding quinoxalino is formed en boil-ing the solution; this pp. in the case of phenyleno and tolyleno o-diamines is coloured deepred by HCl (Hinsberg, B. 18, 1228).

15. If a mixture of bases is treated with sufficient citraconic acid to form the acid salts and the aqueous selution is boiled the primary amines will be ppd. in the form of alkyl-eitraconamic acids, while the secondary and tertiary amines can be obtained by distilling the filtrate with steam (Michael, B. 19, 1390).

 Oxidising agents convert aromatic amines into azo or azoxy compounds. Hence the nitration of such amines by the usual methods regnires previous introduction of acctyl into the amidogen. But by treating the nitrates with cold cono. H.SO, nitro-amines may be prepared, the nitroxyl taking a m position with regard to amidegen (Levinstein, D. P. J. 256, 471).

17. Amines form condensation products with quinones, e.g.

C₆H₂O₂(NHPh)₂; C₆H₂O(NPh)(NHPh)₂

18. Aromatic amines when boiled with fatty amides produce ammonia and alkyl-amides, e.g. OH, CO.NH, + NPhH, - CH, CO.NPhH + NII.

19. Silver salts form additive compounds with aminos (Mixter, A. C. J. 1, 239).

20. The ferrocyanides are obtained by adding the amines to a mixture of aqueous K4FcCy4 and hydrochloric acid; a crystallino pp. of the acid ferrocyanide B'_H, FoCy, xaq, is usually formed (Fischer, A. 190, 184; Eisenberg, A. 205, 265). The ferrocyanidos of tertiary amines are particularly insoluble in water and may be used as a means of isolating those amines. To reoover the amine, the pp. is suspended in water and decomposed by CuSO, and the excess of OuSO removed from the filtrate by baryta.

21. Ohloral hydrate heated with tertiary

aromatic aminss and ZnCl2 forms a condensation product which, when decomposed by aqueous KOH gives an aldshydo dsrivative. ...

 $\begin{array}{c} \text{COl}_{2}\text{CH}(\text{OH})_{2} + \text{C}_{e}\text{H}_{2}\text{NMs}_{2} = \\ \text{COl}_{2}\text{CH}(\text{OH}).\text{C}_{e}\text{H}_{2}\text{NMe}_{2} + \text{H}_{2}\text{O} \end{array}$ $OCl_3CH(OH).C_6H_4NMe_2 = CCl_3H_+HCO.C_6H_4NMe_2$

(Boessneok, B. 18, 1516; 19, 365).

22. Primary aromatic amines in alcoholic solution absorb cyanogen; and the product when boiled with glacial HOAo becomes a di-alkyloxamide:

2PhNH, + C,N, = PhNH.C(NH).C(NH).NPhH PhNH.C(NH).C(NH).NPhH + $2H_2O =$

PhNII.CO.CO.NPhH + 2NH, Aromatic o-diaminos act similarly; the resulting oxalyl-o-diamine may be viewed as a di-oxyquinoxaline (Bladin, Bl. [2] 42, 104).

23. Primary aromatic amines heated with glyccrin (or acrolein), H2SO,, and nitrobenzone (as oxidising agent) produce bases of the quineline series. A similar reaction occurs when paraldehyde) is substituted for glycol (or glyccrin.

24. Nitric oxide passed into an alcoholio solution of tertiary aromatic bases produces azocompounds of the form RR'N.C, H, N2.C, H4.NRR'.

25. For the action of aceto-acetic ether.

v. p. 19.

26. Orthodiamines form orystalline compounds with glucose (Griess a. Harrow, B. 20, 281).

27. Diazobenzene chloride reacts with primary and with secondary amines, forming diazo-amides (v. di-azo-compounds). In the case of the secondary amines the compounds Calla. N. NRR', being easily crystallised and sl. sol. water, may be conveniently used in separation of these bases from tertiary and in some cases from primary bases (Wallach, A. 235, 235).

28. A solution of potassium croconate gives with salts of o-diamines dark-coloured pps. consisting of the corresponding azines (Nietzki, B.

19, 2727).

29. Tertiary aromatic amines form condensation products when heated with arematic acids or alcohols in presence of ZnCl2 or P2O3; water being eliminated at expenso of H para to N (Fischer, A. 206, 85).

OTHER REACTIONS of the amines are described in articles on the several bases, e.g. METHYLAMINE, ETHYLAMINE, ANILINE, PHENY-LENE-DIAMINE. Soe also Amides, Amido Acids, Amidines, and Amidoxims.

AMISATIN v. ISATIN.

AMMELIDE C₃II,N₀O₂ or C₂H₂N₂O₄ i.e. C₂N₃(NH₂)(OH)₂ (?). Melanurenic acid. 'Amidocyanuric acid.' Mono-amide of cyanuric acid. Liebig mado a distinction between ammolide and melanurcnio aoid, but his ammelido was a mixture of ammeline and his melanurenie acid, hence it seems best to transfer the name am melide

to melanuronio acid (Klason, J. pr. [2] 33, 295).
Formation. — 1. From melam and conc.
H₂SO, (Liebig, A. 10, 30; Gabriel, B. 8, 1165; Jager, B. 9, 1554) .- 2. From melamand boiling oono. KOHAq er oono. H₂SO₄ at 150° (K.).— 3. From anmelino and oono. H₂SO₄ at 160° or by heating ammeline nitrats (Knapp, A. 21, 241); the change is incomplete.—4. A product of the dry distillation of urea (Liebig a. Wöhler, A. 54, 371; Laurent a. Gerhardt, A. Ch. [2] 19, 93; Drechsel, J. pr. [2] 11, 289).—5. Among products got by boiling mellon potassium with

aqueous KOH (Hennesberg, A. 78, 246; Liebig, A. 95, 269).-6. From ite ethers or their thioderivatives by gentle heat (K.).—7. From thio-ammelide and KMnO, (K.).—8. From eyanogen bromide and cyanamide at 100° (Ccch a. Dehmel, B. 11, 25).—9. From urea and cyanogen iodido at 150° (Poensgen, A. 128, 339; Hallwachs, A. 153, 294; Schmidt, J. pr. [2] 5, 86) .- 10. In small quantity by action of COCl. on NH, (Bouchardat, A. 154, 355).-11. In small quantity from di-cyan-di-amide by heating with water at 160° or with aqueous ammonio carbonato at 120° (Bambergor, B. 16, 1078, 1703).

Preparation.—Conc. H.SO, (300 g.) ie slowly poured upon melam (100 g.) and the solution heated for a few minutes to 190°. When cold it is ponred into a litre of water, when ammelido sulphate slowly crystallises (Strieglor, J. pr. [2]

33, 163).

Properties .- White crystalline powder; v. sl. sol. water, insol. usual menstrua, sol. mineral acids, insol. acetic acid, v. sol. ammonia. It does not separate when its solution in warm aqueous NaOH is cooled (difference from ammeline). May be crystallised from boiling water. Not attacked by Cl, Br, HI, or AcCl.

Reactions.—1. Boiling dilute acids or alkalis form NH, and cyanurio acid. Baryta-water does not effect this change. - 2. Phosphorus pentachloride forms Cy,Cl, -3. KMnO, in acid solution forms cyannric acid.-4. Water at 170° forms CO2 and NH3.-5. Heated in a current of

moist CO₂ it forms cyanamide.

moist CO₂ it forms cyanamide.

Salts.—(H₂A" = C₁H₁N₃O₄).—
H₂A"H₂SO₄3aq.—H₂A" 2HNO₃.—H₂A" 2HCl.—
Na₂A" 6aq.—Na₄HA" 5aq.—K₂A".—KHA".—
(NH₁).A" 5aq.—(NH₃)HA"1½aq.—GaA"aaq.—BaA"2½aq.—CuA".—NiA"2aq.—Ag₂A".—AgHA"1½aq (Striegler; Volhard, B. 7, 92).

Dimethyl ther C₂H₂N₄O₂ i.c.
C₃N₃(NH₂)(OMe)₂. (212°]. Formed by action of ammonia on trimethyl cyanurate, and ocents as a hypoduct in the preparation of that body

as a by-product in the preparation of that body (Hofmann a. Olshausen, B. 3, 273). Plates, sl. col. cold alcohol, v. sl. sol. ether, sl. sol. cold water. - C, H, N, O, AgNO, : needles.

Di-chyl ether C.H₁₂N₁O₂ i.e. C₂N₃(N11.)(OEt)₂. [97^o]. By-product in the action of CyCl on NaOEt, and formed by heating cyanetholin with aqueous NH3 at 100° (H.a.O.). Prisme.—C.H.₁N₁O.AgNO₃: needlee.—(C.H.₁N₁O.),AgNO₃: needlee.—AMMELINE C₃H₅N₂O i.e. C₃N₃(N11.),(OII).

Di-amido-cyanuric acid. Diamide of cyanuric

acid.

Formation.-1. By boiling melam for a long time with KOHAq or ECIAq or by heating it with cone. H₂SO₄ at 100° (Liebig, A. 30, 24; Klason, J. pr. [2] 33, 286). — 2. From Cy₃(NH₂)₂Cl by alkalis (Lament a. Gerhardt, A. Ch. [3] 19, 92).—3. From thio-ammelino and KMnO.—4. From its ethor or its thio derivative by HCl.—5. Formed by boiling the hydrochloride of 'di-amido tri-chloro-methyl cyanidine' (v. Tri-Chloro-Acetonitrile) with NH, Aq:

 $Cy_s(NH_2)_2(CCl_s)HCl + NH_s + H_2O =$ Cy₃(NH₂)₂(OH) + NH₄Cl + HCCl₃. Also by heating tri-chloro-acetonitrile with NH₃Aq at 120°, or with alcoholic NH, at 170° (Weddige, J. pr.

(2) 83, 85).

Properties. - Minute needles in dendritie groupe (when ppd. from a warm solution). Insol. water, alcohol, ether, and benzene; eol. mineral acide; insol. acetic acid; sol. NH, Aq. Separatte when ite colution in warm NaOHAq is cooled.

Reactions .- 1. Split np by heat into NH. and mellon.-2. Warm H.SO, forms NH, and ammelide .- 3. Boiling dilute nitric acid forms first ammelide, then cyanuric acid (Knapp, 4. 21, 255).

Salts.-Its compounde with acide are decomposed by water. B'HCl: prisms. B'HNOs. −B'AgNO₃∙

Ethyl ether Cy₃(NH₂)₂OEt. [190°-200°]. From cyanetholine and NH₃ (Hofmann a. Olshausen, B. 3, 275). V. sl. sol. alcohol.

Chloride Cy,(NH2)2Cl. 'Chloro-cyanamide.' From Cy, Cl, and NH, Aq (Liebig, A. 10, 43; Laurent a. Gerhardt, A. Ch. [2] 19, 90; 20, 98; Bineau, A. Ch. [2] 70, 254). Powder, incol. water. Decomposed by heating with HCl into mellon and NH₃. Dilute KOHAq convorte it into ammeline. NH₃ at 100° forms melamine. KHS forms thio-ammolino.

air.) Mol. w. 17.01. [-75°. Faraday, Q. J. S. 19, 16]. (-38·5°; pressure less than 760 mm. Regnault.) S.G. 3° (liquid) 6234 (Jolly, A. 117, 181: compare also Andreef, ibid. 110, 1). V.D. 8·5. 181: compare ass America, total. 116, 17: 7.53. S. 8. 0°, 1050; 10°, 813; 15°, 727; 20°, 654; (Bunsen, Gasometry, Engl. ed. 169). S. 0°, 1148 (Roscoe a. Dittmar, A. 110, 140); S. 0°, 1270 (Berthelot, C. R. 76, 1041). C.E. (liquid—11°). to 0°) 00155 (Jolly, A. 117, 181). Refractive power (gas) compared with air = 1, 1.309. $[N,11^3] = 11,890$; $[N,H^3,Aq] = 20,320$; $[N11^3,Aq] =$ 8,430 (Th. 2, 68).

Occurrence.- Ammonia salts occur in the atmosphero and in rain water; in many mineral waters; in eea water; near volcanoes; in many soils; in almost all plants; in the excremente of many animals; among the products of the decay of nitrogenous organic bodies. Free ammonia is not known to occur in nature. Ammonia was distinguished from ammonium carbonate by Black in 1756; Priestley obtained it approximately pure and named it alkaline air: Scheele showed it to contain nitrogen; Berthollet demonstrated its composition in 1785. The word ammonia comes from sal ammoniacum, the name given in the middle ages to ammonium chloride.

Formation.—1. By the action of the induction spark, or the silent discharge (Donkin, Pr. 21, 281), on a mixture of N and H in the ratio N:H, a emall quantity of NH, ie produced (Morren, C. R. 48, 432; Perrot, C. R. 49, 204; Chabrier, C. R. 75, 484) .- 2. According to Ramsay a. Young (C. J. 45, 93) a trace of NH, ie formed when a mixture of moist N and H ie passed through a red-hot tube containing iron filings; 3. Ammonium nitrite is formed, a. when hydrogen is burnt in air (Zöller a. Grete, B. 10, 2145; but against thiev. Wright, C. N. 38, 240); b. by the action of a etrong induction-spark on a mixture of N and H.O (Thengrd, C. R. 76, 983; Johnson, C.N. 48, 258 a. 204). Ammonium chloride is produced when electric sparks are passed for 8 to 10 hours through a mixture of HCl gas, N, and H, the elements being in the

ratio N:H. (Deville, C.B. 60, \$17); or by passing the same gases through a red-hot porcelain tube confaining a metal tube cooled by a stream of cold water (Deville, A. 135, 104) .-4. By the action of a porous body—e.g. spongy platinum, pumice, ferrio oxide—aided by heat, on a mixture of H with an exide of nitrogen or HNO, NH, is produced.—5. By decomposing a compound of H and one of N together, NH, is formed: e.g. by the action of water on nitride of Si, B, Mg &c., SiO2, B2O3, or MgO is produced, and the N and H combine to form NII, again moist NO passed over hot iron filiugs yields NII_x. 6. By strongly heating easily oxidised bodies - e.g. As, Zn, K, &c. - with alkaline oxides, in presence of air .- 7. By strongly beating metallic nitrates or nitrites with hydroxides of the alkali or alkaline earth metals and iron filings or zinc .-8. By heating metallie cyanides with steam (v. Marguerite and Sourdeval, D. P. J. 157, 73 and 316) .- 9. By heating solutions of nitrates or nitrites with KOIIAq and Zn or Fo, or with a Cu-Zn couple. Ammonium sulphate is formed when nitrio acid is dropped into a vessel containing Zn and diluto II 2SO Aq (Kuhlmann, A. 64, 233).-10. By the action of water on chloride, iodide, or phosphide, of nitrogen, or on the amides; in the last cases it is often necessary to use solutions of KOH or NaOH .-- 11. By the dry distillation of many nitrogenous organio bodies-e.g. horn, bones, blood, coal, &c. Ammonia is produced, according to Johnson, when N aud H are passed over spongy Pt (C. J. 39, 128); but this is denied by Wright (C. J. 39, 359), whose experiments seem to prove that the NII, obtained by Johuson was the product of the mutual action of a trace of NO (iu what was supposed to be pure N) and H, in presence of the spongy Pt (but v. also Johnson's pamphlet Elementary Nitrogen, and on the Synthesis of Ammonia [Churchill, 1885]).

Preparation.-1. By gently heating a mixture of 1 part chloride or sulphate of ammonium with 2 parts finally powdered slaked lime; the mixture is covered with a layer of lime to absorb water, and the gas is dried by passage through a oylinder containing lime in small pieces. - 2. By gently heating a solution of CaCl2 in NII, Aq previously saturated with NII,; this mixture may be kept unchanged for long.-8. Pure ammonia is prepared by Stas (I'r. 6, 423) by one of the following methods:—(i.) From pure NH Cl and KOHAq; 10 litres of a boiling cone, solution of NII CI are mixed with 1 litre HNO3Aq, S.G. 14; the boiling is continued so long as Cl comes off, the NII, Cl which separates on cooling is dissolved in hot water, and again boiled with $\frac{1}{10}$ volume of HNO, till Cl ceases to come off, water is then added. and NH, is obtained by decomposing by KOHAq. (ii.) From pure (NII,),SO, and KOHAg; 2 kilos. of (NH₄),SO₄ are heated with 1½ kilos. cone. H2SO4 to the temperature whereat the sulphate begins to decompose with effervescence, small quantities of nitric acid are then added until the liquid becomes quite colourless; the salt which crystallises on cooling is dissolved in warm water and decomposed by KOHAq. [The object of these treatments is to remove the small quantities of substituted ammonias-NH,CH, NH,C,H, &c. - which are present in ammonium ohloride and sulphate.] (iii.) From pure KNO, by the action of Zn and Fe in presence of KOHAq: the KNO, is prepared by heating 1 kilo KNO, with metallic copper, and dissolving out the KNO2 in water; this solution is digested with 15 litres KOHAq-S.G. I 25-31 kilos. granulated zinc free from carbon (Zn obtained by fusing commercial Zn with 5 p.c. PbO may be used), and ½ kilo. iron wire pre-viously strongly heated in air and then reduced by hydrogen; the liquid is poured off and

distilled with gentle challition.

Properties .-- A colourless, strongly-smelling, gas, which turns red litmus paper blue, and turmeric paper brown. Taste, hot and strongly alkaline; poisonous when breathed; it destroys the mucous membrane. Easily liquefied to a limpid, colourless, highly refractive, liquid; best by heating solid 2AgCl.3NII3 in oue end of a strong glass tube, closed at both ends, and beut to an obtuse angle, the other end being surrounded by snow and salt (Faraday, Q. J. S. 19, 16). The silver compound begins to melt at 38°, it is quito liquid at 90°, begins to boil at 100°, and the change is complete at 112°. Liquefied at -40° to -50° ; this may be effected by passing the well-dried gas through a U tube surrounded by a mixture of crystallised CaCl, and snow, or by liquid SO₂ which is rapidly evaporated by a current of air (Loir and Drion, J. 1860. 41). If liquid ammonia is cooled by solid CO2 and ether in vacuo (Faraday), or by rapid ovaporation over H.SO. (Loir and Drion), white transparent crystals of solid ammonia are obtained, which melt at -75° (Faraday). Liquid ammonia vaporises in a closed vessel, the vapour-pressures according to Regnault (J. 1863. 66) being as follows: -30° 866.09 mm. + 40°

I1,595.30 mm.

15,158-33 -10 2144.62 60 19,482.10 3183.34 70 O 21,675.55 +10 4575.08 80 30,843.09 20 6387:78 90 38,109.22 80 8700.97 100 46,608.24 Ammonia gas is very soluble in water (v. Combinations No. 1), alcohol, and ether; it is largely absorbed by charcoal (v. Hunter, C. J. [2] 9, 76; 10, 649) and other porous substances; it is absorbed by many saline solutions, the quantity of NH, absorbed being, as a rule, the less the more concentrated is the solution (v. Raoult, C. R. 77, 1078). Ammonia solution is a strongly smelling, caustic, alkaline, liquid: at -40° it forms long needle-shaped orystals: at -49° it solidifies to an inodorous mass; the B.P. and S.G. increase the less is the quantity of NH3 prosent. Many metallic oxides insoluble in water are dissolved by NII₂Aq, e.g. CuO, Ag₂O, &c.; aqueous NII, also dissolves many fats and resins. Ammonia resembles PH, in its properties and reactions; it is, however, much more stable and less easily oxidised than that compound; an aquoous solution of NH3, which doubtless contains NH,OH, is

50

-20 1392·13

comp. also the arts. Hydrides, Hydrox-IDES, and NITROGEN GROUP OF ELEMENTS.) Reactions .- I. Liquid ammonia does not react with H2SO, at -65°; dissolves alkali metals

characterised by the properties expressed by the word alkali (v. Ammonium Compounds;

at first with red, then blue, colour; the metals erystallise out nuchanged (Gore, Pr. 21, 140); alkaline earth metals and heavy metals do not dissolve (Sceley, C. N. 23, 169; concerning solubilities of other elements and salts v. Gore l.c.).—II. Ammonia gas. 1. Heat decomposes NH, partially into N and H; when the gas is passed through an iron or porcclain tubo, decomposition begins at about 500°; the nature of the hot surface exerts a most marked influence on the extent of decomposition; the decomposition is, however, never quite complete (Ramsay a., Young, C. J. 45, 88). A spiral of Pt hoated by an electric current also decomposes NII, (Grove, A. 63, 1) .- 2. The electric discharge decomposes NH3 slowly, but induction sparks from a largo Ruhmkorff's coil moro quickly; the decomposition is not quite complete (Deville, A. 135, 104; Buff a. Hofmann, ibid. 113, 132).-8. NH, is decomposed, into N and H, by passago over several metals at 700° or so, e.g. Au, Pt, Ag, Fe, Cu, &c.; some metals, c.q. Ti, combine with tho N; the alkali metals set free 1 of the H producing compounds of the form NH,M: the compound NH, K is decomposed at a red heat giving NK, and NH,; water acts on it to produce KOH and NH3 (r. Potassium) .- 4. Mixed with oxygen and submitted to the electric discharge, NH, NO, and NII, NO, are formed (Carius, A. 174, 31).-5. Mixed with oxygen and heated, NH, burns to H.O. H, and N, if the NH, is in excess; and to II,0, N, and NII, NO,, if the O is in excess (v. Hofmann, A. 115, 283; Heintz, ibid. 130, 102). The flame examined spectroscopically shows characteristic lines, especially one near D (Dibbits, P. 122, 521). 6. Ozone oxidises NH, chiefly to NH, NO, and NH,NO₂ (Carius, A. 174, 31).—7. A platinum wire heated in NH₃ mixed with air produces NII, NO, if oxygen is passed into the NII, red fumes of N oxides are also produced.—8. Annonia reacts with N₂O₃ and Cl₂O₄ to form H_O, N, NH₄NO₂ or NH₄NO₃, and Cl.—9. Metallie oxides reducible by H are usually also reduced by NII, with formation of metal, N, and II.O, sometimes with formation of metallic nitrides. 10. NH3 reacts with many metallic oxides and haloid salts to form compounds, either of NII3 with the metallic salt -- c.g. PtCl2.4NII3, CuSO4.2NH3-or compounds in which part of the H of NH, is replaced, e.g. NH, HgCl (v. Ammonium Compounds; also the several metals). 11. Chlorine, bromine, and iodine react energetically with NH₃ to produce NH₄X(X = Cl, Br, or I), and N. NH₃ combines with cooled I to form a brown liquid which is decomposed by water with production of NH, IAq and explosive iodide of nitrogen [? NI3] (v. NITROGEN).-12. Sulphur absorbs NH,; on heating N is sot free and aminonium sulphide formed (Brunner, D.P.J. 150, 371) .- 13. Carbon heated in a stream of NH, forms NH4CN and H, sometimes also CH₄-14. Boron heated in a stream of NH₄ form BN (v. Boron), and H.-15. NH₄ combines with acids (II,SO, HCl, &c. &c.) to form ammonium salts ((NH₁₎,SO₁, NH₂Cl, &c. &c., q.v.; v. also Combinations, No. 4).—16. With many organic anhydrides NH₈ combines to form tho ammonium salts of amio acids, q.v. NH3 also acts on several inorganic anhydrides and acid chlorides to form bodies more or less analogous to the amio acids; thus with SO40H.Cl an

monia forms NH(SO, ONH),—the NH watta imido-sulphnrio acid NH(SO₂OH),—then the salts of this acid are obtained salts of amike sulphuric, or sulphamic, acid—NH₂(SO₂OH). Again by the aciton of NH₂ on the acid chloride SO₂Cl₂ it is probable that the amide of sulphuric acid—(NH₂),SO₂—is produced. So also NH₂ reaots with CO₂ to produce NH₂(CO₂ONH₄)—the ammonium salt of amidocarbonic, or carbamic, acid. These compounds will be described under the various acids (v. Carbamic acid; Sulphamic acid; Sulphur oxyacids, nitrogen derivatives of; &c.).

III. Ammonia solution. 1. Heat decomposes NH₃Aq, the whole of the NH₃ being removed as gas.—2. Chlorine, bromine, and iodins react as with NH₃ gas; ellorine produces a little NII,ClO₃ (Fresonius, Fr. 2, 59).—3. Reacts with acids to form ammonium salts (v. Combinations, No. 4, also Ammonium compounds).—4. With many metallic salt solutions it reacts (similarly to KOHAq) to form an ammonium salt and an oxide or hydroxide of the metal.—5. Heated with sulphur in a closed tube ammonium polysulphides are slowly formed (Flückiger, J. Ph. [3] 45,453).—6. Heated with selenion in a closed tube, ammonium selenide and selenite are formed; with tellurium ammonium tellurite is produced (Flückiger, Lc.).

Combinations.—1. Ammonia gas dissolves very freely in water, the action is attended with production of heat; $[NH^2, Aq] = 8,130$ (Th. 2, 68); a concentrated solution of NH_3Aq diluted with n H_2O develops $\frac{1270}{n}$ units of heat (Berthelot, A. Ch. [5] 1, 209). Thomsen (Th. 3, 86) gives the following data

The mass of NII, absorbed by water at 0° is not directly proportional to the pressure; for pressures varying from 50 to about 1,000 mm. the mass of NII, is less, and for higher pressures it is greater, than that calculated by Dalton and Henry's law (for data v. Roscoe a. Dittmar, A. 112, 349). As temperature increases the mass of NII, becomes more nearly directly proportional to the pressure, until at 100° the proportion is established (for data v. Sims, A. 118, 315). The S.G. of an aqueous solution of NII, varies from 8844 at 14°, corresponding to 36 p.c. NII, to 9991 (at 14°) corresponding to 2° p.c. NII, (Carius, A. 99, 164). Carius gives the annexed table. Determinations made at 14° C.

A solution containing 32 p.c. NH₃ corresponds with the quantity calculated on the assumption that the liquid consists of the compound NH₄OH.H₂O (v. AMMONIUM COMPOUNDS).—2. Dry ammonium nitrate absorbs NH₃ at all temperatures from -13° to +25° with liquefaction of the salt; heated over 25°, NH₄ is evolved and the substance becomes solid; the liquid at -10° and 760 mm. contains 42°5 grams NH₄ and 100 ann NH₄NO, these numbers agree with those calculated from the formula NH₄NO, 2NH₄; the solid at 28°5 contains NH₄NO, and NH₄ in the proportion NH₄NO₂:NH₄ (Divers, 27. 21, 109;

appoint.	F. 0.	Specific	P. C.	Specific	P. C.	
gravity	NH ₄	gravity	NH.	gravity	NH.	
08844	36.0	0.0120	24.0	0.0800	10.0	
0.8848	35.8	0·9188 0·9189	23.8	0.9520 0.9527	12·0 11·8	
0.8852	.85.6	0.9145	23.6	0.9534	11.6	
0.8856	35.4	0.9150	23.4	0.9512	11.4	
0.8860	35.2	0.9156	23.2	0.9549	11.2	
0.8864	35.0	0.9162	23.0	0.9556	11.0	
0.8868	34.8	0.9168	22.8	0.9563	10.8	
0.8872	34.6	0.9174	22.6	0.9571	10.6	
0.8877	34.4	0.9180	22.4	0.9578	10.4	
0.8881	34.2	0.9185	22.2	0.9586	10.2	
0.8885 0.8889	34·0 33·8	0·9191 0·9197	22·0 21·8	0.9593 0.9601	10·0 9·8	
0.8894	33.6	0.9197	21.6	0.9608	9.6	
0.8898	33.4	0.0209	21.4	0.9616	9.4	
0.8903	33.2	0.9215	21.2	0.9623	9.2	
0.8907	33.0	0.9221	21.0	0.9631	9.0	
0.8911	32.8	0.9227	20.8	0.9639	8.8	
0.8916	32.6	0.9233	20.6	0.9647	8.6	
0.8920	32.4	0.9239	20.1	0.9651	8.4	
0.8925	32.2	0.9245	20·2 20·0	0.9662	8.0	
0·8929 0·8934	32·0 31·8	0.9251 0.9257	19.8	0.9677	7.8	
0.8938	31.6	0.9264	19.6	0.9685	7.6	
0.8943	31.4	0.9271	19.4	0.9693	7.4	
0.8918	31.2	o-9277	19.2	0.9701	7.2	
0.8953	31.0	0.9283	19.0	0.9709	7.0	
0.8957	30.8	0.9289	18.8	0.9717	6.8	
0.8962	30.6	0.9296	18 6	0.9725	6.6	
0.8967	30.4	0.9302	18.4	0.9733	6.4	
0·8971 0 ·8976	30·2 30·0	0.9308 0.9314	18·2 18·0	0.9741	6.0	
0.8981	29.8	0.9321	17.8	0.9757	5.8	
0.8986	29.6	0.9327	17.6	0.9765	5.6	
0.8991	29.1	0.9333	17.4	0.9773	5.4	
0.8996	29.2	0.9340	17.2	0.9781	5.2	
0.9001	29.0	0.9347	17.0	0.9790	5.0	
0.9006	28.8	0.9353	16.8	0.9799	4.8	
0.9011	28.6	0.9360	16.6	0.9807	4.6	
0.9016 0.9021	28·4 28·2	0.9366 0.9373	16·4 16·2	0.9815	4.2	
0.9021	28.0	0.9380	16.0	0.9831	4.0	
0.9031	27.8	0.9386	15.8	0.9839	3.8	
0.9036	27.6	0.9393	15.6	0.9817	3.6	
0.9041	27.4	0.9400	15.4	0.9855	3.4	
0.9047	27.2	0.9407	15.2	0.9863	3.2	
0.9052	27.0	0.9414	15.0	0.9873	3.0	
0.9057	26·8 26·6	0.9420	14·8 14·6	0.9882	2.8	
0.3068 0.3068	26.4	0.9427 0.9434	14.4	0.9890	2.4	
0.9073	26.2	0.9441	14.2	0.9907	2.2	
0.9078	26-0	0.9449	14.0	0.9915	2.0	
0.9083	25.8	0.9456	13.8	0.9924	1.8	
0.9089	25.6	0.9463	13.6	0.9932	16	
0.9094	25.4	0.9470	13.4	0.9941	1.4	
0.9100	25.2	0.9477	13.2	0.9950	1.2	
0.9106	25.0	0.0484	13.0	0.9959	1·0 0·8	
0·9111 0·9116	24.8	0·9491 0·9498	12·8 12·6	0.9967	0.6	
0.9110	24.4	0.9505	12.4	0.9983	0.4	
0.9127	24.2	0.9512	12.2	0.9991	0.2	
		1 7	1			

Racult, C. R. 76, 1261.—3. Ammenia gas combines with a great many metallic chlorides, sulphates, &c., to form either double compounds or compounds which are best regarded as substituted ammonium salts. (For a slight general skotch v. Ammonium Compounds. The several com-

pounds are described in the arts. on the different metals).—4. Ammonia gas or solution combines with acids to form well-marked salts isomorphous with the corresponding salts of the alkali metals. The value of the heat of neutralisation of an acid by NII,Aq is always rather smaller than the value when KOHAq or NaOHAq is used; thus Thomsen (Th. 1, 412-421) gives these numbers:

BAq [H°SO'Aq, BAq] [H°Cl°Aq, BAq]
2KOHAq 31,288 27,501
2NaOHAq 31,5,8 27,488
2NH'3Aq 28,152 24,544
[H°N°O'Aq, BAq]
27,514
27,364

27,644
These results are quite in accordance with the view that an aqueous solution of NH, contains the compound (NH,)OH, analogous in composition and properties to the hydroxides of the alkali metals (v. Almontum Compounds).

Detection. - Free ammonia is detected: 1. by its smell; 2, by its action on HCl whereby white clouds of NH,Cl are produced; 3. by its action on paper, a. moistened with neutral HgNO₃Aq, whereby a black stain (Hg₂O) is formed, b. moistened with CuSO₄Aq whereby a sky-blue colour (CuSO₄4NH₃) is produced, c. moistened with MaSO, Aq, whereby brown spots (Mn.O.) are formed, d. steeped in an ethereal solution of alkanna root (Enz. J. 1870. 935), whereby a blue colour is produced (Böttger, J. pr. 107, 146). The presence of ammonia or ammonium salts can be ascertained by various tests; the following may be mentioned -4. Sodium picrate precipitates yellow ammonium picrate. - 5. A solution of sodium molybdate containing phosphorie and nitrie acids forms a citron yellow pp. (Sonnenschein, J. pr. 56, 302).-6. Nessler's solution-a strongly alkaline solution of HgI₂ in KIAq-forms a brown pp., or brown colour in extremely dilute solution, of NHg.I.H.O (Nessler, C. C. 1856, 529). All ainmonium salts are at least partly volatilised when heated; some give sublimates of the original sait, e.g. NH₂Cl; others are decomposed, e.g. NH₃NO₃ and NH₃NO₂.

Estimation.—1. Ammonium salts are sometimes of install in a minimum salts are sometimes of install in a minimum.

times estimated in a mixture, all the other constituents of which are non-volatile, by heating a specified quantity and determining the loss of weight, -2. All ammonium salts are decomposed by heating with KOHAq (or NaOHAq) with evolution of NH4: if organic N-containing conpounds which evolve NII, by the action of alkalis are absent, it is only necessary to add a weighed quantity of the ammonium compound to an excess of KOUAq, or NaOHAq, which has been boiled and cooled, in a flask connected with a condenser and receiver, and to warm on a sand bath; NH, passes over and is received in dilute HClAq; the NH4Cl is then transformed into 2NH, Cl.PtCl. (v. No. 3), or the NII, is received in a measured quantity—excess—of standardised HClAq, H₂SO₄Aq, or H₂C₂O₄Aq, and the residual acid is determined by titration with standard alkali. [A special apparatus is described by Harcourt (Fr. 2, 14).] If N-containing organic compounds are present which evolve NII, by the action of alkalis, Schlossing (A. Ch. [8] 81, 158) recommends to place the substance with excess of milk of lime over a measured quantity of standard H, SO, Aq, nnder a bell jar, for 48 honrs, and then to determins the residual acid by standard alkali .- 3. Ammonium sales the aoids of which are soluble in alcohol may be estimated by conversion into 2NII Cl.PtCl.; an excess of nearly neutral PtCl, Aq, free from IINO,, is added to the solution, the liquid is evaporated at 100°, the residue is washed with alcohol, dried at 100°, and weighed, or it is strongly heated and the residual platinum is weighed. This method is applicable in the presence of salts which form double compounds with Pt soluble in alcohol; it is best that such ralts should be chlorides, to insure this the mixture is evaporated with excess of conc. HClAq (it is best to remove sulphuric acid by Ba(OH), oxcess of Ba(OH), being afterwards removed by CO.). In the case of KCl, which forms a salt 2KCl.PtCl, insoluble in alcohol, the mixed Pt salts are weighed, theu strongly heated and again weighed, the KCl is dissolved out in water, and the residual Pt is weighed. 4. Ammonium salts are decomposed by alkaline hypochlorites or hypobromites giving off all their N, which may be collected and measured.

(2NH,ClAq + 3NaClOAq = N_2 + 3NaClAq + 3H,O + 2HClAq).

Wöhler employed ealcium hypochlorite for the purpose; Kuop (Fr. 9, 225) used barium or sodium hypobromite (prepared by the action of Br on Ba(OH), Aq or on NaOHAq); Schiff has described a special apparatus (Fr. 7, 430); Krocker a. Dietrich (Fr. 3, 64; 5, 40) decompose by excess of brominated NaOClAq, and determine the residual hypochlorite by titrating with au alkaline solution of arsenious oxide. (Comp. also Foster, C. J. 33, 470.)—5. Minute quantities of NH, are determined by the colorimetric process of NH, are determined by the colorimetric process of Nesslerising; a measured quantity of Nessler's reagent—Hgl, in KIAq made strongly alkaline by KOHAq—is added, and the colour is compared with that produced by an equal quantity of Nessler's solution in an equal volume of water containing a known quautity of ammonia.

M. M. P. M.

AMMONIA, ACTION ON ORGANIC BODIES. 1. Ammonia converts alkyl salts of inorganic acids into amines (q. v.).-2. It converts all yl salts of carboxylic acids into amides (q. v.) .-3. It converts the oxides of acid radieles into amide and ammonium salt: $Ac_2O + 2NH_3 =$ AcNH2 + AcONII, -4. It converts ellorides of acid radicles into amides of the corresponding acids: $AcCl + 2NH_3 = AcNH_2 + NH_4Cl. -5$. It unites with eyanic ethers forming ureas.-6. It unites with thio carbimides forming thiopreas .- 7. It unites with aldehydes, but in the case of the higher fatty aldehydes and the aromatic aldehydes water is simultaneously split off .- 8. It unites with ketones and quinones .-E. Alcoholic NII, sometimes removes HCl; thus, it converts isobutylidene chloride Me, CH. CIICl, chloro-iso butylene Me, C: CIICl, and di-chloropropane CH3.CCl2.CH3, into chloro-propylene CH₃CCl:CH₂(Oeconomides, C. R. 92, 1235).—10. For its action on oxy compounds see AMINES, Formation 10.-11. Ammonia can displace O by NH.

AMMONIAC GUM v. Gum.

AMMONIUM COMPOUNDS. Compounds produced by the action of ammonia on acids. The conditions of occurrence in nature, and also of the artificial syntheses, of these salts are reforred to in the article Ammonia, q. v. In that article soms data ars given regarding the absorption of ammonia by water (Combinations, No. 1), and regarding the thermal values of the neutralisation of acids by NII₃Aq (Combinations, No. 4). The products of the mutual actions of NII, Aq, and HClAq, H, SO, Aq, and other acids-the ammonium salts-arc for ths most part white crystalline bodies, easily solubls in water, and many of them soluble also in alcohol; they exhibit marked analogies with the salts of potassium. Corresponding ammonium and potassium salts are isomorphous, hence they probably have similar compositions. The ammonium salts are distinguished by their comparatively great volatility; heated, as solids, they are completely volatilised, if the acid of the salt is volatile; if the acid is non-volatile (e.g. borate or phosphate), ammonia is evolved. They do not exist as gases; when volatilised they are either decomposed e.g. NH, NO, or dissociated e.g. NII,Cl, q. v. (v. also Dissociation). When gaseous NH, acts on gaseous HCl, HBr, or HI, combination occurs with production of much heat and formation of solid compounds NHaHX, thus (Th. 2, 75):

X (NII³, IIX) Cl 41,900 Br 45,020 I 43,460

If the solid products of these actions, NH3HX, are heated to about 450°, a vapour is obtained containing NH, and HX; on cooling this vapour the compound NH, HX is re-formed. Gaseous NII, does not combine with HC!, HBr, or HI, at temperatures above about 450°. These facts establish a difference between the ammonium and potassium compounds. This difference is further exhibited in the reactions of the two classes of compounds; the ammonium salts are easily decomposed, e.g. by alkalis and alkaline earths, with production of NH3. On the other hand the properties of NH, Aq (v. Ammonia, Properties of) are so similar to those of KOHAq, and the reactions of acids with these solutions, whether considered thermally or chemically, are so analogous, that there can be little doubt that the composition of ammonium salts is similar to that of potassium salts. This similarity is at once rendered apparent by formulating the former class of salts as compounds of the hypothetical group of atoms NII, ammonium. Thus we have:

NH, Cl isomorphous with and chemically analogous to K.Cl,

NH, NO₃ isomorphous with and chemically analogous to K.NO₃,

(NII₁)₂.SO₄ isomorphous with and chemically analogous to K₂.SO₄, (NH₄)₂.C₂O₄ isomorphous with and chemically

analogous to K., C.O., NH4, C., H., O. isomorphous with and chemically

analogous to K.C.H.O.

If this analogy of properties is in all cases supposed to accompany analogy of composition; then NH,Aq must be formulated as NH,OHAq.

The compound NH,OH has not been separated

from the solution of NH. in water: but this does not prove the non-existence of the compound in this solution. A chemical compound may, and sometimes almost certainly does, exist as a member of a system, and yet it may be incapable of existence apart from the other members of the system. The existence of overy compound is conditioned by other factors than the eloments which compose it; among theso factors, temperature, and the presence or absence of other compounds, are very important. Compounds closely resembling NH OH, and undoubtedly derived from NH,OH, are known as definite solid bodies; they are obtained by roplacing the four hydrogen atoms in the complex NH₁OH by alcoholio radicles C₃H_{2n-1} or C_nH_{2n-2}; thus N(CH₂),OH, N(C₃H₃),OH, and N((C₂H₃),(C₆H₃),OH, have been prepared. These bodies closely resemble NaOH and KOH in their properties; their existence and properties aro strong arguments in favour of the existence of the compound NII OH in aqueous selutions of NH2. The formulæ NH2.OH, NH2Cl, (NH2)2SO4 &c., then better summarise the properties and reactions, and suggest the analogies, of the ammonium compounds, than the alternative formulæ NH3.H2O, NH3.HCl, (NH3), H2SO, &c. The name ammonium is given to the compound radicle, or group of atoms, NII. We do not know that the composition of the molecule of ammonium chlorido is represented by the formula NH Cl; indeed we do not oven know the molecular weight of this, or of any other, ammonium compound. These compounds seem to exist only as solids, or in solution. We are scarcely justified in applying the term moleculo to the chemically reacting small particles of solids or liquids, unless the term is used in a wider and vaguer senso than is given to it when we speak of the moleculo of a gas (v. ATOMIC AND MOLECULAR WEIGHTS). But in saying that the complex or collocation of atoms which forms the reacting weight of ammonium chloride is a combination of an atom of chlorino with the radicle, or group of atoms, ammonium, we mean to imply that, when this complex of atoms reacts with various other kinds of matter, it behaves as if the four atoms of hydrogen were, in some way, more directly and closely related to the atom of nitrogen than to the atom of chlorine. The fact that when the same complex of atoms is heated it separates into two distinct molecules, HCl and NH2, neither proves nor disproves the correctness of the formula NIL, Cl, and the conception which that formula embodies. Neither does the fact, that no gaseous molecule is known containing a single atom of nitrogen combined with more than three monovalent atoms, disprove the formula NII, Cl; for the solid compound ammonium oldoride presents us with phenomena to which the conceptions regarding the valencies of atoms, which have been gained by the study of gaseous molecules, are not strictly applicable. In councetion with the constitution of am-

In councetion with the constitution of ammonium compounds it is of interest to observe that the compound produced by the union of N(CH₂),QH₂ with C₂H₃ appears to be identical with the compound produced by the union of N(C₂H₃),QH₃ with CH₂I; it eeems as if this compound N[(CH₂),(C₂H₃)] belonged to the

same form or type as NH, 1, NH, Cl, NH, OH, &c. (V. Meyer and Lecco, B. 8, 233 a. 936).

The group of atoms, NH,, is evidently chemically comparable with the atoms K, Na, Li, Cs, or Rb; but these are the atoms of strongly positive metals; hence if the group NH, could be isolated it might be expected to exhibit properties similar to those of the alkali metals. Experiments have demonstrated the impossibility of the existence of NH, uncombined with other atoms; but certain reactions are known which suggest the existence of an alloy of this hypothetical metallic

radicle with mercury. Ammonium Amalgam. If an electric current is passed through conc. NH3Aq, or NH4ClAq, the negativo electrode consisting of mercury and the positive of a platinum wire, the mercury swellsnp, sometimes to 20 times its original volume, and becomes pasty so that it may be lifted by the hand, while nitrogen is evolved at the positive electrode. The same result is obtained as regards the mercury, if a piece of solid NH_Cl is used; also if sodium-amalgam, containing about 1 p.c. Na, is placed in conc. NH, ClAqin this case NaClAq is produced. The peculiar, pasty, lustrous, metal-like, substance formed in these experiments is called ammonium-amalgain; at a very low temperature, obtained by solid CO, and other, it is a dark-grey, solid, crystalline, mass; even at -29° it bogins to evolve NII, and H, and this change proceeds rapidly at moderate temperatures; the two gases

An amalgam of K with Hg is produced by electolysis under conditions very similar to those which attend the production of ammonium-amalgam; the analogy between ammonium and potassium is thus carried out here also. Ammonium-amalgam, it is said, does not reduce salts of Ag, Cu, or Fo, as K amalgam does (Landolt, A. Suppl. 6, 346). When the amalgam is subjected to increased pressure its volume is found to change almost inversely as the pressure; in this respect then it behaves like a gas rather than a semi-solid compound. The following data are given by Routledgo (C. N. 26, 210):—

always come off in the ratio NII, : H.

e.e. of Hg in amalgam.	c.c. of amalgam, pressure 762 mm.	Increased pressure upplied, in um. of Hg.		e.c. of amalgam calculated by Boyle's law.
14·5	21·0	1 52 4	18·0	17·9
24·4	36·2	1524	31·6	30·9
10·4	18·0	1863	14·7	13.7
23·8	42·2	1026	38·8	38.5
23·8	42·0	2015	32·2	31.6
23·8	36·2	1495	32·6	30·6
29·2	39·5	1989	34·4	35·4

As the pressure increases, the surface of the amalgam becomes brighter, until under large pressures it has the appearance and liquidity of moreury (Seeley, C. N. 21, 265). These results point to the existence of gaseous NH₂ or H, or, it may be, gaseous NH₄, in the Hg; but they do not disprove the existence of the atomic group NH₄ in come kind of loose combination with Hg. An alloy of Fe and NH₄

is said by Meidinger (G. C. 1862. 78) to be formed when FeSO₄Aq or FeGL₄Aq, to which a considerable quantity of NH₂ClAq has been added, is electrolysed by a strong current, the negative electrode consisting of a copper wire.

By electrolysing aqueous solutions of ammonium salts using carbon electrodes, Bartoli a. Papasogli (G. 13,281) obtained benzenecarboxylio acids and a compound of C, II, O, and N, re-

eembling mellogen.

Of the ammonium compounds we have here to consider the bromide, chloride, fluorides, todides, sclenides, telluride, and sulphides; tho others will be considered under the headings CARBONATES, NITRATES, SULPHATES, &C. The hydroxide, known only in aqueous solution, has been already treated of in the art. Armonia.

Ammenium bronide. NH₁Br. Mol. w. unknown; does not exist as a gas, but is dissociated by heat into HBr+NH₃. S.G. 4º 2:379 (Schröder, P. 106, 242). S. G. 4.5° 2:327 (Eder, Sitz. W. 82 (2), 1284). V. D. at 440° to 860' 21-4 (Deville and Troost, C. R. 49, 239; 56, 881). S. (10°) 66·2; (16°) 72; (30°) 81·1; (50°) 94·1; (100°) 128·2. S. (alcohol, S.G. 806, 15°) 3·1; (75°) 10·5. S. (ether S. G. 729)·12 (Eder, l.c.). [NH³, HBr] = 45,020; [N, II⁴, Br] = 65,348 (Th. 2, 75). [NH'Br, Aq] = -4880 (Th. 3, 197). S.V.S. 41·7.

Formation.—1. By adding HBr or HBrAq to NH₁ or NH₃Aq. 2. By the action of Br on NH₃Aq; 4NH₃Aq + 3Br = 3NH₄BrAq + N: on evaporation, crystals are obtained.

Properties and Reactions.—White crystals; soluble in water; the act of solution is attended with absorption of much heat. Exposed to moist air turns yellow, and acquires an acid reaction. An aqueous solution gives off ammonia at moderately low temperatures.

Troost (C. R. 92, 715) describes three compounds, NH, Br, xNH, x = 1, 3, and 6, obtained by the action of excess of NH, on warm NH, Br; the dissociation-phenomena of these compounds have been studied by Roozeboom (lt.T.C. 4, 361).

Ammonium chloride. NH.Cl. (Sal ammoniae.) Mol. w. unknown; does not exist as gas, but is dissociated by heat into NH₁+HCl. S.G. 1-52 (Schröder, P. 106, 212). V.D. at 350°=14·4, at 1040°=14·5 (Deville a. Troost, C. R. 49, 239; 56, 891); but vapour consists of equal volumes of NH, and HCl (v. Reactions, No. 1). S.H. (16° to 45°) 373 (Kopp, T. 155, 71). S.H. (23° to 100°) 3908 (Neumann, P. 126, 123). C.E. (oubical, 0° to 40°) 00018764 (Fizeau, C. R. 64, 814). S.V.S. 35·2. S. (0°) 28·4; (10°) 32·8; (110°) 77·2; S. increases approximately 4·4 parts for each 10° (Alluard, C. R. 59, 500). S. (alcohol S.G. 939, 8°) 12·6; (56°) 30·1 (Gerardin, A. Ch. [41 5, 129]. [NH³, HCl] = 44,900; [NH³Aq, HClAq] = 12,270; [N, H¹, Cl] = 75,790 (Th. 2, 75); [NH⁴Cl,Aq] = -3,880 (Th. 3, 197).

[NH·Cl. n H·O, m H·O] (Th. 3, 109) #+m = 50 25 100 200 10 -87 -121-129-12925 - 84 42 - 42 50 8 92 100 O

Occurrence.—In small quantities, in the neighbourhood of volcances, and in fumaroles; in some animal secretions, e.g. saliva.

Formation.—1. By mixing HCl and NH, in equal volumes. 2. By the action of HCl on N and H, under the influence of the electric discharge, or when heated and quickly cooled (for details v. AMMONIA, Formation, Nos. 1, 2, and 3).—3. By the decay, or destructive distillation, of various N-containing organic matters.

The name Sal ammoniacum seems to have been given by the earlier chemists to rock-salt; Geber, probably latter part of 8th century, prepared ammonium chloride from urino and common salt; towards the end of the seventeenth century the name Sal ammoniacum came to be applied to ammonium chloride. The salt was prepared in Egypt by sublimation from the sooty deposit obtained by burning camel's dung. The first manufactory of sal ammoniac in this country was established at Edinburgh in 1756.

Preparation.—Crude ammonium carbonate obtained by the dry distillation of bones, horn, blood, &c., or gas coal, is decomposed by hot milk of line, and the NH₃ produced is led into HClAq, the liquid is boiled down, and the crude NH₄Cl is purified by solution, filtration through animal charcoal, re-crystallisation, and sublimation. For details of preparation of pure NH₄Cl

v. Ammonia, Preparation No. 3.

Properties.—Pure ammonium chloride is a white, inodorous, salt, with a pungent taste; it crystallises from aqueous solutions in small cubes or octahedra which gather together into feathery masses (v. further, Naumann, J. pr. t0, II and 310). By sublimation and rapid cooling it is obtained as a loosely cohering powder consisting of minute octahedra; the ordinary sublimed salt has been partially fused and appears as a semitranslucent mass of fibrous crystals. It is very tough, and cannot be powdered; sal ammoniae is best obtained in fine powder by evaporating a solution to dryness with constant stirring. It is casily soluble in water (v. data at beginning of article); a conc. aqueous solution boils at 115.80 at 768 mm. (Alluard, C. R. 59, 500); S.G. 120 of cone. NH₄ClAq containing 26 p.c. NH₄Cl is 1.0752 (Michel a. Krafft, A. Ch. [3] 41, 471). Tables of S.G. of NH ClAq are given by Gerlach (J. 1859. 42), and Schiff (A. 110, 74). On heating NH, ClAq of 106 p.c. to 37° NII, is given off in tho water-vapour (Lceds, Am. S. [3] 7, 197); as temperature increases the decomposition of NH, Cl probably increases also (v. Fittig, A. 128, 189; Dibbits, B. 5, 820; Brücke, J. pr. 101, 481).
Reactions.—1. When NH,Cl is heated it

Reactions.—1. When NH Cl is heated it vapour is found to consist of HCl and NH (v. Pehal, A. 123, 199; Than, A. 131, 129; Wanklyn, P.M. [4], 29, 112; Wurtz, J. 1859, 30; Deville a. Troost, C.R. 49, 239; 56, 891; Tommasi, B. 14, 353). The density of the vapour is 12-9 according to Bineau (A. Ch. [2] 68, 416); 144 at 350°, and the same at 1010°, according to Deville and Troost (C.R. 49, 239; 56, 891). Than (l.c.) has shown that HCl does not combine with NH, at 350° or higher temperatures.—2. NH ClAq is decomposed by chlorine with formation of HCl and nitrogen chloride (q. v.).—3. Heated with iron, zine, or better with potassium, NH Cl is decomposed with production of metallic chloride, NH,, and H; in presence of air and moisture the reaction proceeds rapidly with formation of metallic chloride.

or oxychloride and NH .- 4. Many metallic | oxides decompose NH Cl with formation of chlorides, and NH2; in some cases-e.g. oxides of Hg, Pt, Au &o.—the chloride combines with a portion of the NH₄Cl to form a deuble compound. b. Alkaline carbonates decompose NII, CI when heated with volatilisation of ammonium car-Calcium carbonate, especially when freshly precipitated, dissolves readily in NII, ClAq; on heating, ammonium carbonato is evolved .-6. Crystallised sodium sulphate partially decomposes NH Cl when the two are mixed togethor by rubbing, NaCl and (NH₄)₂SO₄ being formed and partly dissolving in the water which comes from the sodium sulphate crystals .-7. Sulphuric anhydride vapour is absorbed by powdered NII,Cl, on warming HCl is evolved and SO,NH,ONH, is formed; if water is added (NH₄)₂SO₄ is produced.—8. The reaction between acids and NH,Cl follows the ordinary course of the interaction of acids with salts of other acids.

Combinations. - I. NH4Cl combines with many metallic chlorides to form double compounds; e.g. PtCl, 2NH, Cl; HgCl, 2NH, Cl; CuCl, 2NH, Cl &e. (v. the several Metals). -2. With an aqueous solution of ICl,, the compound NH₄Cl.ICl₃ is produced (v. Ammonium iodide; Reactions, No. 2) .- 3. According to Troost (C. R. 88, 578) when NH, Cl is heated with a large excess of pure dry ammonia, at least two compounds are formed: HCl.4NII, melting at 7°; and HCl.7NH, formed at -31° and 750 mm., melting at - 18°.

Ammonium flueridss (Marignac, Ann. M. [5]

15, 221).

I. NEUTRAL SALT. NH.F. Mol. w. unknown. [NH*Aq, HFAq] = 15,200 [NH*, HF] = 30,100 (Gunta, C. R. 97, 1483). Formation.—1. By mixing HF and NH₂.—2. By heating KF or NaF with NH4Cl. Preparation .- 1. By gently heating a dry finely powdered mixture of I part NH,Cl with 31 parts KF in a platinum crucible covered with a lid, which is kept cold by dropping water on to it; the NHL sublimes on to the lid.—2. Ordinary HPAq is saturated with NH₃Aq, a little (NH₄)₂CO₃Aq is added, the clear liquid is decanted and evaporated in a platinom dish with repeated additions of small pieces of solid ammonium carbonate. Properties.—Hexagenal prisms with strong saline taste; unchanged in dry air, but deliquescent in moist air; oasily soluble in water, less soluble in alcohol; an aqueous solntion gives off NH, and acquires an acid reaction; the dry salt absorbs NHa but gives it off again on heating; sublimes readily with previous fusion; etches glass, and must be kept in platinum, silver, or gutta percha, vessels. Reactions ... I. When moist, or in solution in water, NH,F decomposes silicates with formation of SiF4; the same decomposition is effected by the dry salt by heating it with silicates.—2. An aqueous solution is decomposed by heat, with formation of the acid salt NH, F.HF and evolution of NH, (v. infra).

NH.F.HF. Mol. w. nn-II. ACID SALT. known. S.G. 1.2I. Formation. - I. By evaporating an aqueous solution of NH,F at 36° to 40° in a platinum dish .-- 2. By adding excess of NHAq to a solution of fluosilioie acid, and evaporating: as thus obtained the salt is mixed with silica. Properties.—Colondess prisms:

easily soluble in water; slightly delignescent; easily volatilised, vapour being very acrid.

Ammonium iodides.

I. NH,I. Mol. w. nnknown; does not exist as gas, but is dissociated by heat into NH, + HI. S.G. 11 ° 2 498. S.V.S. 58. V.D. (440° to 860°) 38.8, but vapour consists of equal volumes of III and NH₃. [NH³, HI]=43,462; [N,H⁴,I]=49,313 (Th. 2,75). [NH⁴I,Aq] = -3,550 (Th. 3, 197). Formation.-I. By mixing equal volumes of III and NH3, or by acting on HIAq with NH, Aq. — 2. By decomposing Fe.I. Aq by (NH) CO.Aq. or BaJ.Aq by (NH,), SO.Aq. Pre-(NH₄), CO₃Aq, or BaJ₂Aq by (NH₄), SO₄Aq. paration .- I. Hot saturated solutions of (NH4)2SO4 and KI, equal equivalents, are mixed; after cooling, alcohol equal to 15 p.c. of the water used is added; the liquid is filtered after I2 hours, and evaporated with addition of a little NH₃Aq from time to time (Jacobseu, C. C. 1864. 192).—2. A solution of 27½ parts KI in 48 parts H.O is mixed with a solution of 22 parts tartaric acid in 48 parts water, the mixture is placed in a freezing mixture to separato KHC, II,O, the filtrate is evaporated at 100° with addition of a little (NH₄)₂CO₃ (Beyer, D. P. J. 171, 467). Properties.—Colourless cakes, very soluble in water and alcohol; deliquesces in moist air, and becomes yellow through separation of I, and loss of NII,; may be sublimed unchanged in absence of air. Reactions .- I. NH, IAq is casily decomposed by dilute acids; the solid compound is decomposed by dry HCl gas at high temperatures. at 360° about 10th, at 440° about 1th, at a dark red heat about this, of the NII,I being decomposed (Hautefeuille, C. R. 64, 704).—2. Chlorine led into saturated NII, IAq produces long, goldenyellow, needles of NH Cl.ICl, (Filhol, J. Ph. 25, 441); this compound is decomposed by gentle heating into ICl, and NH Cl.—3. By heating in ammonia, the compounds NH, LxNII, v = 1, 3, and 6, are obtained according to Troost (C. R. 92, 715).

II. According to Guthric (C. J. [2] I, 239) a compound of NII, I with I-NII, 1.1-is obtained, as a brownish-black liquid, soluble in alcohol, ether, CS2, and KIAq, less soluble in CHCl2, when I is added in small quantities to a cone, solution of NH, NO, with which 1 equivalent of KOH has been mixed. NH,I.I casily separates in dry air into NII, and I; water or dilute alkali produces iodide of nitrogen, NII, I, and HI; mercury acts on it to form HgI, and NH,; aqueous solutions of acids produce NII, salts and separate I.

Johnson (C. J. 33, 397) describes a compound of NH, I and I the composition of which agrees with the formula NH,I3. It is produced by adding I to NII, I in presence of a little water until no more I dissolves. It forms dark-blue, somewhat deligy scent, prisms; S.G. 3.749; soluble in a little water, decomposed on dilution with precipitation of I. This compound seems to form a double salt with KI, viz. 5NH, Iz. KI, obtained by passing NH_a into the mother liquor from which KI, has separated.

Ammenium selenidss (NII,)2Sc, and (NH,)SeH (Bineau, A. Ch. [2] 67, 229). Neither has been gasified and therefore mol. ws. are unknown. NH, has no action on Se, but readily combines with H2Se; when excess of NH, is used 2 vols. combine with 1 vol. H2Se and produce (NH4)2Se. when excess of H.Se is used equal vols. of the

gases combine and form NH, HSe. These compounds are white solids which soon turn red by exposure to air or when dissolved in air-containing water; both smell of NH, and H, So and appear easily to undergo change; their aqueous solu-tions probably contain polyselenides although none of these has been isolated; the products of the distillation of K.Sc with NH,Cl probably

also contain anmonium polyselenides.

Ammonium telluride NH, HTe. White leaf-shaped crystals; easily soluble in water, volatilised at 80°. Formed by the direct union of NH, and H.Tc (Bineau, A. Ch. [2] 67, 229).

Ammonium sulphides, and Sulphydrate or Hydrosulphido. Five solid sulphides, and a hydrosulphide, of ammonium are known; their compositions are expressed by the formula NH,HS, (NII,) 2S, (NII,) 2S, (NII,) 2S, (NII,) 2S, (NH,) 2S, (NH,) 2S, in the gaseous state; the first and second, which have been more studied than the others, are dissociated by heat, into NH3+H2S, and 2NH3+H2S, respectively. All the ammonium sulphides are soluble in water, they very easily undergo change at ordinary temperatures, usually giving off NII, and H.S which is often partly decomposed with precipitation of They are all decomposed by dilute acids with precipitation of white amorphous S, evolution of H.S. and formation of an ammonium salt of the reacting acid. These sulphides act as salt-forming or basic compounds towards such acidic sulphides as As S, As S, Sb S, &c. (v. infra: also Arsenic, and Antimony, thio-acids). According to Berzelius any one of the ammonium sulphides, except (NH4)2 S7, can be prepared by gently heating the corresponding sulphide of potassium with sal ammoniae; in every case except that of K2S, the NII Cl must be in execss, else part of the ammonium sulphide formed is decomposed with production of S (NH₃ and II) which combines with the potassium sutphide to form K.S. Little or nothing is known of the physical coustants of these compounds; the following thermal data are given, but, by reason of the instability of the sulphides and the indirect methods by which the numbers have been obtained, they must be accepted with caution:-

H.F. of solids from gascous N and H, and solid S. $(NH_1)_2 S_2 = 69,000$ $(NH_4)_2 S_4 = 69,600$ (Sabatier, C. R. 91, 53) $(NH_4)_2 S_5 = 69,400$ (Sabatier, A. Ch. [5] 22, 73).

The tetra- and penta-sulphide dissolve in water with absorption of about 8,000 gram-units of heat per formula-weight of the sulphide.

The sulphides of ammonium have been studied chiefly by Fritzscho (J. pr. 24, 460;

Preparation.—NH,HS is prepared by the reaction of equal volumes of NH, and H,S at the ordinary temperature, or at temperatures not lower than - 10°. An aqueous solution of NH,IIS is obtained by saturating NH,Aq with H,S in absence of air.

(NH₁).S is prepared by cooling a mixture of 2 vols. NH₂ and 1 vol. H.S to -18°; or by distilling K.S with excess of NH Cl and cooling tho distillate to -18°.

(NII,) S2 is obtained by passing vapour of S and of NH Cl through a hot porcelain tube and

then into a well cooled reselver. An aqueous solution may be prepared by dissolving S in

(NH₁),S Aq in the proportion (NH₁),S.S. (NH₁),S₀: when NH₁HSAq (v. supra) is digested with S, the solution saturated with NH₁, and then with H2S, more S added, and saturation with NH, and then with H,S repeated, the whole liquid sets to a crystallino mass; if this is heated to 40°-50° a clear liquid is produced from which, on gradual oooling, largo orystals of (NH4), S. separate out.

(NII4)2S4: if the mother liquor from the orystals of (NH₄)₂S₅ is surrounded by a freezing mixture, and treated first with NH, and thon with H.S a orystalline magma is formed; on warming a clear liquid is produced from which

crystals of (NII₁)₂S₄ are deposited.

(NH,) S, is obtained by the gradual decomposition of (NH,) S in dry, slightly warm, air; it is also formed when a solution of (NH1)2Sa in its mother liquor (v. supra) is placed under a large bell jar for some time.

Properties and Reactions.—NH,SH: hard, white, plates or needles; very soluble in water, and easily volatilised. V.D. at 56° 12.8, which corresponds with equal vols. of ILS and NH. (Deville a. Troost, C. R. 56, 891). Aqueous solution is colourless, but soon changes in air from absorption of O, which decomposes a part of the NH, HS with formation of H.O, NII₃, and S; some of the S acts on the remaining NH, HS to form (NII,)2S2, another part of the S is oxidised to $\Pi_{a}S_{a}O_{a}$, and a portion of it is usually deposited. This process proceeds if exposure to air is prolonged; the (NII₁),S₂ is slowly decomposed, until finally a solution of (NII₁)₂S₂O₃ in NII,Aq, mixed with solid S, is the result. NII,IISAq reacts with most metallic salts in solution to form sulphides of the metals; it also reacts with acidic metallic sulphides to form ammonium thio-salts, with evolution of II.S. e.g. $As_2S_3 + 2NH_4SIIAq = 2NH_4AsS_2Aq + II_2S$ (v. Ausenic, Thio-Acids of).

(NH.) S: white, lustrous crystals; easily soluble in water, forming a colourless liquid which easily decomposes with evolution of NH₃ and formation of NH₄IIS. V.D. 18.2 (calculated for 2 vols. NH₂ + 1 vol. H₂S = 17.0) (Deville a. Troost, C. R. 56, 891). Reacts as a strongly marked base towards acidic sulphides to form ammonium thio-salts.

(NII,)2S4: sulphur-yellow erystals, easily soluble in water and alcohol. Stable only in au atmosphere saturated with NH, and H2S; easily decomposed in air with evolution of NH, and H.S. A saturated aqueous solution is fairly stablo; more dilute solutions, and solutions in alcohol, soon precipitate S. When heated, NH HS and S are formed.

(NH₁)₂S₅: orango-rod prismatio orystals, casily soluble in water and alcohol. These solutions are very unstable, decomposing into (NH_d)₂S₂, S, H₂S, and NH₃, and after some time also (NH_d)₂S₂O₃. In dry air the crystals give off NH, and NH, HS and are changed into (NH,),S,.

(NH4)2S7: ruby-red crystals, much more stable than any of the lower sulphidss; decomposed avabout 300°; soluble in water forming a fairly stable liquid which is only slowly decompossd by HClAq.

The liquid known as fuming liquor of Boyle,

AMYI.

or volatile liver of sulphur, chiefly consists of a mixture of various ammonium polysulphides; obtained by distilling a mixture of 1 part S, 2 parts NH,Cl, and 3 parts CaO. It is a dark yellow, strongly smelling, and etrongly furning, liquid. It dissolves sulphur and then no longer fumes in air.

Ammonia reacts with many metallic salts to form compounde, soveral of which behave as if they were derivatives of ammonium chloride, sulphate, &c., rather than double compounds of ammonia with the metallic salts in question. Thus NH3 and PtCl2 form the crystallino compound PtCl., 4NII, by the action of H.SO, on this, HCl is evolved, and there is produced PtSO₄4NH₃; decomposed by Ba(OH)₂Aq this compound yielde Pt(OH)₂4NH₃, which loses H₂O when heated, with production of PtO.4NH3. The compound Pt(OH)2.4NH3 is a markedly alkaline body, rosembling NaOH or KOH; it neutralises 2 equivalents of a monobasic acid. The compounds PtCl. 4NH₃, PtSO₄4NH₃, and Pt(OII)₂4NH₃, can ecarcely be regarded as double compounds of ammonia and platinum salts; their reactions are better suggested by supposing them to be derivativee of animonium compounde, obtained by replacing part of the hydrogen by platinnm. The name platinam monium has been given to the hypothetical radicle

(Pt<NH3) N2HPt.

The chloride of this radicle would be N.H.PtCl,; if two hydrogen atoms in the group N₂H₆Pt are supposed to be replaced by two ammonium groups (NH,) we get the hypothetical radicle a mmonium-platinammonium N.H.(NH.).Pt. The componeds PtCl_4NH₃, PtSO_4NH₃, and Pt(OH)₂.4NH₃ may be regarded as compounds of this radicle; thus N_H_(NH₁)_Pt.Cl₂, N_2H₄(NH₁)_2Pt.SO₄, N_2H₄(NH₁)_Pt.Cl₂, the string of

Compounds are obtained by the action of NH3Aq on Hg.Cl., and HgCl., respectively, which have the composition Hg.NH.Cl and HgNH.Cl; these react as derivatives of NILCl in which II, is replaced by Hg, and by Hg, respectively. The name merouro-ammonium is sometimee given to the hypothetical radicle NIL IIg., and the name mereuri-ammonium to the hypothetical radicle NII, Hg.

A great many bodies are known the reactions and relations of which can be gathered together into one point of view by considering them as compounds of various hypothetical radicles derived from NII, N₂H₈, N₂H₁₂, &c., by replace-ment of part of the hydrogen by various metals. Theee compounds will be described under the headings of the various metals (v. more particularly the chrom-ammonium-, cobalit-ammonium-, COPPER-AMMONIUM-, MERCURY-AMMONIUM-, RIId PLA-TINUM-AMMONIUM-, COMPOUNDS; in the articles Chromium, Cobalt, Copper, Mercury, and Pla-TINUM respectively).

Ammonium salts, i.e. derivatives of acide obtained by replacing H by the group NH, are described under the various headings CARBONATES, NITRATES, SULPHATES, &c. &c. The principal salts are the following:—Antimonate; arsenate, arsenite; borate; bromate, &c.; carbamate; carbonates; chlorates, chlorite, tc.; chromates; cyanates; cyanide; iodate, per-

iodate, do.; molybdates; nitrate, nitrite, &c.; phosphates; selenite, &c.; silicates; sulphamate, dc. : sulphates, sulphites, dc. : tantalate : tellurates; thioarsenates, thiocyanates, thiosulphates, &c. For an account of the general properties of these salts v. beginning of precent article. M. M. P. M.

AMOXY .. Contraction for amyl derivative of Oxy-

AMYDECYLENIC ACID v. Decenoid acid. AMYGDALIC ACID C20II26O12 or C20H28O12 Formed by boiling amygdalin with baryta. Deliquescent crystallino mass, insol. alcohol, and ether. By boiling with H.SO, and MnO, it yields formic acid, CO2, and benzoic aldohyde (Liebig a. Wöhler, A.22, 11; 66, 240; Schiff, A. 154, 348).

Acetyl derivatives C20H2, Ao, O18 and

 $C_{20}II_{21}\Lambda c_{*}O_{13}$ (S.). AMYGDALIN $C_{20}II_{27}NO_{11}3aq$. [200°]; after solidifying it melts at 125°-130° (Wöhler, A. 4I, 155). Mol. w. 511. S. 8.5 at 12°. [a] = 35.5°.

Occurrence. - In bitter almonds (Robiquet a. Boutron, A. Ch. [2] 44, 352); to a small extent in sweet almonds; in laurel leaves (Cerasus lauro-cerasus); in the leaves, blossoms, and bark of the birdcherry (Prunus padus); in young choots of the apple tree; and in the kernels of apples, pears, and peaches (Riegel, A. 48, 361; Wicke, A. 79, 79; 81, 241; Lehmann, N. R. P. 23, 449).

Preparation.—The almond-cake from which the fatty oil of almonde has been removed by pressure is extracted with boiling alcohol; the filtrate is concentrated and the amygdalin ppd.

Properties .- White pearly scales or thin prisms (from water). Insol. ether.

Reactions.-1. Under the influence of emulsin or of boiling dilute H.SO, it ie split up into beuzoic aldehyde, prussio acid, and glucoso (Liebig a. Wöhler, A. 22, 17): $C_{20}H_{27}NO_{11} + 2II_{2}O = C_{7}H_{6}O + CNH + 2C_{6}H_{12}O_{6}$ 2. KMnO, forms cyanic and benzoic acids.-3. Potash or baryta form amygdalic acid.-4. Conc. HCl gives mandelio acid, glucose, aud NII₃. - 5. PCI₅ gives CyCl and benzylidene chloride. -6. Zn and diluto hydrochlorie acid

chloride. -6. Zn anu unito give C.H., CH., CH., CH., NH., (Fileti, B. 12, 297). Acetyl derivative C₂₀H₁₀Ae,NO₁₁. Long needles (from alcohol); insol. water (Schiff; A. I54, 338).

Amorphous amygdalin has been described by Winckler (B. J. 20, 428), Neumann (B. J. 23, 503), Simon (A. 31, 263), and Lehmann (loc. cit.).

AMYL C.H., I'entyl. A monovalont baeylous radiclo which can occur in eight forms: n-amyl, CH₃.CH₂.CH₂.CH₃.CH₄.CH₂; iso-butyl-carbinyl (CH.) CH.CH...CH. tert-butyl-carbinyl (Cll_)CH(C.H_).CH_; tert-butyl-carbinyl (Cll_)_3C.CH_; methyl-n-propyl-carbinyl,CH_,CH_,CH_,C(CH_)H; methyl-isopropyl-carbinyl, (CH1) CH.C(CH2)H; di-ethyl-carbinyl (C2H5).CH; and di-methyl-ethyl-oarbinyl $(CH_3)_2(C_2H_3)C_4$ Ordinary amyl alcohol is a mixture of iso-butylcarbinol and sec-butyl carbinol, and it is from this mixture that most of the amyl compounds

have been prepared. The term 'iso-amyl' com-pounds will, for the sake of brevity, bs used in

this dictionary to denote the mixture of amy!

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compounds prepared from this source. Inasmuch as the proportion of the two conetituents of 'isoamyl' alcohol varies with ite ecurce, 'isoamyl' compounds prepared by different chemiste can hardly be expected to possess identical physical charactere. Amyl derivatives of hydroxylio carbon compounds are described under the compounds of which they are the ethers.

Di-amyl (C,II,)2 or C, the v. Decane.

AMYL-ACETYLENE v. Heptinene.

AMYL ALDEHYDE v. VALERIC ALDEHYDE.

AMYL ACETATES C, H, O, Pentyt acctates. Mol. w. 130.

Preparation. - Similar to that of ethyl

acetate, p. 14.

n-Amyl acetate, (147·6°) (Gartenmeister, A. 233, 260); (148·4°) at 737 mm. (Lieben a. Rossi, A. 159, 74). S.G. § 8948; § 20 8774 (G.). C.E. (0°-10°) 00106 (G.). S.V. 173·8 (G.). Prepared from n-amyl iodide and silver acctate.

Iso-amyl acetate. (137.6°) at 745 min.; (188.9°) (R. Schiff, A. 220, 110). S.G. 4.8762 (Mendeléeff, J. 1860, 7); $\frac{26}{4}$ ·8561 (Brühl). $\mu_{\mathcal{B}}$ 1·4088 (B.). $R\infty$ 59·7 (B.) S.V. 174·6 (S.). Is largely used as a flavouring agent to imitate jargonelle pears.

Methyl-propyl-carbinyl acetate. (133°...135°) (Wurtz, A. 148, 132); (134°-137°) (Schorlemmer, A. 161, 269). S.G. 2 922 (W.).

Methyl-isopropyl-carbinyl acetate. (125°)

(Wurtz, A. 129, 367). Di-ethyl-carbinyl acetate. (132°) at 741 nm. S.G. 2 909 (Wagner a. Saytzeff, A. 175, 306).

Tert-amyl acetate. (124°) at 750 ium. S.G. º 891 (Flawitzky, A. 179, 318). Decomposed by heat into amylene and acetic acid (Menschutkin, C. R. 95, 618).

AMYL ALCOHOLS C, IÍ, O. Mol. w. 88. Theory indicates 8 amyl alcohols (v. Amyr.), viz.: 4 primary, 3 secondary, and 1 tertiary. One of these, tert-butyl carbinol, is unknown.

n-Amyl alcohol Cl13.CH2.Cl12.CH2.CII...O11. (137°) at 740 mm. (Lieben a. Rossi, C. R. 71, 370); (137·9° i. V.) (Zander, A. 224, 81). S.G. 6 8282 (Z.). C.E. (0°-10°) ·00091 (Z.) S.V. 123.4 (Z.).

Occurrence.-In fusel oil (Wyschnegradsky, A. 190, 350).

Formation.-1. From n-valeric aldehyde and sodium amalgam (L. a. R.).-2. From n-amyl

chloride (Schorlemmer, A. 161, 268). Inactive amyl alcohol

(CH_a)₂CH.CH₂CH₂OH. (130·5°-131·2°) (Lachowiez, A. 220, 171); (131-5° cor.) (Perkin). S.G. 15° 8135; 25° 8078 (P.). M.M. 5-959 at 18-6° (P.). Fusel oil is a mixture of active and inactive amyl alcohol; they can he more or less eeparated either by passing HCl into the boiling alcohol, when the inactive alcohol is converted into amyl chloride more readily than its isomeride (Lc Bcl, C. R. 77, 1021); or by means of the barium salts of the two amyl-sulphurio acids, C,11,1SO,II, the active salt being the more coluble in water (Pasteur, C. R. 41, 296). The simplest way to obtain an inactive amyl alcohol is carefully to fractionate fusel oil (L.). The same alcohol can be prepared from isobutyl alcohol by converting it first into valeric soid (Balbiano, G. 6, 229). Iso-amyl alcohol.

(CH₃)OH(O₂H₃).OH₄OH. Fermentation amy! alcohol. Fusel oil. [0.-134] (Olszewski, M. 5, 128). (130·5°-131°) (R. Sohiff, A. 220, 102). S.G. 29 ·8104 (Brühl, A. 203, 23). S.H. 102). S.G. 2 8104 (Bruhl, A. 208, 23). S.H. 679 (Diaconoff, Bl. [2] 38, 172). Latent heat of vaporisation 123·8 (D.). S. 2·5 at 16·5. H.F.p. 74,890 (Thomsen). H.F.v. 71,700 (T.). μ_{β} 1·4124. R._{\omega} 43·08. S.V. 122·7 (S.). Critical temperature 307° (Pawlewski, B. 16, 2634).

Occurrence.-Formed in small quantity in the alcoholio fermentation of saccharine liquids. Isoamyl angelato and isoamyl tiglato occur in Roman oil of chamomile (Köbig, A. 195 99).

Properties.—Poisonous liquid with powerful odour. Its detection in alcohol is described on

p. 96. It burns with smoky flame.

Decomposition .- 1. Its vapour led through a red-hot tube produces acctylene, cthylene, propylene, and butylene (Wurtz, A. 101, 212).-2. S.Cl. gives amyl chloride and amyl sulphite (Carius a. Frics, A. 109, 1).-3. PCl, and PCl, form amyl chloride.-4. Potash-lime at 220° gives hydrogen and potassic valerate. - 5. ZnCl, produces amylone (q. v.). 11ot H.SO, and P.O. act similarly .- 6. Poured upon bleaching powder, it reacts in less than an hour; the distillate decomposes with evolution of Cl., and HCl, and then contains amyl alcohol, valeric aldehyde, and amyl valerate (Goldberg, J. pr. [2] 24, 116).

Compounds.—(C₃H₁₂O)₂SnCl₄. Deliquescent erystalline plates, decomposed by water (Bauer a. Klein, A. 147, 249).—C₂II₁₂O₂SbCl₂.—Crystalline.-- $(C_s\Pi_{12}O)_3$ CaCl₂ (Heindl, M. 2, 209).

Sodic amytate, C,II, NaO,2C,II, O (Fröhlich, A. 202, 295). At 165° it combines with CO forming sodic isovalerate and the sodium salt of an acid $C_{10}\Pi_{18}O_2$. CO passed over a mixture of NaOC₂H₁₁ and NaOAo at 180° produces sodic formate and the sodium salts of a variety of acids, the principal being iso-heptoic acid (q.v.) formed by substitution of H of acctic acid by C₅Π₁₁. Another product is oxy-vinyl-heptoio or oxy-ennenoio acid (q. v.). An acid orystallising in needles C₁₁H₁₂O₄ [139°] is also formed; its empirical formula is that of di-acetyl-heptoio acid (Poetsch, A. 218, 56).

Potassium amylate C, II, OK. White silky crystals (do Forcrand, C. R. 104, 68).

Thaltium amytate C, H, OTl. S.G. 25. An oil obtained by heating thallium ethylate with amyl alcohol.

Aluminium amylate Al(OC, H11)3. [70°]. S.G. 4 9801. Formed by action of AlI, and iodine (Gladstone a. Tribe, C. J. 39, 7; v. ALUMINIUM IODIDE, p. 148).

Active amyl alcohol CEtMeH.CH,OH. (128°) (Pedler, A. 147, 243); (127°-128°) (Just, A. 220, 149). $a_{\rm p} = -2.3^{\circ}$ (J.); -4.4° (Lo Bel; Pierre a. Puchot).

Occurrence.-In fermontation amyl alcohol, which is thus condered more or less lævorotatory Preparation. - Described under inactive amyl

alcohol.

Properties .- In a sample for which a was only -1.14°. Perkin (C. J. 45, 470) found S.G. 18.8150; 25.8091; and M.M. 5.94 at 20°. A rotation of more than 4.4° to the left (in a tube 100 mm. long) has been observed by Loy (~11.5°), and by Pedler (-8.6°

Reactions .--1. A dilute solution mixed with mixture of (CH₂)₂CH.CH₂CH.OH and yeast, penicillium glaucum, and a little H.SO. becomes dextrorotatory. The new dextro-rotatory amyl alcohol forms a laworotatory iodide (Le Bel, Bl. 81, 104).—2. On oxidation it yields a dextrorotatory valeric acid, boiling at 170° (Pedler).—3. Hot NaOH renders it inactivs.

References. - Pastour, C. R. 41, 296; A. 96, 255; Popoff, B. 6, 560; Lsy, B. 6, 1362; Erlenmeyer a. Holl, A. 160, 257; Pierre a. Puoliot, C. R. 76, 1332; Bakhoven, J. pr. [2] 8, 272; Le Bol, B. 6, 70; 9, 358, 732; C. R. 82, 562; Bl. 25, 545; Pedler, A. 147, 243; Chapman a. Smith, Pr. 17, 308.

Msthyl-n-propyl-carbinel Pr.CMcH.OH

(119°). S.G. 2 ·824. S. 13·7.

Formation.-1. From its iodide.-2. By reducing methyl propyl kctons with sodiumamalgam (Belohoubck, B. 9, 924). So prepared it is inactive, but if it be dissolved in 20 pts. water and penicillium glaucum bo introduced, it becomes lavorotatory (a -5.5°) (LoBel, C. R. 89, 312).—3. From acetyl chloride and zino propyl (Markownikoff, Bl. [2] 41, 259).

Reactions. - 1. Oxidation gives methyl propyl ketone .- 2. Gives the iodoform reaction.

Pr.CMeH.OH. Msthyl-isepropyl-carbinel. (113°). S.G. 2 ·833 (Wischnegradsky, A. 190, 338). Formation.-1. From methyl isopropyl ketone with sodium-amalgam (Münoli, A. 180, 839) .- 2. By adding water to the product of the

action of zinc methido on bromo-acetyl bromido (Winogradoff, A. 191, 125), or chloro-acetyl chloride (Bogomoletz, A. 209, 86; Bl. [2] 34, 330).

Reactions .- 1. Conc. H.SO, forms tri-methylethylene, Mc.C:CMcH, which may be converted by warm cone. HI into the iodide of tertiary amyl alcohol .- 2. Oxidation gives methyl isopropyl ketono, acetone, acetic acid, and CO2. 3. PCl_s forms a chloride (87°).

Di-ethyl carbinol Et, CH.OH. (117°). S.G.º 832. Formed by adding water to the product of the action of zino ethide on ethyl formate (Wagner a. Saytzeff, A. 175, 351). The first rsaction may be written HCO.OEt + 2ZnEt = HCEt(OZnEt)Et + ZnEt(OEt). Water then displaces OZnEt by OH. It gives di-cthyl ketone on oxidation.

Tsrtiary amyl alcebel Et.CMe2.OII. Di-methyl-ethyl-carbinol. Amylene hydrate. [-12°]. (102° cor.) (Perkin, C. J. 45, 471). S.G. $\frac{1}{15}$ *8144; $\frac{25}{25}$ *8070 (P.). M.M. 5°99 at 19°. H.F.p. 84,510. H.F.v. 81,320 (Th.). S.V. 121.3 (R. Schiff, A. 220, 102).

Formation .- 1. From tertiary amyl iodide (q. v.) .- 2. From zine methido and propionyl chloride (Popoff, A. 145, 292; Jermolajeff, Z. 1871, 275; Wysolinegradsky, A. 190, 336).

Preparation. — Amylene (1 vol.), prepared from ordinary amyl alcohol, is shaken with an ice-cold mixture of water (1 vol.) and H2SO, (1 vol.).

Reactions .- 1. On oxidation it gives riso to acetone and acetio acid.-2. When introduced into the stomach (of a rabbit) it is excreted as a glyenronate, $C_{11}H_{20}^{*}O_{7}$, which is split up by acids into the alcohol and glyouronic acid (Thierfelder a. Mering, II. 9, 515).—3. Slowly decomposed by heat at 220° into II.O and amylene, this decomposition does not take place unless traces of HOl or HI are present (Wolkoff a. Bougaieff, J. R. 1885, 276).

AMYLAMINES C.H. N. Normal Amylamine

CH2.CH2.CH2.CH2.CH2.NH2 (108°).

Formation .- From the amide of normal hexoic acid by the action of bromine and potash (Hofmann, B. 15, 770). A mixture of 1 mel. proportion amide and 1 mel. bromine is run into excess of 10 p.c. solution of potash at 60°.

AMYLAMINES FROM AMYL ALCOHOL OF FER-MENTATION :-

Ordinary Amylamine

(Cll₃)₂.CH.CH₂.ClP₂.NH₂ (95°-96°). S.G. 13 · 7503.

S.V. 126.84 (Schiff).

Formation .- 1. Amyl cyanate or oyanurate with potash (Wurtz, A. Ch. [3] 30, 447; Brazisr and Gossleth, A. 75, 252).—2. Dry distillation of animal substances (Anderson, A. 105, 335) .-3. Dry distillation of leucino (Schwanert, A. 102, 225). -4. Amylsulphate of potassium with alcoholic ammonia at 250° (Berthelot, A. 87, 372).-5. Distillation of horn with aqueous potash (Limpricht, A. 101, 296).—6. Caustio potash on ilannel (Gr. Williams, Chem. Gaz., 1858, 310) .-7. Amide of isohexoic acid (isobutylacetamids) with bromine and aqueous potash (Hofmann, B. 15, 770) .- 8. In the decomposition of yeast (Müller, J. 1857, 403).

Preparation.—Amyl bromido is heated to 100° with alcoholic ammonia in large excess, the alcohol evaporated and the residual hydrobromides decomposed with potash. The oily layer which consists of mono-, di-, and some tri-, amylamine, is dried with caustic baryta and fractionated. Or, potassium amyl sulphato is distilled with potassium cyanate; the resulting amyl cyanate and cyanurate distilled with strong potash; the distillate neutralised with hydric chloride, evaporated and crystallised; and the amylamine obtained by distilling the hydrochloride with lime (Wurtz; Silva, Bl. [2]

Properties .- Colourless liquid, miscible with water and alcohol.

Reactions.—1. Oxidised by chromic acid to isovalerio acid (Chapman a. Thorpe, A. 142. 177).—2. With ClCO₂Et yiolds ethyl-amyl-car-bamate, C₅H₁₁NHCO₂Et (amylurethane) (Custer, B. 12, 1329).

Salts.-B'HCl: scales; sol. alcohol.-B'2H2PtCla: scales; sol. hot water, insol. alcohol. -Aurochloride: seales.

Combination .- With carbonic disulphide it forms C11H26N2S2(2C1H13N + CS2) white shining scales, insol. water and other, sol. alcohol (Hofmann, C. J. 13, 60).

Active Amylamine .- The amylamines obtained from active amyl alcohol, probably (C.H.)(CH.)CH.CH.OH ()11 (Erlenmeyer; v. AMYL ALCOHOLS; Sauer, B. 8, 1037), are optically active, and their salts are much more difficult to crystalliso than are those of the corresponding inactive compounds (Plimpton, C. J. 39, 332). Amylamine from alcohol rotating 4° for 10 cm. rotated 3° 30′ to the left; (96°-97°); S.G. 2° 17725.—Hydrochloride: deliquescent; feobly dextrogyrate. -- Platino-chloride: scales; scl. hot water. S. 2.4 at 14°.—Aurochloride: sol. alcohol; separates on slow ovaporation in lozenge-shaped orystals with the acute angles truncated.

Inactive Amylamine from inactive amylabloride. (96°-97°). S.G. \(\frac{2}{3}\) 7678; \(\frac{21^3}{3}\) 7501. —Hydrochloride: crystallises well.—Platinochloride S. 1.7 at 14°: scales.—Aurochloride: sol. alcohol; lozengo-shaped crystals with one acute angle truncated.

Diamylamine (C,H11)2NH. (186°-187°).

S.G. 2 . 7825 (Silva).

Fornation.—1. From ord. amylamine and ord. amyl bromido (Hofmann, A.79, 21).—2. From amyl oyanide and potash (Silva, Z. 1867, 457). 8. From amyl bromide and alcoholio ammonia (Custer, B. 12, 1329; Plimpton, C. J. 39, 332; Bell, B. 10, 1867).—4. From amyl chloride and aqueous NH₃ at 140°-165° (Mallot, C. R. 104, 993).—5. From the nitroso-compound (Custer, B. 12, 1333).

Properties.—Oily liquid; sl. sol. water. Reacts with Cl.CO₂Et yielding diamylurethane (Custer).

Salts.—B'HCl: laminæ; crystallises well from hot water.—(B'HCl),PtCl; i sol. alcohol, sl. sol. water. Auro-ohloride: sol. alcohol, insol. water.

Active Di-amylamine. (182°-184°). S.G. 3° 7878. From active anyl bromide. (V. active amylamine) (Plimpton, loc. cit.) Dextrorotatory

(5° 15' for 10 cm.).

Hydrochloride: soluble in water, alcohol, and ether. Much more soluble in cold water than the inactive salt. Solution rotates to the right. Platinochloride: sol. alcohol insol. water. Crystallises from dilute alcohol in octahedrons. Auroshloride: insol. water, sol. alcohol.

Inactive Di - amylamine. (186°–187°). S.G. 4.7878; 14.7776. From inactive amyl

chloride (Plimpton).

Hydrochloride: laminæ; rotates when thrown on the surface of water. Insol. ether.—
Platinochloride: sol. alcohol, insol. water. Crystallises easily from dilute alcohol in rectangular prisms.

Triamylamine (C₃H₁₁)₃N. (237°). From diamylamine and annyl bromide or from amyl bromide and ammonia (Hofmann, A. 79, 22). Amyl cyanate and potash (Silva, Z. 1867, 458).

Oily liquid, insol. water.

Salts. — B'HOI: crystalline mass with Instre of mother-of-pearl. Platinochlorido: rhombio prisms; insol. water, sol. alcohol.

Active Triamylamine. (230°-237°). S.G. 4° '7964. Prepared from active amyl bromide (3°) and active diamylamine (5° 15'). Rotates 44° 15' to the right for 10 cm. (Plimpton, loc. cit.).

Hydrochloride: syrup which solidifies over sulphurio acid. Solution strongly dextro-

over sulphurio acid. Solution strongly dextrogyrate.

Aurochloride: needles; insol. water, sol.

alcohol.
Inactive Triamylamine. (237°). S.G. 4.788.
From inactive amyl chloride and ammonia.

"Hydrochloride: crystallisos from water n prismatic needles, from ether in pearly scales.—Aurochloride: needles; sol. alcohol. Inactive triamylamine may be separated from nactive diamylamine by treating the hydro-hlorides with ether which dissolves the triamylamine salt.

Tetramylammonium Salts.

Iodide (C,H₁₁),NI. From ordinary amyliodide and triamylamine or amyl iodide and ammonia (Hofmann, C. J. 4, 316). The mixture of triamylamine and amyl iodide is boiled and after three or four days solidifies on cooling into an unctuous crystalline mass. Monoolinio laminæ (Lang, J. 1867, 491). Dissolves sparingly in water forming an extremely bitter solution from which it is ppd. in a crystalline form by alkalis. Boiled with silver oxide it yields a very bitter alkaline solution of Tetramylammonium hydroxide. mixing the liquid with potash or on concentrating, the hydroxido separatos as an oily layer, which gradually solidifies. By evaporating a solution of the hydroxido in an atmosphere free from carbonio acid, crystals containing several molecules of water are obtained. When heated these crystals melt and give off water, triamylamine, and a hydrocarbon which is probabiy amyleno.

 $(\ddot{\mathbf{C}}_{s}\mathbf{H}_{11})_{s}$ NOI: lamina with palm-like ramifications. — $((\mathbf{C}_{s}\mathbf{H}_{11})_{s}$ NOI)_sPtCI₁: orange - yellow needles. —Sulp hate: long capillary threads.—Nitrate: needles. —Oxalate: large deli-

quescent plates.

Amylamine. Corresponding to methyl propyl carbinol. (CH₃)(C₃H.)CHNH₂. (89°-91°). By reduction of methyl propyl ketone phenyl hydrazide in alcoholic solution with sodium amalgam and acetic acid (Tafel, B. 19, 1924). Mobile liquid, smelling strongly ammouiacal, miscible with water, alcohol, and other.

Hydrochloride: silky needles.—Platinochloride: yellow needles; sol. water and hot alcohol, less so in cold alcohol.—Oxalate: crystallises from hot alcohol in scales.

Tertiary amylamine. (OII_3)_*(C_2H_5)UNH_2. (77:5^2-78^2). S.G. 2.7611; 15-2.7475. Formerly considered to be (CH_3)_*CH(CII_3)CHNH_2 on account of its formation from the cyanate corresponding to Wurtz's amylene hydriodide and amyleno hydrate, then regarded as isopropyl-methyl carbinol, and now shown to be dimethyl-ethyl carbinol (Flavitzky, A. 179, 310).

Formation.—1. From pseudoamylurca and strong potash (Wurtz, Bl. [2] 7, 143).—2. By the action of dimethyl cthyl carbinol iodide on the cyanides of potassium and mercury, and treatment of the nitrile so obtained with hydric chloride (Wyschnegradsky, A. 174, 60).—3. By treating the product of the action of the same iodide upon silver cyanate with strong hydric chloride (Rudnew).

Properties.—Odour ammoniacal; pps. copper salts but does not redissolve the ppd. cupric

hydrate.

Hydrochloride: efflorescent scales, or octahedrons, from alcohol and ether. — Platinochloride: fine crystals derived from a monoclinio prism, easily soluble water and alcohol. — Aurochloride: large yellow crystals, monoclinio.

Reactions. — With bromine it forms bromamyl-amine C₅H₁₂BrN which can be distilled with steam (Wurtz). R. T. P.

AMYL-ANILINE C₁₁H₁,N i.e. C_eH₃,NHC₈H₁₁. (258°). Mol. w. 163. From aniline and isoamyl bromide (Hofmann, C. J. 8, 297). Smolls, when cold, like roses. When its hydrochloride is

heated at \$20° it changes to the hydrochloride of amido-phenyl-pentane, C,H11.C,H4.NH2 (Hof-

mann, B. 7, 529).

Isoamyl-aniline C.H. NHC.H., (243°) at 720 mm. Colourless oil. V. sol. alcohol, ether, and benzene, insol. water. Is a by-product of the action of isovaleric aldehydo and HCl upon aniline. Salts.-B'HC1: colourless prisms, v. sol. water. The nitrate and oxalate are sparingly soluble. The picrate is a reddish-yellow oil.

Acetyl derivative C.H.N(C.H.)Ao, (278°) at 720 mm., colourless fluid, v. sol. alcohol

and ether; insol. water.

Nitrosamine CoH, N(CoH11)NO: oil; volatile with steam; v. sol. alcohol and ether, insol. water (Spady, B. 18, 3376).

Di - isoamyl - aniline (275°-280°) (Hofmann, A. 74, 156). -B'₂H₂PtCl₆. Iso - AMYL - ANTHRACENE C₁₀H₂₀ i.e.

 $C(C_bH_{11})$ C.H. [59°]. From amyl-VCH

hydro-anthranol by boiling alcoholic HCl.

Preparation.—Anthraquinone (30 g.), zinc dust (100 g.) NaOH (50 g.), water (150 g.) are boiled together for 5 hours and then amyl bromide is added. The liquid is poured off, and the pp. dissolved in alcohol, reppd. by water, and boiled with alcoholic HCl (Liebermann, A. 212, 104; B. 14, 796).

Properties.—Long sea-green needles with blue fluorescence (from alcohol). Conc. Il SO, gives a green solution. V. sol. benzone, CS2, chloroform, or benzoline. Picrate forms red needles [115°], CrO₃ in HOAc gives amyl-oxanthranol. Forms a bromo compound

C₆H₄ C(C₃H₁₁) C₆H₄. [76°]. Picrate [110°]. Forms also a corresponding chloro-derivative [71°]. l'icrato [108°].

Iso-Amyl-anthracene-di-hydride

 $C_{19}H_{22}$ i.e. $C_6H_4 < \frac{CH(C_5H_{11})}{CH_2} > C_6H_4$. (350°) ; (292°) at 570 mm. S.G. 18 1.031. Prepared by reduction of amyl-oxanthranol with P and HI (Liebermann, B. 14, 457; 15, 1000; A. 212, 79). Clear fluorescent liquid. Miscible with alcohol, ether, benzene, and acetic acid, in all proportions. On oxidation with HNO, authraquinone is formed.

Iso-AMYL ARSENATE (C.H.,),AsO, (Crafts,

Bl. 14, 101).

ISO. AMYL ARSENITE (CaH11), ASO3. (288°)

(Crafts, Bl. 14, 105).

n-AMYL-BENZENE. Phenyl-pentane. C,H i.e. Ph.CH₂.CH₂.CH₂.CH₂.CH₃. Mol. w. 148. (201° uncor.) at 743 mm. S.G. ²² 8602. From benzyl bromide, n-butyl bromide, and sodium (Schramm, A. 218, 388). Pleasant smelling oil.

Reaction.—Bromine vapour at 150° gives

*Ph.CHBr.CH_.CH_.CH_.CH_.CH, (?) *which on
distillation gives Ph.CHI.CH..CH_.CH_.CH. distillation gives Ph.CII:CH.CH₂.CH₃.CII₄ (210°-215°) which combines with Br₂ forming

Ph.CHBr.CHBr.CH, CH, CH, . [54°].

Iscamyl-benzene (193°) at 736 mm. S.G. 12 859. From bromo-bonzeno, isoamyl bromide, and Na (Fittig a. Tollens, A. 129, \$369; 131, 313; Bigot a. Fittig, A. 141, 160; Schramm, A. 218, 390). Also from isoamyl chloride, benzene, and AlOl, (Friedel a. Crafts, A. Ck. [6] 1, 454).

Reaction.—Bromins exposer at 150° gives Ph.CHBr.CH., CH(OH.), (?) which on distillation gives HBr and Ph.CH.CH.CH.(CH.), phenylisoamylene, which forms a dibromide [129°].

District of the property became Ph. CHE. (179°).

Di-ethyl-oarbinyl-bsnzene Ph.CHEt2. (178°).

Formation.—1. From benzylidene chloride and zine ethide (Lippmann a. Luginin, Z. 1867, 674).-2. From benzo-trichloride, Ph.CCl, and zine ethide (Dafett, M. 4, 153, 616).

Tert-amyl-benzene Ph.CMc.Et. (c. 187°). S.G. 2 874. From tert-amyl chloride, benzene,

and AlCl₃ (Essner, Bl. [2] 36, 212).

Di-iseamyl-benzene C_{1a}H₂₀ i.e. (C₃H₁₁)₂C₆H₄. (o. 265°). S.G. 2·887. From benzene, isoamyl chloride, and AlCl, (Austin, Bl. [2] 32, 12). SULPHONIC

AMYL-BENZENE C11H16SO3 i.e. C5H11 C6H4SO3H.

Isoamyl-benzene sulphonic acid. Deliquescont crystalline mass (Fittig a. Tollens, A. 131, 315). Salts.—KA' aq.—BaA'₂: hair-likeneedles.

Di-ethyl-carbinyl-benzene sulphonic acid. CHEt, C, H, SO, H. Salts. BaA', 1 aq: poarly leaflets, sl. sol. water and alcohol (Dafert, M.

4, 617).

p-I_SO-AMYL-BENZOIC ACID C₁₂H₁₀O₂ i.e. C₆H₁(C₅H₁₁).CO₂H [1:4] [158°]. Formed by saponification of the nitrile. Sublimes in flat colourless needles. Sol. alcohol, ether, and hot water, sl. sol. cold water. Salt: AgA': small colourless needles, sl. sol. cold water (Kreysler, B. 18, 1710).

p-Iso-AMYL-BENZONITRILE

C.H.(C.II.).CN (260°-263° uncorr.). Colourless oil. Formed by heating tri-isoamylphenylphosphate with dry KCN; yield -20 p.c. (Kreysler, B. 18, 1709).

Iso - AMYL BORATE C₁₅H₃₃BO₃ (C,II,O),B. (254°). S.G. 2.872.

Iso-amyl borate (C, H₁₁, O)BO. S.G. 2 971. Oil. n.AMYL BROMIDE G₂H₁₁Br i.e. CH₂CH₂CH₂CH₂CH₂Br. Mol. w. 151. (129°). S.G. 21246. From n-amyl alcohol (Lieben a. Rossi, A. 159, 73).

Inactive amylbromide (Cll2)...Cll.CH2.CH2Br (120.6° i. V.) at 734 mm. S.G. 2 1.026 (Lacho-

wicz, A. 220, 171).

Iseamyl bremide (118.5°) at 756 mm. (R. 1if, B. 19, 563). II.F.p. 31,000 (Berthelot). Schiff, B. 19, 563). S.V. 138.6 (S.); 143.8 (Ramsay). For a specimen which rotated + 52° in 100 mm. Perkin (C. J. 45, 458) found: (120.5° cor.); S.G. 15. 1-2193; 25. 1-2083; M.M. 9-01 at 17°.

Active amyl bromide (117°-120'); S.G. 11 1.225 (Le Bel, Bl. [2] 25,545). Dextrorotatory;

 $a = +3.75^{\circ}$.

n-Sec-amyl bromide CH, CH, CH, CHBr.CH, (113°) (Wurtz, A. 125, 118). Formed when isoannyl bromide is heated at 230° (Eltekow, B. 8, 1244).

Iso-sec-amyl bromide (CH3)2.CH.CHBr.CH4 (116°) (Wyschnegradsky, A. 190, 357).

Tert.amylbromideCH3.CH2.CBr(CH3)2.(100°). AMYL-BROMO- v. BROMO-AMYL

Iso-AMYI-CARBAMIC ETHER C,H,,NO, i.e. C,H,,NHCO,Et. Amyl-urethane. (218°). S.G. 93. From isoamyl-amine and ClCO,Et (Custer, B. 12, 1328). Oil; sol. alcohol and ether.

Di-iscamyl-carbamic ether (C,H11)2N.CO2Et. (247°). From di-isoamyl-amine and ClCO₂Et(C.).

Yor. I.

Iso-AMYL CARBAMINE O.H., N i.s. C.H., NO. 187°). Mol. w. 97 (Hofmann, A. 146, 109).

180°.AMYL CARBONATE C₁₁H₁₂O₃ i.e.
(C₈H₁₁),CO₄ (229° cor.). S.G. ¹² · 91.

180°.AMYL CETYL OXIDE C₂₁H₄₄O i.e.
(C₈H₁₀O₄), S.G. ¹⁸ · 91.

180°.AMYL CHIVE C₈H₁₀C i.e.
(C₈H₁₀C C₈C, C₈C). Plates.

7.AMYL CHLORIDE C₈H₁₁Cl i.e.
(CH₂CH₂CH₂CH₂CH₂Cl. Mol. w. 106°5. (106°).

18 G. 22°. 873° (Jachawicz 4 290° 1011). 802° (106°).

S.G. 22 873 (Lachowioz, A. 220, 191); 883 (L. a. R.). Formed from n-amyl alcohol (Lieben a. Rossi, A. 159, 72; G. 1, 314) or by the chlorination of n-pentane (Schorlemmer, A. 161, 268).

Inactive amyl chloride (CH₃)₂CII.CH₂.CH₂Cl. (99.8°-100.5°). S.G. 29.870. From iso-pentane

(Lachowitz).

Iso-amyl chloride (99.5°) (R. Schiff, B. 19, 562). S.V. 134.4 (S.); 136.5 (Ramsay). In a specimen which rotated +5.8° in 100 mm. Perkin (C. J. 45, 452) found: (97-99° cor.); S.G.

ls 8801; 25 8716; M.M. 7:17 at 19:5°.
Formation.—1. From isoamyl alcohol and HCl (Balard, A. Ch. [3] 12, 294), S₂Cl₂ (Carius a. Fries, A. 109, 1), or PCl₃ (Cahours, A. 37, 164).

Reactions .- 1. Converted into amyl alcohol by water at 100° (Butlerow, A. 144, 34), or better at 120° (Niederist, A. 186, 392).—2. H.SO., forms HCl and C.H., SO., H (Oppenheim, J. pr. 102, 339).

Active amyl chloride

CH, CH, CH(CH,).CH,Cl. (99°). $a = +1.24^{\circ}$

S.G. 15 ·886 (Lo Bel, Bl. [2] 25, 546). n-Sec-amyl chloride CH₂.CH₂.CH₂.CHCl.CH₃. (104°). S.G. 21 ·891. From n-pentano by chlorination (Schorlemmer; Lachowioz). CH3.CH2.CH:CH.CH3 and HCl (Wagner a. Saytkeff, A. 179, 321).

Iso-sec-amyl chloride (CH.), CH.CHCl.CH. (91°). S.G. 2° 88. From (CH₃), CH. CHICH, and HCl (Berthelot, C. R. 56, 700; Wurtz, A. 129, 368; Wysohnegradsky, A. 190, 357).

s-Sec-amyl chloride Et,CHCl. (103°-105°). S.G. 21 895. From the alcohol (W. a. S.).

Tert-amyl chlorids EtCMe₂Ol. (86°). S.G. 19 870. By action of PCl, on tert-amyl alcohol or on methyl-isopropyl-carbinol (Wyschnegradsky, A. 190, 336; 191, 331).

AMYL-CHLORO- v. Chloro-AMYL.

CYANATE O₈H_{II}NO Iso - AMYL C.H., N.CO. (135°). Prepared by distilling amyl-carbamic other with P2O5 (Custer. B. 12, 1330), or from C₃H₁₁SO₄K and potassium cyanate (Wurtz, A. Ch. [3] 42, 43). With ammonia it yields amyl-ures, and with potash it yields amylamine.

AMYL CYANIDE v. AMYL CARBAMINE and mitrile of Hexoro Acid.

AMYLENE C.H. Pentene. Mol. w. 70. n-Propyl-ethylene CH, CH, CH, CH; CH; CH;

Formation .- 1. Together with amyl acetato when KOAc and Ao₂O act on n-amyl chloride at 200° (Schorlemmer, A. 161, 269).-2. Together with di-allyl (principal product), pentane, and other bodies, in the action of zino ethide on allyl iodido (Wurtz, A. 123, 203; 127, 55; 148, 131).

Properties .- Liquid; insol. H2SO, (2 vols.)

diluted with water (1 vol.).

Reactions .- 1. Gives Pr.CHI.Me with HI. 2. Alkalins KMnO, gives succinio, butyric, oxalic, and formio acids (Zoidler, A. 197. 253).

Isopropyl-ethylene (CH_a), CH.CH:CH₂. (21°). Formed together with EtCMe:CH₂ by action of alcoholic KOH on isosmyl iodide (Wysohnegradsky, B. 10, 81; A. 190, 328).

Properties .- Liquid; insol. at 0° in HaSO.

(2 vols.) mixed with water (1 vol.).

Reactions .- Does not combine with HI at -20°, but at 20° it combines slowly forming (CH₃)₂CH.CHI.CH₃.

s-Mathyl-sthyl-ethyleue CH, CII:CII.Et.

Formation.—1. From CH3.CH2.CHI.CH2.CH3 Waguer a. Saytzeff, A. 175, 373; 179, 302), or CH, CH, CH, CH, CH, (Wurtz), and an alcoholic solution of KOH.—2. From ethyl-crotonic acid, CH₂.CH:CEt.CO₂H, by combining it with IIBr and noutralising the resulting β-brome-diethyl-acetic acid: CH3.CHBr.CEtII.CO2Na=

NaBr + CO₂ + CII, CH:CEtH (Fittig, A. 200, 27). Reaction.—HI forms CH, CHI.CH, CH., CH., CH., Et.C(CH₃):CH₂. u-Mathyl-sthyl-sthylene (32°). S.G. 2 670. From active amyl iodide and alcoholic KOH (Le Bel, Bl. [2] 25, 546).

Properties.-Liquid; dissolves in H,SO.

(2 vols.) diluted with water (1 vol.).

Reactions.—III forms CH3, CH2, C1(CH3).CH3 Tri-methyl-ethylens (CH3)2C:CH.CH3. (36° S.G. 15 ·6704; 25 ·6614. M.M. 6·121 at 13·2° (Perkin, C. J. 45, 448).

Formation.—1. From CH3.CH2.CI(CH3)2 (Ermolajeff, Z. [2] 6, 275) or CH3.CH1.CH(CH3)2 (Wy.) and alcoholic KOH.—2. From ethyl isoamyl oxide and P₂O₅ (Flavitzky, A. 169, 206).

Properties.-Liquid; solublo at 0° in H2SO4 (2 vols.) diluted with water (1 vol.)

Reaction.—HI forms (CH₃)₂C1.CH₂.CH₃.

Iso-amylene. (36°). S.G. 9-9 ·661; 20 ·648. H.F.p. 10,600 (Berthelot); 18,970 (Th.). H.F.v. 16,650 (Th.). V.D. 2-47 (for 2-42). S.V. 110 (R. Schiff, A. 220, 89); 110-8 (Ramsay). μ_{β} 1:3813. R_{∞} 39-29 (Brühl). A mixture of trimethyl-ethyleno (90 p.c.) and u-methyl-ethyl sthyleno (10 p.c.) with a small quantity of isopropyl-ethylene (Flavitzky, A. 179, 340).

Preparation.-From isoamyl alcohol and ZnCl2, many other hydrocarbons being also formed (Etard, C. R. 86, 488; Wyschnegradsky,

C. R. 86, 973).

Properties.—Absorbed at 0° by H₂SO₄ (2 vols.) diluted with water (1 vol.), with production of tertiary amyl alcohol. A more dilute acid (2 pts. H.SO, to 1 pt. water by weight) forms methyl-isopropyl-carbinol (Ossipoff, B. 8, 542, 1240).—NOCI forms a compound C₃H_{1e}NOCI which may be reduced to amylamino (Tonnies, B. 12, 169).—3. C.H., K.PtCl, aq is formed by boiling isoamyl alcohol with PtCl, and then adding KCl (Birnbaum, A. 145, 73); deliquescent plates.

Other References .- Balard, A. Ch. [3] 12,820; Frankland, C. J. 3, 35; Bauer, Sitz. B. 44 [2] 87; Z. 1866, 380, 667; Bauer a. Klein, Z. [2] 4, 386; Guthrie, A. 121, 108; Lippmann, A. 129, 81; M. 5, 559; Eltekoff, B. 6, 1258; Linnemann, A. 143, 350; Buff, A. Suppl. 4, 143; 148, 349; Thorpe & Young, A. 165, 7; Flavitzky, A. 165, 157; Le Bel, Bl. 17,3; 18, 166; Borthelot, A. Ch. [4] 9, 442; C. R. 44, 1350; Renard, A. Ch. [6] 1, 227; Markownikoff, Z. [2] 2, 502.

Oxidation of amylenes .- Examined by Zeid.

ler, A. 186, 245; 197, 253; Truchot, C. B. 63,

274 : Berthelot, C. R. 64, 86.

Di-amylene $C_{10}H_{20}$. (156°). S.G. $\frac{19}{2}$ 780. R_{\odot} 76.58 (Nasini a. Bernheimer, G. 15, 93). R₂₀ 76:58 (Nasini a. Derinionio, ... S.V. 211:18. Occurs in the product of action of ZnCl2, H2SO4, or P2O3, on isoamyl alcohol; and is also formed by shaking amyleno with H2SO.

Reactions.-1. Bromine forms C10 H20 Br2-2. Chromic acid mixture produces a methenic acid C, H14O2 (Schneider, A. 157, 213; Pawlow,

J. R. 9, 75).

Combinations.—C₁₆H₂₀S₂Cl₂; from amyleno and S₂Cl₂ (Guthric, C. J. 12, 112; 13, 35; 14, 128). Distilled over KOH it forms C10H1082, $(112^{\circ}),$ S.G. 13 ·880. ZnEt₂ gives C₁₄H₃₀S₂ (240°-250°).

References .- Balard, A. Ch. [3] 12, 320; Bauer, Bl. 1863, 332; 1867, 341; Berthelot, C. R. 56, 1242; Walz, Z. [2] 4, 315; W. v. Schneider, A. 157, 185; Wyschnegradsky, B. 8, 434; Lebedcff, J. R. 7, 246; Tugolessoff, B. 12, 1486.

Triamylene C1, H30. (248°). S.G. 81. V.D. 7.6 (for 7.4). Among products of action of ZnCl₁ on isoamyl alcohol (Bauer, Sitz. B. 44 [2] 87; A. 137, 249; 147, 254). Forms a bro-[2] 87; A. 161, 242; 171, 252; Frans a bounded of the service of t

isoamyl alcohol (Balard; Baucr).

AMYLENE DI-ACETIN v. di-Oxy-pentane. AMYLENE BENZOATE v. di-Oxy-pentane. AMYLENE BROMIDE v. di-Bromo-propane.

AMYLENE TRI-CARBOXYLIC Ether.—Et, A." (151°). Obtained by introducing allyl into ethane tri-carboxylio acid (Hjolt, B. 16, 333). At 160° it splits up into CO2 and allyl-succinic acid (q. v.).

AMYLENE CHLORHYDRIN v. CHLORO-

AMYLENE CHLORIDE v. di-CHLORO-PENTANE. AMYLENE - CHLORO - SULPHIDE v. di-AMYLENE, Combinations

AMYLENE IS-ETHIONIC ACID v. Oxy-

PENTANE SULPHONIO ACID.

AMYLENE GLYCOL v. di-Oxy-pentane.

AMYLENE GUANAMINE C.H. 15N5. [178°]. Formed by heating guanidine caproate (hexoatc) at 225° (Bandrowski, B. 9, 243). Crystals; v. sl. sol. water, v. sol. alcohol. Salt.—B'HCl.

AMYLENE HYDRATE. Tertiary

ALCOHOL (q. v.).

AMYLENE HYDRIDE. PENTANE (q. v.). HYDROCHLORIDE. AMYLENE

CHLORIDE (q. v.).

AMYLENE NITRITE CaH10N2O4 O₃H₁₀(NO₂)₂. From amylene by treatment with NO, or furning HNO, (Guthrie, C. J. 13, 45, 129). Tables; decomposed at 95°.

AMYLENE OXIDE C.H.,O.

Isopropyl-ethylene oxide Pr.CH

(82°). By action of potash on chloro-amyl alce-Pr.OHCl.CH.OH or PrCH(OH).CH.Cl (Eltekoff, Bl. [2] 40, 23; J. R. 14, 355). Heated with water for 50 hours at 100° it forms Pr.CH(OH).CH₂OH. Does not combine with NaHSO.

Tri-methyl-ethylene oxide Me.C (76°). S.G. 2 ·829.

Preparation .- By action of potash on the chloro-amyl alcohol obtained by the union of Me₂C:CHMe with HClO (E.).

Properties. - Liquid; readily unites with cold water to form di-oxy-pentane. Does not combine with NaHSO.

Methyl-cthyl-ethylene oxide MeCH-CHE (80°). Prepared by action of potash on the chloro-amyl alcohol resulting from union of Me.CH:CHEt with HCIO (E.). Unites at 100° with water forming MeCH(OH). CHEt(OH).

Di-amylene oxide $\rm C_{10}H_{20}O.~(170^{\circ}-180^{\circ}).~From$ di-acetyl-di-oxy-decame (q.~v.) (di-amylene diacetin) and solid KOH (Bauer, Sitz. B. 45, 276) Oil. Reduces ammoniacal AgNO₂.

Di-amylene oxide (198°-203°). (for 5.4). Obtained as an oil by the action of potash on a mixture of amylene and Bz,O, that has been heated at 110° (Lippmann, M. 5, 562). Does not reduce ammoniacal AgNO, or combine with NaHSO3.

Di-amylene oxide (180°-190°). From diamylene and ohromic mixturo (Schneider, A. 157, 221). Reduces ammoniacal AgNO. Oxidiscd to amethenic acid, C,H1,O2.

Di-amylene oxide (193°). From di-amylene bromide (v. di-Bromo-Decane), water, and PbO (Eltekoff, J. 1878, 374).

Two or more of the preceding di-amylene

oxides may bo identical.

AMYLENE SULPHIDE C₅H₁₀S. (c. 200°). S.G. 12 907. Formed by boiling O₁₀H₂₀S₂Cl₂ (v. Diamylene) with zine (Gnthrie, C. J. 14, 128). Colourless oil.

Amylene snlphide (?) O₅H₁₀S. (130°-150° V.D. 3.2 (calc. 3.5). Formed by action of acids or of heat upon the product of the union of ZnEt₂ and CS₂ (Grabowsky, A. 138, 165). Alcoholic HgCl2 forms plates of C5H108 HgS HgCl2; alcoholic AgNOs forms CsH10OAg2OAgNOs.

AMYL ENNONYL KETONE (?) C18H24O i.s. Et_CH.CO.C,H₁(O,H₃), (?). (280°-300°). One of the products got by passing CO over a mixture of NaOEt and NaOAc at 250° (Geuther a. Fröhlich, A. 202, 312).

AMYL ENNYL KETONE O, H28O i.e. C.H., CO.C. H., (?). Amyl - valerone. (209°). S.G. 12 ·845. One of the products of the passage of CO over sodium iso-amylate at 100° (Geuther a. Fröhlich, A. 202, 301). Liquid; smells like quinces. Does not combine with NaHSO.

AMYL ETHER v. AMYL OXIDE.

AMYL FLUORIDE *C,H,F. (72°-92°). mixture of amyl fluorido and polymerides of amylone is formed by saturating amylene at 0° with HF (S. Young, C. J. 39, 489).

AMYL-GLYOXALINE C.H., i.e.

 $CH \leqslant^{\rm NH}_{\rm CH.N} \geqslant C.C_5H_{11}\ (?). \ \ \textit{Glyoxal-anathyline.}$ [84°]. From cenanthol-ammonia and glyoxal (Radziszowski, B. 16, 748). Thin glistening needles. Sol. alcohol, al. sol. ether, insol. water.

Iso-amyl-glyoxaline C.H., N. s.c. $OH < N(O, H_{11}) > OH (?)$. (240°-245°).

B.G. 11.94. From glyoxaline and amyl bromide (Wallach, A. 214, 322; B. 15, 651). Liquid; v. sl. sol. water, sol. even in vory dilute alcohol. Salt .- B'2H2PtCla: plates (from alcoholio

HCl); v. sl. sol. cold water or cold alcohol.

Iso-AMYL HEPTYL OXIDE C12H26O C₅H₁₁,O.C₇H₁₅. Amyl-ananthyl ether. (221°). S.G. ²² 668. V.D. 6.57 (celc. 6.45). From sodium heptylate and isoamyl iodido (Wills, C. J. 6, 316).

(Py. 2:3)-AMYL-HEXYL-QUINOLINE

C₂₀H₂₉N i.e. C₆H₄ CH:C(C₅H₁₁) N:C(C₆H₁₃) (320°-360°).

Oily fluid. Formed by the action of conanthic aldehyde and HCl upon aniline (Docbner a. Miller, B. 17, 1719)

Salts.—B'2H2Cl2PtCl4: large yellow plates. -B'CeH2(NO2)3OH: yellow needles; sl. sol. water and cold alcohol.

AMYL HYDRIDE v. PENTANE.

Iso-AMYL-HYDRO-ANTHRANOL C19H22O or $C_6H_4 < \frac{C(C_5H_{11})(OH)}{C_6H_4} > C_6H_4$. [74]. Formed, as

a by-product in the treatment of anthraquinone with zinc-dust and amyl bromide (Lichermann a. Tobias, B. I4, 801; A. 212, I02). Crystallino solid. Insoluble in water, extremely soluble in other solvents. On boiling with alcoholic IICl it gives amyl-anthracenc.

Iso - AMYL - HYDROQUINONE. From isoamyl-arbutin and dilute H.SO., glucose being also formed (Schiff a. Pellizzari, A. 221, 365). Needles. Gives a crystalline nitro-derivative.

AMYLIDENE - ACETO - ACETIC ETHER

v. p. 24.

Iso-AMYLIDENE-m-AMIDO-BENZOIC ACID C₁₂H₁₅NO₂ i.e. C₄H₉ CH:N.C₆H₄.CO₂H. [c. 130°]. From valeric aldehyde and m-amido-benzoio acid (Schiff, A. 210, 119)

AMYLIDENE ANILINE C₁₁H₁₃N i.e. Me.CH.CH.:CH:NPh. [97°]. From valerio aldehyde aud aniline in the cold (Lippmann a. Strecker, B. 12, 74). Prisms. — B'HCl. — B'HCl. — Valeric aldehyde and aniline at B'₂H₂PtCl₆. Valeric aldehyde and aniline at 100° form di-amylidene-di-phenyl-diamino, a neutral oil, C., H., N. (Schiff, B. 12, 298).

AMYLIDENE BIURET C, H, N,O2. From valeric aldehyde and cyanio acid (Baeyer, A.

114, 164)

AMYLIDENE BROMIDE v. di-BROMO-PENT. ANE.

AMYLIDENE-DI-CARBAMIC ETHER

C11H22N2O4 i.e. Mo2CH.CH2CH(NH.CO2Et)2. Amylidene urethane. [126°] From carbamic ether, valeric aldchyde, and conc. HCl (Bischoff, B. 7, 633). Needles. Split up by hot dilute acids into valeric aldchydo and carbamio ether.

AMYLIDENE CHLORIDE v. di-CHLORO-

a AMYLIDENE GLYCOL v. ortho-Valerio al-DEHYDE.

AMYL IODIDES C.H., I. Mol. w. 198. n. Amyl iodide CH., CH., CH., CH., CH., I. (156° cor.) S.G. 2 I 544; 22 I 517. From the chloride and HI (Lieben a. Rossi, A. 159, 74).

Iso-amyl iodide. (I48°). S.G. 15 1.510; Na (Leone, G. 12, 2 1.498. M.M. I3.20 at 19.6° (Perkin, C. J. 45, palo yellow needless.

482). S.V. 151-08 (B. Schiff, B. 19,564). From isoamyl alcohol (4 pts.), iodine (5 pts.), and P (Cahours, A. Ch. [2] 70, 81; Grimm, J. pr. 62, 885). From amyl-chloride and Cal, 3 aq at 100° (Van Romburgh, R. 1, 151). Partially converted by heating with EtOH into Et1 and isoamyl alcohol (Friedol a. Crafts, A. 130, 198).

Active amyl iodide EtCHMc.CII,I. (1440-145°). S.G. $\frac{1}{4}$; I·5425 (Just, A. 220, 152). $\alpha = 3.76$ ° for 100 mm.at 16° (J.); 5·2° (Le Bel, Bl. [2] 25, 542). From the alcohol by HI. Reduced in alcoholic solution by Sn and conc. HCl. to inactive isopentane (J.)

u-n-Sec-amyl iodide CH3.CH2.CH2.CH1.CH3. (I44°-145°). S.G. 2 I.539. Formod by union of III with CH, CH, CH, CH; CH; (Wagner a. Saytzeff, A. 179, 313; Wyschnegradsky, A. 190, 347) or CH3.CH2.CH:CH.CH3 (Wurtz, A. 148,

132).

Iso-see-amyl iodide (CII,)2CH.CHI.CH, (137°-139°). From (CH₃)₂CH.CH:CH₂ and HI (Wy.). Water and PbO convert it into (CH₃)₂C(OH).CH₂.CH₄.

s-n-Sec-amyl iodide CH₂.CH₂.CIII.CH₂.CH₃. (145'-146°). S.G. 2 I 528; 22 I 50. From di-

ethyl-carbinol and HI (W. a. S.)

Tert-amyl iodide (CH₃)₂Cl.CH₂CII₃. (I29°), S.G. ⁹ 1·524; ¹⁹ 1·50. From iso-sec-amyl iodide and HI (Winogradoff, A. 191, 132); also from tert-amyl alcohol and HI (Wy.). By shaking with water for 1 hours it is almost completely converted into tert-amyl alcohol (Bauer, A. 220, 158). With McOH at 100° it forms McI and tert-amyl alcohol; MeOAo at 110° gives amylenc, McI, and HOAc.

DI-n-AMYL KETONE C₁₁H₂₂O i.e. (C₅H₁₁)₂CO. Caprone. [15°]. (226° cor.). S.G. 20 ·826. Prepared by distilling calcic caproate. Does not

combino with NaIISO,

Reactions .- 1. Conc. HNO3 forms caproio nitro-valeric, and oxalic acids. - 2. CrO, forms caproic and valeric acids (E. Schmidt, B. 5, 601; Lieben a. Janecck, A. 187, 134; Herez, A. 186,

n - AMYL - MALONIC ACID CaH14O4 i.e. (C.H.,)CH(CO.H)2. [82°]. Formed by saponifying the product of the action of KCy upon a-bromo-lioptoio ether (Hell a. Schüle, B. 18, 626). Split up at 140° into CO2 and n-hexoic acid.

Salts.-CaA": S. .04 at 18°.-SrA": S. .09 at 16°.—BaA": S. ·6 at 10°.—CdA".—PbA": S. ·008 at 20°.—Ag₂A".

TRI-Iso-AMYL-MELAMINE C18H36Na i.e. C3II3 (C3II11)3N6. Formed by desulphuration of isoamyl thiocarbimide (Hofmann, B. 3, 264). Thick oil.—B"H,PtCl

Iso-AMYL MERCAPTAN C,H,S C, H, SH. Mol. w. 104. (120° i.V.) (Beckmann); (I18°) (Nasini, G. 1883, 302). S.G. 29 8348. R. 31 94 (N.). From isosmyl chloride and KHS (Balard, A. 52, 313) or C.H. SO.K (Krutzsch. J. pr. 31, 1).

AMYL MUSTARD OIL v. AMYL THIO.

AMYL-NAPHTHALENES C1, H18.

(a) -Ist - amyl - naphthalene $C_{10}H_{11}C_{3}H_{11}$ [1]. Formed by heating an othereal solution of (a). bromo-naphthalono and isoamyl bromide with Na (Leone, G. 12, 209) .- Piorate, [85°-90°] : (6) - Lac-amyl - naphthalene C₁₀H₁, C₂H₁, [3], (c. 290°). From naphthalene, isoamyl chloride, and AlCl₁ (Roux, Bl. [2] 41, 379).—Picrate C₁₁H₁₀O₂H₂(NO₂)₂OH. [105°-110°]. Amyl-naphthalene (?), (305°). From lapachic acid, HI and P (Paterno, G. 12, 369).—Picrate

[141°]: orange needles.

Iso-AMYL NITRATE C.H., NO. Mol. w. 133. (147°). S.V. 153.59 (R. Schiff, B. 19, 567). From urea nitrate (10 g.), isoamyl alcohol (40 g.) and HNO₂ (30 g.) (P. W. Hofmann, A. Ch. [3] 23, 374). Liquid, smelling like bugs.

Jso-AMYL NITRITE C, H, 1NO, Mol. w. 117. (96°) (B.). (99°) (Guthrie, A. 111, 82). S.G. 9. H.F.p. 48,140. H.F.v. 44,660 (Th.).

Preparation. — 1. Nitrous vapours (from As O, and HNO, of S.G. 1.52) are passed into isoamyl alcohol (Balard, A. Ch. [3] 12, 318; Hilger, Ar. Ph. [3] 4, 485; Williams a. Smith, Ph. [3] 16, 409).— 2. By distilling together KNO₂, isoamyl alcohol, and dilute H₂SO₄ (Rennard, Russ. Zeitschr. Pharm. 1874, 1). Yellowish liquid, smelling like nitrous ether.

AMYLNITROUS ACID, so-called. C.H, N.O. Obtained by action of HNO, on di-amyl ketono (Chancel, C. R. 94, 399). Liquid; may be

reduced to n-valeric acid. Salt.—C, H, KN,O,: greasy-looking plates.

AMYLODEXTRIN v. DEXTRIN.

AMYLOID v. STARCH. AMYLONITROPHOSPHOROUS ACID, socalled. C_{1e}H_{zz}PNO₄ (?). An oil, got by action of P₂O₂O₃ on isoamyl nitrite (Guthrio, A. 111, 65).

AMYL OXALATE v. OXALIC ACID. Iso-AMYL OXAMIDE C,H₁₁N₂O₃ i.e. NH₂·CO.CO.NllC₃II₁₁. [181°]. From isobutylisoanuyl glyoxaline and H2O, (Radziszewsky a.

Szul, B. 17, 1296). Di-iso-amyl oxamide C₁₂H₂₄N₂O₂ i.e. O₅H₁₁.NII.CO.CO.NHC₅H₁₁. [129°] (Wallach a. Schulze, B. 13, 516). [1395] (Wurtz). Silky needles; insol. water. From isoamylamine and ethyl oxalate.

AMYL-OXANTHRANOL v. OXANTHRANOL.

AMYLOXIDEC, II 220. Amylether. M. w. 158. Iso-amyl oxide $(C_sH_{11})_2O$. (173°). S.G. $\frac{15}{15}$ *7807; \$\frac{1}{25}\$ '7741. M.M. 11 168 at 15 6° (Perkin, C. J. 45, 474). From potassium isoamylato and amyl iodido.

Iso-sec-amyl oxide (Pr.CMeII) O. (163°). From Pr.CMeHI and Ag.O (Wurtz, A. 129, 366). Iso-AMYL-PHENOL C₁₁H₁₀O i.e.

C₃H₁₁.C₆H₄.OH [1:4]. [93°]. (250°). Formed by heating phonol with isoamyl alcohol and ZnCl₆ at 180° (Liebmann, B. 15, 151) or by the action of nitrous acid upon amido-phenyl-isopentane (Calm, B. 15, 1616). Long needles; sl. sol. water.

Benzoyl derivative C10H15OBz. [81°]. (349° cor.). Flat needles, formed by distilling tri-isoamyl phosphate with NaOBz (Kreysler, B. 18, 1717)

AMYL-PHENYL. v. PHENYL-AMYL-.

Iso-AMYL-PHENYL PHOSPHATE

C13H45PO4 i.e. (C5H11.C4H4.O)3PO. (above 400°). Formed by heating isoamyl-phenol with POCI, (Kreysler, B. 18, 1701). Thick oil; v. sol. ether, sol. alcohol.

ISO-AMYL-PHENYL BILICATE C44H60SiO4 i.e. (C₃H₁₁.C₆H₄.O)₄Si. (c. 394°) at 118 mm. From isoamyl-phenol and SiOl, (Hertkorn, B.

18, 1692).

AMYL PHOSPHATES.

Iso-amyl-phosphoric acid (C,H,O).PO(OH), From syrupy phosphoric soid and amyl alcohol at ordinary temperature (Guthrie, C. J. 9, 134). Deliquescent crystalline mass; v. sol. water, and alcohol, insol. ether. Salts.—K.A".— (NH₄)₂A".—BaA".—PbA".—CuA".—Ag₂A".

Di-iso-amyl-phosphorio aoid (C,H11O)2PO(OH). From amyl alcohol and bromide of phosphorus (Krant, A. 118, 102). Salts.—CaA'₂: S. 1·6 at 18°.—AgA'.—AgHA'₂. AMYL-PHOSPHINES v. PHOSPHINES (Hof-

mann, B. 6, 297).

His. 5, 50, 1919. The C.H., PH. (107°). Di-iso-amyl phosphine (C.H.,). PII. (c. 218°). Tri-iso-amyl phosphine (C.H.,). P. (300°). $Qxide (C_5H_{11})_3PO.$ [e. 65°].

Iso-amylo-iodide (C.H.,).PI.
Iso-AMYL PHOSPHINIC ACID C.H.,PO. i.e. C, H11.PO(OH)2. Pentane phosphinic acid. From isoamyl phosphine and HNO. (S.G. 1.35). Pearly plates (from water) Salt .- Ag2A": amorphous pp. (Hofmann,

B. 6, 305).

AMYL PHOSPHITES.

Iso-amyl phosphorous acld (CsH10)P(OH)8. Formed together with di-isoamyl-phosphorous acid by shaking with water the product of the action of PCl, on isoamyl alcohol. Dilute Na₂CO₃ dissolves mono- but not di-, amyl phosphite (Wurtz, A. Ch. [3] 16, 227).

Chloride C.H., O.PCl. (173). S.G. 21:109 (Mensehutkin, A. 139, 348).

Di-iso-amyl-phosphorous acid (C,H,O),P(OH). S.G. 12 .97.

Tri-iso-amyl phosphite (C,H,1O),P. (236°), in hydrogen. From PCl, and NaOC, H₁₁ (Williamson a. Railton, C. J. 7, 218).

AMYL-PIPERIDINE C10H21N i.e. $C_3H_{10}N(C_3H_{11})$. (188°). Colourless liquid, nearly insoluble in water. Formed by digesting piperidine with amyl bromide and aqueous KOH.

Methylo-iodide. B'Mel. [195°]. Thick prisms. By moist Ag₂O it gives an alkaline hydrato which on dry-distillation yields methylamyl-piperidine (Schotten, B. 15, 421).

Iso-AMYL-PYRROL CaHINN i.e. CaHINCAH (c. 182°). S.G. 10 879. Formed by distilling isoamylamino mucate (C. A. Bell, B. 10, 1866).

Iso-amyl-pyrrol carboxylio acid, isoamylamide C.H ... NC, H .. CO.NHO, H ... [77°]. Prisms. Formed along with the isoamyl-pyrrol (B.).

Iso-AMYL SILICATE C20H44SiO4 Si(C₃H₁O). (324°). S.G. 22° 868. V.D. 15°2 (6216. 13°0). From SiCl, and isoamyl alcohol (Ebelmen, A. 57, 331). Oil, very slowly decomposed by water.

AMYL SULPHATES.

Iso-amyl snlphnrio acid C₂H₁₂SO₄ i.e. C₂H₁₁SO₄H (Cahours, A. Ch. [2] 70, 86; Kekulé, A. 75, 275).

Salts. — NH,A.—NaA'laq. — KA'aq.— MgA'₂4aq.—CaA'₂2aq.—SrA'₂2aq.—BaA'₂2aq: flat tables, S. 9·7 at 10° (Balbiano, B. 9, 1487); S.G. 1623 at 21:2° (Clarke, B. 11, 1506).— ZnA', 2aq.—HgA', 2aq.—PbA', aq.—MnA', 4aq.— NiA', 2aq.—CuA', 4aq.—AgA'.

Iso-amyl sulphate (C.H.1), SO. Formed by passing SO, into warm amyl nitrite (Chapman,

B. 8, 920).

AMYL SULPHIDES.

Di-iso-amyl-sulphide (C₃H₁₁)₂S. Mol. w. 174. (214° i.V.). S.G. ²⁰ ·8431. R $_{\infty}$ 54·2 (Nasini, G. 13, 802). Amyl alcohol (131°-132°) is converted by PCl, into amyl chloride and this is mixed with alcoholic K2S (from half saturation of alcoholio KOII with H2S) and heated in closed vossels for 10 hours at 100°. Product fractionated (Beckmann, J. pr. [2] 17, 440). Also from potassium amyl-sulphate and K.S (Balard, A. Ch. [3] 12, 303).

DI-iso-amyl disulphide (C.H.,) S. (250°). S.G. 19 .918. From potassium amyl-sulphate and K2S2 (O. Henry, A. Ch. [3] 25, 246; Spring a.

Legros, B. 15, 1938).

Iso-AMYL SULPHITE (C₅11₁₁O)₂SO. (230°-250°). From SOC'₂ or S.Cl₂ and isoamyl alcohol (Carius, A. 106, 291; 111, 97). Oil; decomposed by water or KOHAq into amyl-sulphurous acid and amyl alcohol.

AMYL SULPHOCYANIDE C.11, NS i.e. C.41, S.CN. (197°). S.G. 22 905. Got by distilling potassium amyl-sulphate with potassium sulphocyanide (Heury, A. Ch. [3] 25, 248;

Medlock, A. 69, 214). Di-iso-AMYL SULPHONE [31°]. (295°). Di-iso-amyl sulphoxide (5 pts.) is heated with water (20 pts.) till it melts, a solution of KMnO, (3 pts.) in hot water (30 pts.) is added with constant agitation. The sulphone is extracted with ether. The yield is that indicated by theory (Beckmann, J. pr. [2] 17, 411).

Properties. Long needles, grouped in tufts. Sl. sol. hot water; sol. alcohol, ether, benzene, CHCl, and CS. Soluble in 1LSO, HNO, and acetic acid, but precipitated by water from these solutions. Not reduced by Zn and H.SO, by sodium-amalgam or by HI.

AMYL-SULPHONIC ACID v. Pentane

BULPHONIC ACID.

DI-iso-AMYL-SULPHOXIDE DI-iso-AMYL-SULPHOXIDE (C.H.1). SO. [37°]. From di-amyl sulphide (1 pt.) and fuming HNO, (2 pts.). Crystallised from ether (Saytzeff, A. 139, 854; Beckmann, J. pr. [2] 17, 441). Flexible fatty-looking crystals. Chlorine acts on it in presence of water forming pentanc sulphonic acid, chloro-pentane sulphonic acid, di-isoamyl sulphone, valeric acid, chlorovalerio acid, tri- and tetra-chloro-pentanes, &c. (Spring a. Wiussinger, Bl. [2] 41, 307).

AMYL - SULPHURIC ACID v.

SULPHATE.

Iso-AMYL TELLURIDE $Tc(C_3H_{11})_2$. (0.198°). Got, in impure state, by distilling calcium amyl sulphate with TeK2 (Wöhler a. Dean, A. 97, 1).

Iso - AMYL - DI - THIO - CARBAMIC ACID C.H., NS. i.e. C.H., NH.CS.SH. Iso amy lamine salt C.H., NH. HA'. From isosmylamino and . CS₂ in ethereal solution (Hofmann, J. 1859, 379). Laminæ.

AMYL THIOCARBIMIDES CoH, NS i.e. C.H., N.CS. Amyl mustard oils. Mol. w. 129.

Iso-amyl-thio-carbimide (183°). S.G. 17 .942. Obtained by boiling the preceding compound with aqueous HgCl, (Hofmann, B. 1, 173; Buff, B. 1, 206).

Tert-amyl thio-carbimide EtCMe, N.CS. (166°). From EtCMe2NH2 by successive treatment with CS2 and HgCl2 (Rudneff, Bl. [2] 33, 300).

AMYL THIO-PHOSPHATES.

Iso-amyl thio-phosphate (C.H.1)H.PSO. From isoamyl alcohol and PSCl. (Chevrier, Z. 1869, 413).

Tri-iso-amyl thiophosphate (C,H₁₁)₃PSO₃, S.G. 12 85. From C₃H₁₁ONa and PSCl₃ (C.). Oil. Tri-iso-amyl thiophosphate Di-iso-amyl di-thio-phosphate

(C₅H₁₁)₂HPS₂O₂. Salt. -PbA'₂ [70°] Tri-iso amyl-tetra-thio-phosphate

(C,H1),PS. Formed, together with the preceding, when P.S. acts on isoamyl alcohol (Kowalewsky, A. 119, 310).

Iso-AMYL THIOSULPHATE.

Iso-AMYL The salt Na(C₃H₁₁)S₂O₃2aq is formed by acting with isoamyl iodido on sodium thiosulphate. It

crystallises in laminæ (Spring a. Legros, B. 15, 1938).

Iso-AMYL THIO-UREA C.H., N.S C.H., NH.CS.NH. Monoclinic crystals (Arzruni, P. 152, 284).

AMYL-TOLUENE C12H18 i.e. Cl13.C6114.C3H11. Methyl-amyl-benzene

o-Iso-amyl-toluene (?). (201°). S.G. 2 895. From tolucne, isoamyl chloride, and zinc dust (Pabst, B. 9, 503).

m-Iso-amyl-toluone. (208°). S.G. 22 868. From toluene, isoamyl chloride, and AlCl_s (Essner a. Gossin, Bl. [2] 42, 213). KMnO, gives isophthalic acid.

p-Iso-amyl-tolueno. (213°). S.G. 2 ·864. From p-brome-toluene, isoamyl bromide, and Na (Bigot a. Fittig, A. 141, 160). CrO, produces terephthalic acid.

AMYLUM v. Starch.

AMYL-UREA C6H11N2O i.e. C5H11NH.CO.N1H2-Iso-amyl-urea [91°]. From amyl eyanato and hot alcoholic NH3 (Custer, B. 12, 1330; cf. Wurtz, C. R. 32, 417; Bl. [2].7, 141). Crystals; sl. sol. water.

Iso-Hexoyl-derivative C₂H₁₁NH.CO.NH.CO.C₂H₁₁. [94°]. From the amide of isohexoic acid, Pr.CH₂.CH₂.CO₂H₁, by means of Br and NaOII (Hofmann, B. 15, 758).

Tert-amyl-nrea. [151°]. S. 1.26 at 27°. From tert-amyl cyanate and NH, (Wurtz, A. 139, 328).

n-Hexoyl derivative. [97°]. Formed by action of potash on a mixture of n-hexamide and bromine (II.). Plates; sol. alcohol, and other, insol. water.

Di-iso-amyl-urea C₃ll₁₁NH.CO.NHC₃H₁₁. [39°]. (270°). Formed by boiling isoamyl cyanate with isoamyiamine and alcohol (C.). Needles; insol. water, sol. alcohol and ether.

Di-tert-amyl-urea. Formed by action of KO11 upon tert-amyl cyanato (W.). Needles:

may be sublimed.
Tri-iso-amyl-ursa (C,H₁₁)₂N.CO.N11.C,H₁₁. (260°). From isoamyl cyanate and di-isoamylamine (C.). Liquid.

Tetra-isb-amyl-urea (CsH11)2N.CO.N(CsH11)2 (241°). Obtained by the action of Cl.CO.Et upon a mixture of di- and tri- isoamylamine (C.).

AMYL URETHANE v. AMYL-CARDAMIC ETHEB. AMYL-VALERONE v. BUTYL ENNYL RETONE.

Iso-AMYL-XYLENE C13H20 i.e. C₄H₄Me₂.C₅H₁₁. Di - methyl - isoamyl - benzene. (233°). S.G. 2 · 895. From bromo-xylene, isoamyl bromide and Na (Fittig a. Bigot, d. 141, AMYRIN. A crystalline resin, difficultly soluble in alcohol, contained in some specimens of elemi, and in arbol-a-broa resin (Buri, Neues Repert. für Pharm. 25, 193; Hesse, A. 192, 179). According to Hesse its formula is $C_{it}H_{rs}(OAe)_2$. Bromine forms a complicated bromo-derivative.

ANACARDIC ACID C₂₂H₂₂O₃. [26°]. Occurs in the fruit of Anacardium occidentale (Staedeler, A. 63, 187). Crystals; insol. water, v. sol. alcohol and other. Salts.—CaA"aq.—BaA".—PbA".—HA"PbOAc.—A"FeOH aq.—AgHA".

ANALYSIS. To analyse a thing means to resolve it (ἀναλύειν) into its components. This term, however, has a very wide meaning, which stretches far beyond the outermost limits of our resources of even virtual analysis. So well is this understood by all that even when we speak of a complete analysis we refer only to as complete a solution as the science affords of one or other of three special problems. One of theso is the actual or virtual resolution of the body into its component chemical species or perhaps genera; another, the determination of its elements; the third, the determination of what, in the sense of some imagined general mode of decomposition, are its primary radicles. This (the last named) problem has received a partial solution in the sense that we have ready-made methods for the determination of the acids and bases that may be contained in a solution of salts of a certain low order of complexity. These methods include only a minority of the nonmetallic salt radicles, but they include all the better-known clements as such; and as we have general methods for converting any kind of substance into salts of low order of complexity, these latter methods, conjointly with the former, constitute a complete solution of the problem of ultimate analysis. In regard to the first problem, our powers are very limited. That we have methods for the proximate analysis of certain classes of substances need not be specially affirmed; without these, vegetable and animal ohemistry could have no existence—but a general exposition of their principles would resolve itself into the retailing of commonplaces. We prefer to give a brief summary of what we have of means and ways for seeing whether a substance presumed to be pure really is one substance or is a mixture. In a sense there is only one method: we subject the substance to some physical or chemical process of fractionation, which, while sure not to produce transmutations, gives the several proximate components a chance of parting from one another; and we then compare the several products with one another and with the The form which the original substance. mothod assumes depends largely on the state of aggregation of the substance under operation.

I. Gases. The onenoss of a gas can in general be proved by (a) fractional diffusion through a septum of gypsum or graphite; if the gas is a mixture of, say, two species, the lighter one diffuses out faster than the other; with mixtures of gases of the same specific gravity, the method, of oourse, breaks down: (b) partial absorption. This method is discussed fully under gas-analysis' (q. v.).

II. Solids. These may be susceptible of fractionation by (a) partial fusion; (b) partial solution in suitable solvents; (c) partial freezing of the liquefied body; (d) partial crystallisation out of solutions; (e) partial volatilisation. (See III.) For the comparison of the several fractions, the determination of the fusing points comes in as a handy, and in general sensitive, test

III. Liquids. For these the methods given under (b) and (c) for solids may be available. In the case of distillable liquids we generally resort to fractional distillation, taking care to observe the temperature of the (saturated) vapour, during the progress of the operation. A mixture may have a constant boiling-point, and may besides remain undecomposed on distillation; as a rule, however, it is not so. The volatility of each component depends chiefly on the value for it of the product mp, where m is the molecular weight (or vapour density), and p the vapour-pressure at the prevailing temperature of chullition. For two components, the respective products $m_i p_i$ and m_2p_2 have in general different values. Hence it is not necessarily the lowest boiling component which comes over first; because a large m may make up for a small p. As a mere test for purity, the determination (at a series of suitable temperatures) of the vapour-pressure by the statical method goes considerably further than the determination of the boiling-point curve. In a pure substance, the pressure, p, at to is a function of t only; in a mixture of (say) two liquids, p depends (in a given trial) on the volume of vapour produced, because the ratio of the weight of the vapour to that of the unvolatilised residue changes with this volume. If this ratio is very small, we have an approximation to the vapour-pressure of the more volatile component; if the ratio is large, the pressure approaches the value characteristic of the mixture as such. Any of the many mixtures of constant boiling-point, when subjected to this test, at a suitable temperature, is sure to reveal its complexity. Unfortunately the operations involved are somewhat troublesome, and the results are liable to be largely vitiated by the presence of absorbed air in the sample.

The second and third of our three general problems, qualitatively considered, form the body of what is customarily being taught as

QUALITATIVE ANALYSIS.

The resources of qualitative analysis—apart from mere methods of identification of named species, which we leave on one side—may be arranged under three heads:—

I. Flame Teets. A set of methods for the detection of elements as such, which, being all founded upon ultimate or penultimate dissociations at high temperatures, are in a high degree independent of the constitution of the substance operated upon. Another specific feature in these tests is that they are easy and rapid of execution, and demand only very small quantities of substance.

II. A set of what we will call methods of chemical disintegration (each general in reference to a large class of bodies), by means of which compounds of high chemical complexity oan be, so to say, opened up, and their elements brought within the range of our routine methods of sait analysis (n. imfra). These methods might, each, be indexed, by reference to a certain element, generally of high valency; as a rule it is a non-metallic element, and in the analyst's sense, sui generis, i.e. not susceptible of detection

as one of a group.

III. The systematic methods for the radical analysis of a solution of salts which were referred to in the introduction. In addition to these analytical methods, the analyst naturally discounts all that may help him towards the solntion of his problem. Our three classes of methods alone, it is true, if judiciously employed, would enable one in general to perform an exhaustive ultimate analysis; but a purely ultimate analysie, in the majority of cases, is not what we want. The ideal which we aim at in all analyses made for practical purposes is (as good an apology as is attainable for) a proximate analysis; a recipe, so to say, for composing the substance from things of known constitution. Hence, if the substance presents the aspect of a mixture—mechanical or physical—we naturally begin by trying to effect a separation of the several things from one another: by mero picking out, or eintriation, in the case of an obvious mixture; by distillation in the case of a solution in a volatile solvent. We will assume, however, that this division of the given substance into two or more substances has been effected (if called for, and possible), and that the substance to be analysed is a solid (which virtually includes the case of a liquid); gases demand special methods, which lie beyond our programme. In this, as in any analogous oase, we naturally begin by a close observation of at least the general properties of the body; it may be expedient to supplement this observation by the exact determination of certain physical properties, such as hardness, specific gravity, crystalline form, optical constants, &c., &c. Everything here depends on the nature of the case, and -of the operator. A mineralogist, for instance, by such determinations, may be able virtually to analyso a mineral; but everybody cannot do this. After the more purely observational stage, it is cxpedient to study the behaviour, at a graduated succession of high temperatures, of (a) the substance itself, (b) the substance and atmospheric oxygen, (c) the substance plus added reagents. We therefore begin by heating a few centigrams of substance in a sublimation tube over a Bunsen lamp-first gently, then more and more strongly -on the chance of obtaining a readily recognisable sublimate, gas or vapour, or residue. Of the identifiable residues, charcoal is the most important, because its formation proves the presence of organic matter in the sample, although all organio matter does not yield charcoal. If 378 substance presents a metallic, or semi-

Alic, aspect, it is expedient to roast a frag-CH, N. a draught-tube, on the chance of Lso-an white arsenic, iodine, &c., or an tained by sulphurous acid, &c. If the subleous Hgs partially volatile or partially 1000. we prepare a supply of the fire-proof the fire proof the fire the fire the fire-proof of the fire-p I. Flame Toris.

By "flame-tests" we mean dry way tests, in which the substance, or substance plus reagents, is heated directly in the flame. These tests were introduced by Gahn, and subsequently extended and brought to high perfection chiefly by Berzelius and Plattner. In accordance with the modes of operating which these authorities found it convenient to adopt, a blowpipe flame used to be universally employed as a heating medium. But Bunsen, some twenty years ago, showed that most blowpipe tests can be done more easily and conveniently in the flame of the gas lamp which bears his name, and that this flame can also be employed for certain new teste introduced by him, which tests could not be conveniently done with the blowpipe. Many chemists prefer the Bunsonian modus operandi, but the blowpipe has not by any means become obsolete; it will continue to be used, because it offers certain specific advantages of its own. For the purposes of this article it will be sufficient to give the following enumeration of the more important of the characteristic flametest operations:

(1.) A few mgms. of the substance are placed on an asbestos-stick, and exposed to the several regions of a Bunsen-flame, proceeding from lower to higher temperatures, to determine degrees of fusibility and volatility.

(2.) A few nigms., fixed to one end of a hairfino platinum wire, are exposed first to colder, and then to the hottest, part of the flame-mantlo of a Bunsen, in order to see whether the flame is thereby coloured. If a colour is produced, it is analysed optically (by means of a spectroscope), with the view of thus effecting a chemical analysis of the glowing vapour. Of elements identifiable by their spectra (or flame colours), the following may be named:—TI, II, Rb, Cs, K, Na, Li, Ba, Sr, Ca in most of the ordinary states of combination, Cu as haloid salts, B as boric acid or fluoride, P as free phosphoric acid or PH₂.

Spectrum analysis, as everybody knows, was invented by Bunsen and Kirchoff. Somo years ago Bunsen brought it into a new form, in which a spark current, produced by means of an induction-coil is used for the volatilisation of the substance; at the very high temperatures thus produced, a great many elements, besides those named, become identifiable by their spectra.

Flame Tests with Reagents.—Of these sodium carbonate is most extensively employed; either platinum wire or charcoal being used as

a support.

On platinum; (1) as a mere flux, it identifies SiO₂, given as such or as highly acid silicate; (2) in conjunction with oxygen used as air, or introduced as nitre—it detects Cr and Mn with certainty, converting the former into (yellow) chromate, the latter into (green) manganate.

When nsed on oharcoal; in the reducing flame, it may bring to light, (1) S, Se, Te. Any non-volatile form of these yields a fused mass, containing Na₂S &c., recognisable by the black stain (Ag₂S &c.) which it produces when placed on a silver coin, and moistened with a drop of

water.—(2) One or more of the following metals; As, Hg, Zn, Cd, Pb, Bi, Sn, Cu, Ag, Au, Fe, Ni, Co, Pt. The oxides and many of the salts of these metals when subjected to the operation under discussion, are reduced to the elementary state. The metal thus liberated may assume the form of a visible fused bead, or remain concealed in the form of fused scales or an unfused powder or sponge; these, however, can in all cases be isolated and brought to light by elutriation with water in an agate mortar. Part of the metal, in general, volatilises, and in passing through the flame becomes oxide. If the reduction is effected in the blowpipe flame on a block of charcoal (old style), part of the oxido in general settles down on the charcoal as a ring, and by its colour may aid in identifying the metal. When operated upon as described, compounds of As yield only vapour of oxide, which in most cases is lost altogether. Hg; vapour of metal, which is also lost. Zn, Cd; little or no metal, but abundant oxide rings; (ZnO is white, CdO brown). Pb, Bi; easily fusible metal, and tangible quantities of yollow oxide. Sn; easily fusible metal, and little (white) oxide. Cu; not easily-Ag; more easily —Au; not easily—fusible metal or scales, and no oxide. Fe, Ni, Co; unfused powdery, or spongy, metal, which follows the magnet. No oxide. Pt; like Fe, but the metal is not magnetie and is unacted on by HNO, Aq.

In Bunsen's mode of operating—which consists in heating the mixture of substance and soda on a slender stick of charcoal in the reducing part of the 'zone of fusion,' the oxide is lost, but all the respective metals fall within the

range of

Bunsen's Film Tests .- When the airholes of the Bunsen are partially closed, a luminous tip forms somewhere near the apex of the flame. Many oxides suffer reduction when held in the centre of this tip on an asbestos stick; and the reduced elementary substance can be collected on a Berlin basin (filled with water to keep it cold), held over the sample across the flame. The elements thus appear as films resembling the stains of As and Sb produced in Marsh's test. The following elements chiefly yield films: As, Sb, Te, Se; hardly attacked by nitric acid of 20 p.c. Bi, Hg, Tl; vory slowly dissolved by nitric acid of 20 p.c. Pb, Cd, Zn, In: instantly dissolved by nitric acid of 20 p.c. By a very obvious modification of the process, oxide films can be produced in lieu of inetallic ones; but we cannot go any further into this

Borax is always used as a bead fused to the end of a platinum wire. Such a bead dissolves most metallic oxides at a moderately high temporature, forming glasses, the colours of many of which are characteristic of the metal. Often one metal gives two colours according to whether the fusion is effected in the oxidising or in the reducing flame; this affords additional means of discrimination.

Microcosmic salt (or rather the fused Na.O.P.2O. produced by its decomposition by heat) acts on metallic oxides pretty much as borax does; but its specific function is the detection of silica. If a splinter of a silicate is treated in a fused meta-phosphate bead, the

bases dissolve out, the silice remains in the characteristic form of an unfused 'skeleton' of the splinter.

Whatever the flame-tests may have brought out by way of positive results, their negative results count for very little.

II. Methods of Chemical Disintegration.

Substances may be divided into two classes. as regards the operation's required to bring them within the range of our systematic methods of salt-analysis. (1) Such as are simple salts (we mean salts which can be analysed by our routine methods), or can be made into solutions of such by the application of the ordinary mineral solvents, such as water, dilute mineral acids (qua acids), nitric acid or aqua regia (qua oxidants). This class comprises many minerals, and ordinary chemical bodies, but unfortunately (and naturally) we have no general test for the recognition of these bodies as a class. (2) Such as demand special methods of disintegration. Of the more commonly occurring chemical genera, the following may be named as falling within this class:—(a) Fluorides; these although perhaps of the simplest constitution, demand special methods because hydrofluoric acid and all acid fluoride-solutions attack glass and porcelain. (b) Most silicates: silico-fluorides. Cyanides, especially metallocyanides. (d) Salts of certain complex organic acids (not oyanides); in the sense that they exhibit abnormal metalreactions. (e) Organic compounds generally; in the sense of ultimate analysis generally. An-orthophosphates. (g) Certain classes of sulphur compounds.

This list does not pretend to be complete, but it includes most bodies which the practical analyst is likely to come across. For the second class of substances as a class, we of course have not a general test any more than we have for the first, but we have general tests for the ssveral genera, in this sonso at least that we have general methods for the detection of their cha-

racteristic elements.

The following section is compiled partly with the view of supplying the necessary information in this direction.

General methods for the detection of certain elements (mostly non-metals) and for the ultimate analysis of their compounds.

Silicon is always isolated in its highly characteristic form of silica, SiO2, which is easily identified by the blowpipe tests given above, and by its convertibility into volatile SiF, by the action of HF. Silicon and metallic silicides, when fused with caustic alkali, yield alkaline silicates (q. v.). Alkaline silicates (even if so acid as R₂O.4SiO₂) dissolve in water, forming alkaline solutions. Mineral silicates, Slags, Glasses, &c. fall within two classes according to whether they are, or are not, decomposable by hydrochloric acid. Those of the first class are finely powdered and digested in conc. hot HClAq until disintegrated, evaporated to complete dryness (to convert the colloidal part of the silica into the insoluble form), drenched with HClAq, allowed to stand (to re-ohlorinate the Al,O, and Fe,O

produced), treated with water, and filtered. The silica remains on the filter; the solution contains the metals as chlorides. Of those of the second class, some are disintegrable by hot semi-conc. H2SO4Aq (ex. the clays). The general mothod is to fuse the finely powdered silicate with KNaCO, nntil all is dissolved, and to analyse the fused rosiduc as a silicate of the first class. Alkalis must be tested for in another portion of the silicate, after evaporation with NH,FAq, whereby Si is removed as SiF, and the bases remain as fluorides easily convertible into sulphates by H.SO, Aq (comp. Fluorino).
Aluminium.—Only the forms of Al₂O₃ in-

soluble in acids need be considered here; these if finely enough divided, all dissolve at a redheat in fused KOH, becoming aluminates soluble

Chromium.-All non-volatile compounds, when fused (in silver) with KOH and KNO, yield alkaline chromate, recognisable by its yellow colour and the very intensely yellow colour of its aqueous solution. This operation constitutes a general method of disintegration for the forms of Cr.O. and chromites insoluble in acids; it goes a certain way even with chrome iron orc, but the complete disintegration of this mineral demands special methods.

Titanium. - TiO, stands between SiO, and Al₂O₃. Unlike the former it is not volatilised by evaporation with HFAq. Titanates are de-composed by fusion with KHSO₄; the cold aqueous extract after fusion includes the TiO2 which is precipitated on boiling, as such.

Tin.—The forms of SnO, (including tinstone) which are insolublo in acids yield Sn when fused on charcoal with NaHCO, and KCN. They may be disintegrated (1) by fusion with KOH; the SnO₂ becomes stannate soluble in water: (2) by fusion with six pts. S and six pts. Na₂CO₃; the aqueous extract after fusion contains the Sn (also any As and W that may be present) as thiosalt, and consequently falls in with a certain stage of the routine method of metal-analysis (v. infra).

Carbon in any state of combination is convertible into CO., which is readily identified. It is distinguished from HCl and SO. by its scanty solubility in water, and inertness towards oxidising agents; from N, H, &c. by its abundant solubility in solutions of basic hydrates; with CaO₂H₂Aq and BaO₂H₂Aq it gives a characteristic white pp. of carbonate. Carbonates (almost without exception) are decomposed by mineral soids with evolution of CO2. Elementary carbon (in all forms) barns in oxygen to CO2.

Combustible Carbon Compounds .- The methods of organic analysis (q. v.) are easily translated into general methods for the detection of combustible carbon as CO2. It is necessary to purify the CuO or PhCrO, immediately before nse by heating it to redness in air until it ceases

to give off CO...

 All non-volatile carbon compounds can be burnt by heating them with conc. H2SO,Aq and CrO. Many volatile organic bodies unite readily with conc. H.SO,Aq to form non-volatile compounds, and thus fall within the range of the method which obviously suggests itself for the detection of combustible carbon heside carbonate.

Analysis of Carbon Compounds.-I. Organic acids proper (COOH compounds) need here be considered only in regard to the extent to which they interfere with the routine methods for the detection of the metals in a solution of salts. Some (including formic, acctie, succinic, and many others) interfere only in this sense that, in their prosence, the preci-pitate obtained by H.S in the presence of free acid, may include Zn, Co, Ni, and perhaps other metals of the iron group. This difficulty is easily overcome. A large class of non-volatile acids, including the ordinary fruit acids, prevent the precipitation of Fe₂O₃, Al₂O₃, Cr₂O₃, CuO₄ and other metallic oxides by alkalis, and that of Al and Cr even by sulphide of aminonium. In all difficult or doubtful cases, it is best to destroy the organic part of the salt, which can be done in two ways :- (1) By incineration : which, of course, had better be postpoued until after the elimination of the copper and arsenie groups by sulphuretted hydrogen; if this has been effected, Zn, of all the metals left, is the only one which may be lost by volatilisation .-(2) By treatment (of the dry salts) with oil of vitriol. The ultimate product contains the metals in the form of sulphates.

II. Cyanides. a. Hydrocyanic acid; easily recognised by its volatility and specific smell and reactions; regarding the latter, see b. b. The simple cyanides of the mero positive metals (K to Ca inclusive). These are all soluble in water. The solutions are alkaline, and give off HCN with acids. AgNO₃Aq in excess precipitates AgNC, insoluble in dilute HNO₃Aq. When mixed with (1) excess of alkali, (2) ferrosoferric salt, (3) excess of HClAq, they yield a blue precipitate (or green suspension) of Prussian blue. c. Cyanide of moreury, Hg(NC)₂. Soluble in water. Exhibits anomalous reactions both as a mercuric salt and as a cyanide. But is decomposed by H.S into a pp. of HgS and a solntion of HCN. d. Heavy metallic cyanides, and metallocyanides. Some give off part of their cyanogen as HNC, when distilled with dilute HClAq or H₂SO₄Aq. Many (e.g. prussiates) recognisable by specific tests. Solutions of metallocyanides mostly give pps. with AgNO, Aq, insoluble in dilute HNO, Aq, in which the charac. teristic metals of the radicles can be detected (v. Halogens).

A general method for the detection of the metals in cyanides, cyanates, and thiocyanates, is to heat the dry substance with conc. H.SO.Aq until completely decomposed. The cyanogen becomes ammonia-salt, and CO; the metals remain as sulphates. About the detection of non-metallie elements in carbon compounds, see

sect. on S, P, &c.

Boron occurs chiofly in the form of borate. The presence of boric acid does not interfere with the rontine methods of metal analysis.

Phosphorus is always isolated and identified as orthophosphate. I. Orthophosphates, as far as not soluble in water, are mostly soluble in HClAq. To search for phosphoric acid, we supersaturate the solntion strongly with ammonia, and (after filtration, if necessary) add magnesia mixture (NH,Cl and MgCl, in NH,Aq), crystalline PO,MgNH, 6H,O gradually forms. insoluble in dilnte NH.Aq. A pp. formed by NH.Aq generally contains part, sometimes the whole, of the phosphoric acid. To detect the latter we dissolve the pp. in HNO.Aq, add excess of a nitrio solution of molybdate of ammonia, and allow to stand at 40°C. All the phosphorio acid comes down gradually as a yellow powdery pp. of phospho-molybdate of ammonia, insoluble in excess of reagent, but solublo in excess of aoid phosphate; soluble in ammonia. Both reactions are very delicate, and, in the absence of arsenio acid (which in the circumstances behaves like phosphoric), highly characteristic. Phosphates in any other state of combination can be brought into the orthophosphate form by suitable operations. II. Meta- and pyrophosphates (which besides being different in their own reactions from orthophosphates, oxhibit anomalous metal-reactions); by long-continued boiling with mineral acids, or (what is better) fusion with carbonate of alkali. III. Elementary phosphorus, and all oxidisable phosphorus compounds; hy treatment with HNO, Aq of the proper strength at the proper temperature. Many organic phosphorus compounds, it is true, cannot be thus completely oxidised, but in their case, we need only neutralise the nitric liquor produced with potash, evaporate to dryness, and fuse the residue with KOH, to convert all the phosphorus into orthophosphate.

Sulphur.-Analytically speaking, sulphuric acid is to sulphur what orthophosphoric acid is to phosphorus. I. Sulphates, in an aqueous or IICl solution, are separated out completely by BaCl.Aq, as white, powdery, BaSO, insoluble in aqueous mineral acids, and thus distinguished from all the haryta-pps., except the selenate and fluosilicate. BaSeO₄ is decomposed by boiling HClAq with formation of Cl and SeO, while BaSO, is not so decomposed. The fluosilicate yields no sulphide on fusion with Na₂CO₂ on charcoal; and dissolved fluosilicates give no pp. with SrCl.Aq, while sulphates yield a pp. of SrSO, slightly soluble in diluto acids. II. Acidinsoluble sulphates are disintegrated by fusion with alkaline carbonate, and treatment with water: a solution of alkaline sulphate, and a residue of the respective carbonato, oxide, or metal, are obtained. III. Metallic sulphides .-Many are decomposed by HClAq with evolution of H.S. IV. The salts of the lower sulphur acids, when heated (in solution) with alkaline permanganate are completely exidised with ppn. of manganite, MnO2.R2O. The excess of oxidant used is brought into the same form by addition of a few drops of alcohol. The filtrate contains all the sulphur as snlphate. Only dithionio acid does not yield roadily to this process of oxidation. All the sulphur compounds III. and IV., including dithionates, and many organic sulphur compounds, are oxidised completely by hot, sufficiently cono. HNO₂Aq. Volatile som-pounds (such as CS₂) must be manipulated in a sealed glass tube. From some organic bodies only sulphonio acids are produced; but these, when fused with KOH and KNO, all yield up their sulphur as sulphate.

All oxidisable sulphur compounds are completely oxidised to sulphates by the action of basic reagents (like Na₂CO₃, CaO, &c.), and KNO₃, or even oxygen-gas, at a red heat. All non-volatile sulphur compounds yield alkaline

sulphide when fused with Na₂CO₂ on charcoal in the reducing flame (v. Flame Tests).

Selenion and Tellurinm are closely allied to sulphur, but must be passed over.

Nitrogen, in all states of combination, is susceptible of elimination as nitrogen gas, recognisable by the methods of gas-analysis. Another less general, yet widely applicable and more convenient, method is based upon the conversion of the element into ammonia.

I. Ammonia; "ecognisable by its smell, its great solubility in water, its ready union with HCl to form solid NH4Cl, &c. The least traces of NII, or NH, salt in water, are detected by Nessler's reagent (a solution of HgI, and KI in KOHAq); iodido of mercurammonium separates from moderately dilute solutions, as a brown pp., and even in the most dilute solutions is visible as a brown or yellow colour. II. Ammonia salts; many amides (including all acid-amides) when distilled with caustic alkali, yield NH, which passes into the distillate. III. Nitrates and nitrites in alkaline solutions are reduced by nascent hydrogen (KOHAq and Al) to NII. IV. Metallic nitrides, and all organic nitrogen compounds not containing their nitrogen in the form of exygenated radicles or in the diazo-form, when burnt with soda lime yield their N as NH₃.

Fluorinc. Most metallio fluorides, when treated (as powders) with cone. ILSO Aq in a platinum cruciblo at a gentle heat, give off HF, recognisable hy its etching glass and even rock-crystal. For the purpose of metal-detection, the mass must be evaporated until a tangible quantity of sulphuric acid has gone off as a heavy vapour. The bases remain as sulphates.

Mixtures of fluorides and silicates, when heated with conc. H₂SO₄Aq givo off SiF₄, decomposed by water into H₂SiF₆Aq and a gelatinous pp. of SiO₂, which, however, may be invisible. To detect the fluorine, add excess of ammonia to bring down all the silicon as silica (which filter off), and evaporate the filtrate in platinum on a water-bath nearly to dryness. Residue is fluoride of anmonium.

Fluosilicates. Those of the most basylous metals when heated dry broak up into SiF, and a residue of fluoride. Fluosilicates generally behave to boiling alkali solutions us if the silicon were a basylous metal.

The Halogans (Cl, Br, I). I. The elementary substances are recognised by their very characteristic properties. When treated with zinc and water, they all dissolve as haloid salts of zino (ZnCl., &o.). II. Haloid salts; mostly dissolve in water or in HNO.Aq. Evon from the latter solution the halogon is completely ppd. as haloid salt of silver, insoluble in dilute mineral acid. III. The oxygen acids of the halogens. (Periodio acid ignored.) Of these only bromio and iodic give silver pps. insoluble (or solublo with difficulty) in cold, dilute, HNO.Aq. All the rest form soluble silver salts. Then alkali and alkalino-earth salts when heated dry give off oxygen and become haloid salts. With the only exception of perchloric acid they are all reduced by SO.Aq to halogon hydride (a.g. HClO., to HCl). Hence an obvious IV. Relatively general method for the detection of halogen in a solution of salts. The solution

(which we will assume to be neutral or sold) is mixed with excess of SO, Aq and AgNO, Aq, the pp. is allowed to form, and then treated with HNO, Aq, to remove foreign ealts (including Ag. 80, which is not very readily dissolved). The pp. contains all the halogen of the solution, (except that of the perchlorie acid); but it may beeides contain-if it does not consist of -cyanide, thio-cyanate, and metallo-eyanatee, of silver, (not to mention the sulphide which is easily kept out). An analysis can be effected by calcining the dry pp. with chemically pure eoda-lime, preferably in a current of moist hydrogen. The nitrogen of the cyanogen radicles goes off as ammonia, which is easily identified. The residual product contains the metals of the metallo-cyanates as oxides, the silver as metal, the sulphur of the eulphocyanogen as alkaline sulphide, and the halogens as alkaline haloids. V. Organic halogen compounds. All these, when burnt with quick-lime in a combustion tube, yield up their halogen as haloid salt of calcium, extractable by cold, dilute, HNO.Ac.

III. Methods for the Systematic Examination of a Solution of Salts for its Metals

can be given only on the basis of restrictive assumptions. We assume, in the outset at least, that the solution is so constituted that it might have been prepared by dissolving n set of basic or acid metallic oxides in nqueous mineral acid or alkali, and that certain rare oxides and certain rare combinations of things are absent. Some of the cases lying beyond this programme are dealt with in appended notes to which reference is made in the context. For the sake of generality, however, we assume that all the more ordinary metallic radicles may be present. It evidently would not do to search for them individually and seriatim; the only course one could reasonably think of is to begin by splitting up the given complex group of nietals into a number of groups, so that each of these shall contain the whole of, and nothing but, certain metals, A, B, C, . . .; to then apply the same principle to the groups; and then to the groups of the second order; and so on until one arrives at last at either the individual metals, or at groups of such emallness that the side by-side recognition of their members offers no difficulty. This, at any rate, is the course which is adopted by every chemist. The table on p. 221 in its first vertical column names the generic reagents which are customarily used for the formation of primary groups, and shows how these act on solutions of the groups of oxides named in the successive column headings. For the separation of the groups from one another it is obviously expedient to begin by eliminating the silver group by means of hydrochlerie acid, which must be added in inetalments until the solution is decidedly acid, and, if a permanent pp. appeare (which with us can consist only of these three chlorides), antil the ppn. ie completed. The pp. containe

In an attains solution of saits ornerally, HOLAq may pro-duce a great variety of permanent pps, other than silver-troup chlorides. For the purpose of a mere metal-analysis

all the silver and merourosum as AgOl and Hg,Cl,, but only part (if any) of the lead; a small quantity of thie metal always passing into the filtrate. From the filtrate the copper and arsenic groups are ppd. conjointly by meane of sulphuretted hydrogen. Before applying this reagent, however, we must make sure of the at least relative absence of nitroue, nitric, and ohlorio, acid and other oxidieing agents, which, while not easily or completely reducible by H2S would at least tend to oxidise it and impede its normal action. Any of the three oxidants named can be expelled by repeated evaporation to a small volume with conc. hydrochloric acid.2 last residue ie diluted with the proper proportion of water, and (heedless of any insoluble oxychloride that may separate ont) treated with eulphuretted hydrogen, first at about 70° to make sure that As₂O₃ ie completely reduced to As₂O₃, and ite metal ppd. (ae Ae₂S₃ + S₂), and then again after cooling, or else part at least of the cadmium and other copper-arsenio group metals, whose sulphidee are rather unstable in opposition to aqueous acide, would escape ppn.

On account of the metale just referred to, we must eee that the quantity of free mineral acid is not excessive, but is sufficient to prevent the ppn. of the zinc, which from only feebly acid solutions is liable to pass into the cul-

phuretted hydrogen pp The ppd. sulphides are collected on a filter and washed with very dilute sulphuretted hydrogen water, to constantly re-sulphurise what may have become sulphate by the action of the air; the first inetalments of wash-water being acidified to the extent of the mother liquor, to prevent ppn. of the zinc. In order now to separate the two groups, the pp. ie digested on a water-bath heat with undiluted yellow sulphide of ammonium; an excess of sulphur in this reagent being necessary, chiefly on account of the stannous sulphide, SuS, which becomes ecluble only through conversion into stannic, SnS2. To effect a complete separation, the treatment with sulphide of ammonium may have to be repeated with the first residue. The copper-group sulphides are filtered off and washed with wnrm water mixed with a little eulphide of ammonium. From the filtrate the arsenic-group sulphides are reproduced by acidification with dilute sulphuric acid; after expulsion of the dissolved culphnretted hydrogen by a gentle heat, they are filtered off, and washed with plain water (sulphuretted hydrogen water would dissolve sulphide of nrsenic Ae, S.). The pp. is liable to be contaminated with eulphide of copper; this can be climinated by treatment with warm dilute cauetic potash, which dissolves the arsenio-group sulphides

a pretty safe rule is this. If a solution on adding HClAq a pretty safe rule is this. If a solution on adding HClAq gives an abnormal-looking pp., repeat the experiment with iNO.Aq; if no permanent pp. is produced, HClAq will act normally as a chloride; if a pp. is formed, it must be filtered off and analysed for the metale that may be in it, (as sulphidea, e.g. As.S.; or otherides such as AgCl, &o., &c.). The solution, as a rule, is now fit for treatment with hydrochioric acid, &c.

In evaporating a solution of metallic oxides with a reappraying a solution of metallic oxides with FICAq, it is as well to remember the volatility of AsCl., SbCl., SbCl., SnCl., BiCl., The evaporation is best conducted in a retort, and these volatile chiquides are searched for in the distillate.

for in the distillate.

If the solution contains Ti, the metal passes for the most part into the pp., where it is easily detected hy spectrum analysis. The characteristic solubility of its chloride in Na₂CO₂Aq enables one to separate it from the ordinary silver-group chlorides.

GENERIC REACTIONS OF MINERAL-ACID SOLUTIONS OF GROUPS OF WETATITIC OVERSES

Magnetium Alkali Group. MgO. Kideli, O.		ation of white	Precipitation of white Man, or Man, or Man, or Man, or Man, or Colourless, phorio, No change, r, acids,		No visible change.	METALS, REMAIN DISSOLVED, even in presence of moderate quanti-	No precipitates.			
		ROp, with precipits My Mi		In the absence of phosphorio, oxalic, and certain other, acids, no visible action.		In presence of NH,Cl, no precipitate.	METALS REMA even in presence o	Slowly but completely precipitated as PO,MgNH ₄ +6H ₂ O.		
Rarium Groun	Barium Group. Bao, 8ro, Cao.		METALS REMAIN DISSOLVED. reduced to lower oxides Ro, Gro, Cro, Cro, Red to green or violet.		In the absence oxalic, and cert no visibl		Metals precipitated as R"CO, inso:nble in NH,Cl.	Ea, Sr precipitated as RSO,: no Ca dil. sol.: no pp.; on addition of alcohol, GaSO, completely precipitated.		
4	R ₂ O ₂	A10, Cr.0.	Metals remain dissolved.	Oxides RO, are reduc	rgely diluted acid.* Precipitate soluble in soluble	Cr and Al are precipitated as hydrates; Fe as Fe ₂ S ₂ + S.	Fe ₂ O ₃ , Al ₂ O ₃ , Cr ₂ O ₃ precipitated as hydrates,		ol ; excepting SnO	monium sosphate, Many of these oxides are precipitated, more or less completely, as phosphates, Prophetely Precipitated as ProMRNH, +6H_0.
	B.'0.	FcO, MnO, Zno, NiO, CoO.	METALS	The following rulphur, and cha		insoluble in	Mn, Zn, Ni, and Co, no precipitate.			
Arsenic Group.		A3,0, 8b,0, 8n0,		insoluble in		niam sulphide.† senic Group, no pr rest precipitated s excess of precip	The rest precipitated as sulphides natural excess of precipitant. Metals behave differently.			
Copper Group.		Hgo, Cuo, Biso,, Cdo[Pbo].		tated as sulphides, rgely diluted acid.*					even in pres	ny of these oxides a
Silver Group.		Ag.O. Eg.O.	Metals precipitated as chlorides.	Metals precipi la	insoluble in an	Ar The insoluble in	Metal		Hg.0 and PbO precipitated as sulphates.	Ms
	Reagents.		Hydrochloric acid.	Sulphuretted	hydrogen and . free acid.	Ammonium sulphide in neutral or alkaline solutions.	Sal-ammoniac and excess of ammonia.	Ammonium carbonate.	Dilute sulphuric acid.	Ammonium phosphate, NH,Cl, and NH,

only. From the filtrate, these sulphides can be recovered by acidification, in their original form. After elimination of the cepper and arsenic groups, the barium-group may be separated out by means of sulphuric acid. The barium comes down at once (as BaSO4), the strontium gradually, on standing and keeping warm. Frem the filtrate frem these two sulphates the calcium can be ppd., after due concentration, by judicious addition of alcehol, and allowing to stand for, say, 12 hours. The calcium sulphate is filtered off and washed, first with dilute, and lastly with strong, alceliol. The filtrate, after removal of the alcohol, is ready for the elimination of the iron-group, &c. This method is the best that can be adopted if an analysis for the barium-group metals is our principal object; it also offers certain ether specific advantages; yet the majority of chemists prefer (after application of sulphuretted hydregen in the presence of acid) at once to separate ont the iron-group, by means of sulphide of ammoninm. The addition of this precipitant must of course be preceded by the neutralisation of the free mineral soid of the solution with ammonia; if a sufficiency of sal-ammoniac is not thus produced incidentally, some sal-ammoniae must be added, to bring the pp. into a fit condition for filtration. But we have ne space for these technicalities, and accordingly assume the pp. to have been filtered off and washed with warm water mixed with some sulphide of ammenium, se as to remove the whele of the mother-liquor. This liquor, by theory, contains the whole, in practice it may be assumed in general to contain the bulk, of the barium-group metals and of the magnesium, in addition to the whole of the alkalis. For its analysis, the barium-group is ppd. by means of carbonate of ammonia added to a warm solution. In the presence of ammonia-salts, of which as a rule there is more than enough, only the barium-group metale are ppd. as carbonates; the pp. is collected on a filter, and washed with het water. Part of the filtrate serves for the detection of magnesia by means of ammonium phesphate. The rest of the filtrate is evaporated to dryness, and the residue calcined. The ammenium salts volatilise, or at least their ammonia does, and there remains a residue containing only magnesium and alkalimetals, which latter can be detected without elimination of the magnesium by euitable methods.

In regard to the analysis of the groups, our space decs not permit us to do mere than shortly indicate how the sulphide of ammonium pp. (which may be of very complex composition), can be split up into miner groups. Before deing so, let us state that in the presence of representatives of a certain group of acids which includes H,PO,HF,H,BO, and certain organic acids, e.g. oxalic, the pp. is liable to contain part, or all that there is, of barium-group metals and of magnesium, as salts of the acids named. A thoughtful analyst takes care to keep these inconvenient acids out of the solution; but the introduction of phespheric acid is eften unavoidable, and wo therefore assume it to be present. Whether this acid is present or not, the cobalt and nickel can be eliminated, approximately at least, by treatment of the pp. with cold, dilute ' HClAq, and removed by filtration. The filtrate is next tested for iron, best by adding a few granules of chlorate of potash and boiling, when the iron assumes the form of ferric salt, and becomes visible by the intense yellow colour of its hot hydrochlorio eclution, and at the eame time assumes the right form for the next step, which aims at a separation of the metale present as Fo₂Cl₆, Al₂Cl₆, Cr₂Cl₆, and the phesphates, from the metals (manganese, zino, &c.) present as dichlorides. Of the various methods which we have for their separation, the most convenient for general purposes is the following:-

After having made sure of the complete reduction of the manganese (Mn,Cl,) to manganeus chleride by euflicient boiling with hydrochlorie acid, we allow to coel, dilute pretty considerably, and next add (sal-ammeniae if necessary, and) ammenia, drep by drop, until the mixture is alkaline. We then (without lesing time and giving the oxide of manganese much chance to get per-exidised), boil until the vapours cease to smell of ammonia, and filter. The pp. contains all the iren, aluminium, and chromium, and all the phospherio acid as lime-salt, or in other forms; the filtrate centains at least part of the zino, manganese, and in general part of the rest of the protexides. If the sesquiexide-pp. is bulky, it must be redissolved (after a few washings) and re-produced by a repetition of the first operation. From the pretexide filtrate, the zinc, after acidification with acctio acid, can be ppd. pure by fractional ppn. with sulphuretted hydrogen-water in the cold. The manganeso, traces of nickel and cebalt, and in general much lime, baryta, and strontia, remain disselved.

The scsquiexides-pp.2 (if chromium be pre-

* The cobalt-nickel pp. never contains the whole of these ** The coord micret pp. never contains the whole of these metals; part passes into sointion, and ultimately finds its way into the 'protoxide filtrate.' In addition to its normal components it is liable to contain sniphide of zinc, and perhaps traces of other Iron-group metals, and any cadmium, antimony, &c., that may have been allowed to slip into the filtrate from the sulphuretted hydrogen pp.

If the solution contains urunium, this, in our scheme

If the solution contains uranium, this, in our scheme of analysis, goes with the iron, and consequently has to be looked for in the sequioxide pp.; from which it can he extracted hy digestion with warm, concentrated, solution of carbonate of ammenia. To pass now to a number of rare metals, which we have so far entirely ignored:

Palladium, in our system, holongs to the copper-group. It is characterised chiefly by the utter insolubility and black colour of its ichief, and the arsenic group: only the

black colour of its leddle.

Platinum and gold go into the arsenic group; only the sulphides are not easily soluble in alkaline sulphides. In almost all practical cases they can be kept outside the solution intended for the detection of the metals by snit-able methods. If they are unavoidably present, they ere best separated out; the gold by ferrons chierdle (as metal); the platinum, by means of solid sal-ammoniae added to the concentrated solution, as PtCl, Nil, , which must be washed with the least possible quantity of a solution of the precipitant.

Titanium, as TiO₂₇ in the analyst's sense stands hetween SiO₂ and Al₂O₂. In our system it goes with the Al₂O₃.

Al₂O₃.

Beryllium behaves to our group-reagents like Al₂O₃, but it is far more easily soluble in sal-ammoniae than alumina is. Unlike it, it dissolves in carbonate of ammonia, and

is. Unlike it, it dissolves in carbonate of ammonia, and does not form an alum.

The rare earth metals, cerium, lanthanum, &co, &co, must all be passed over here.

Lithium! (easily detected by spectrum analysis) behaves on the whole like K and Na, but unlike them forms an insoluble phesphate produdile by ovaporating its solution with phosphate of soda pins caustic soda (i.e. with Na, PO), to dryness, and treating the residen with water, when it remains. From magnesia (if ammonia salts are absent) is

sent) is best analysed by fusion with caustic | potash and nitre in a silver dish, and treatment of the fused mass with water. Should the solution be green from manganate, this must be reduced (to MnO₂) by addition of a few drops of alcohol and heating. The mixture is then filtered. The filtrate contains the chromium as (yellow) chromate, the aluminium as aluminate, and part, in general, of the phosphoric acid as phosphate. The residue contains oxide of iron, magnesia (MgO), and possibly barium-group

metals as phosphates.

The Determination of the Non-Mctallic Components.—Our systematio mothods for the detection of the metals contained in a solution of salts are far less hampercd by oncrous conditions than are most of our methods of acid detection. Hence the general rule to first complete the analysis for metals before attempting the systematic and oxhaustive search for the non-metallio components. How far the solutions obtained in the disintegrations are available for the latter purpose, and the respective methods of procedure generally, depend chiefly upon whether we merely aim at the detection of the nonmetallic elements as such or at that of the acid radicles contained in the substance. All we could say in regard to the fermer case is anticipated in the section on the 'Detcction of certain elements, &c.' (p. 217) and the latter is not susceptible of being treated instructively in general terms. In regard to it we must refer to the special hand-books.

Quantitative Analysis.

The general problem of quantitative analysis defines itself. Its, solution, scientifically at least, assumes its simplest form, if the thing to be analysed is given as a free substance and the (let us say one) component to be determined can be separated out exhaustively in the form in which it is meant to be reported. In such a caso all that is required, in addition to the analysis proper, is the numerical definition of the two quantities concerned. Of the several direct methods which we have for this purpose, only two need be mentioned; onc is to measure the volume of the body under stated conditions of temperature and pressure, the other is to de-termine its weight. The former method is contined in practice to gases and liquids; the latter is applicable, and indeed is applied, to bodies of all kinds, and, when we have choice, applied preferably. The volume of a body is a function of temperature and pressure, and its numerical statement is consequently encumbered with the necessary reference to-in generaltwo corresponding standards; the weight of a given body depends only on the intensity of gravity, and even this variable in practice is out of court, because, in ohemistry we always use that well-known method of relative weighing which measures the weight of the body not in terms of a unit-force but as a multiple of tho weight, at the time and place, of an adopted standard mass, viz. the unit-piece of our set of

is separated by solution of baryta in a warm liquid which pps. only the magnesium as Mg(OH).

Rubidium and casium, in any scheme of analysis, follow potassium to the end.

weights. The result of such a weighing is independent of the prevailing force of gravity, and consequently not subject to any variation; it consequently, at least, indexes the mass with perfect definiteness. We, moreover, know that it is independent of any chemical change within the body (or set of bodies) weighed. A mass of, for instance, sulphide of copper weighs precisely as much as the two components did conjointly. Hence for chemical purposes our method of weighing might safely be viewed as a method of true mass-measurement, even if Newton had not proved that equal weights (chemically determined) correspond to equal inertias.

The method of direct quantitative analysis explained is the only one which suggests itself when the component to be determined is a chemically indefinito mixture (such as for instance the mixture of salts contained in a natural water); it applies to a good many other cases; but of courso breaks down whenever the body to be determined is an imaginary radicle such as SO₄ or ClO₃, &c. In such cases (and many others as a matter of expediency) wo determine the component by one or other of our indirect methods of mass measurements; which, however, all come to this, that instead of the unknown mass x, we measure some other quantity q, which bears to x a known fixed relation, x = f(q, m), where m is a mass which must be measured directly, although the analyst does not always do this at the time, or himself at all.

Most of our determinations in analysis are uncertain by at least 0.001 of their values, and a higher degree of relative precision is afforded by any fair ordinary balance. But the nature of our methods compels us, in general, to work on small quantities—we rarely care to start with more than one gram of a given solidand besides the products to be weighed can in most cases not be placed on the bare pan, but must be shut up in apparatus weighing perhaps 100 or 1000 times as much as themselves. The nct weight then comes to us only as a small difference between two large weights directly determined. So it comes that even for the ordinary routine of quantitative analysis, we need a balance which to be generally useful should carry about 100 grams on each side, and with this charge turn distinctly with anything greater than, say, 0.2 milligrams,

The Chemical Balance.1

In its present form the chemical balance is nothing more than a refinement upon the ordinary beam and scales to be seen in any grocer's shop; it is a more perfect realisation of the same ideal machine. There is an absolutely rigid beam, suspented so that while it oscillates freely about a certain axis, every point of which is fixed in reference to the stand, it cannot perform any other motion. From two points which lie in the same plane with the axis of rotation-one a near the left, the other B near the right, end,—the pans are suspended by means of absolutely flexible linear strings. and n are equidistant from the axis of rotation. The form of the (ideal) beam is arbitrary; so in

Partly abstracted from the writer's memoir: Ucher die Waage des Chemikers (Zeitschrift für Instrumentenkunde, 1881, 313 et mg.).

- distributed that, supposing the line A B to be horizontal, the centre of gravity of the empty beam lies vertically below, though very near, the axis of rotation. Let us at once add that in the actual instrument the weight of the beam should be no greater than is necessary to ensure to it sufficient stability of form in all circumstances, because the greater the weight of the beam, the greater (cat. par.) the friction in the axis of rotation, and the greater the time of vibration.

Of the difficulties involved in realising the ideal machine, that of producing a light and yet practically inflexible heam seems to have rested most heavily upon the minds of the earlier makors; but there can be no doubt that many of their offorts in this direction (which occasionally resulted in what we should now call fantastical beam forms, such as hollow ellipsoids or double cones, monstrous skeleton forms, &c.) must be traced back to their inability to reach a sufficient degree of precision in the geometric adjustment of the three pivots, and their thus charging against the flexibility of the beam what was really owing to these defects in the adjustment. As these difficulties were overcome, beams assumed less fantastie forms. Sacré, of Brussels, we believe, never uses any but plain rod-shaped beams for even his finest instruments; most balance-makers, however, prefer the form of a largely perforated rhombus or flat isoseeles triangle; and thereby attain all that is needful even for the best instruments without offending the eye by unduly stretching the maximum section, or without using anything more rigid, intrinsically, than hammered brass or some kind of hard bronze.1

In all modern balances the axis of rotation is sought to be realised in a straight knife-edge ground to a prism of hard material, which is firmly fixed to the beam traversing it cross-wise, and rests-in the best balances along its entire length - on a horizontal, plane, (and equally hard) bearing fixed to the stand. The arrestment is so contrived that, besides doing its primary duty, it secures to each point of the knife-edge a fixed position on its bearing whenever the balance works. In former times both bearings and knives used to be made of hard steel; subsequently agate bearings came to be combined with steel edges, and this is still the most popular combination; although Robinson long ago introduced agate knives in conjunction with agate bearings. The agato knife adds nothing to the precision or mechanical durability, but for laboratory balances offers the great advantage of rendering the system proof against acid-vapours; accordingly it is gaining more and more in popularity. Quite lately an American has introduced as a material for both knives and bearings that very hard (and acid-proof) alloy of osmium and iridium which is used for the tipping of stylograph

The point pivots, A and B, used to be realised visibly by means of two oircular knives fixed to the end of the heam so that their working-edges were parallel to the axis of rotation. From the

a sense is its mass, which, however, must be so lowest points, the pans were suspended by meaus of ? shaped hooks of steel wire. In this way a vsry high degree of precision can be attained, and the system when well executed is more durable than one would think, but with balances used for heavy charges it cannot possibly last for many years.

In the better system introduced by Robinson of London half a century ago each extremity of the beam is provided with a knife-edge similar to the central one (except that it is turned upwards); on each knife-edge rests a stirrup-shaped (or J-shapod) contrivance, terminating in a ring below, and from this ring the pan is suspended by a hook. This, of course, comes to the same as if the pan were suspended from the projection of the working point of the hook-and-eye arrangement on the respective knife-edge; so that the latter need not be absolutely parallel to the axis of rotation. Flat ond-bearings demand a somewhat cumbrous and expensive appendage to the arrestment to secure to each point of every edge a fixed position on its bearing in the working instrument. Hence Staudinger, and many others, prefer to combino (long) end-knives with roofshaped bearings, which, in virtue of their shape, fall into their prescribed positions without external aid.

In now passing from fundamentally important to subsidiary points, the arrestment ought to be taken up first; but we could not possibly do justice to this (practically all-important) subject without workman-like drawings and lengthy descriptions. We therefore pass on at once to the needle which serves to define the position of the beam in reference to the plumb-line.

In the precision-balance the needle is made to point downwards towards a scale fixed to the lowest convenient point of the pillar. The zeropoint defines the 'normal position' of the beam, i.e. that position in which its centre of gravity lies vertically below the axis of rotation. The scale is so divided that the radii drawn from the axis of rotation through the marks divide the tangent to the circle described by the oscillating ncedle, at the zero point, into pieces of equal length, which in most practical cases means

into degrees of equal angular value.

To avoid the use of small weights, each arm of the beam, in most balances, is divided into ten equal parts in the sense that the projections of the marks on the line a B convecting the two point-pivots divido the distance from the oentral pivot to (say) B into ten equal parts. rider weighing ten mgms., when suspended on mark 1, 2, 3, &c., acts like 1, 2, 3, &c., mgms. placed on the pan. In most balances, however, points 0 and 10 are inaccessible. Sons, of Rotterdam, avoid this inconvenience by dividing each arm into twelve equal parts, and providing a rider of twelve mgms, weight. Some makers make the top bar of their beams straight, and exactly parallel to the plane of the three pivots, and let it project beyond the terminal edges, besides keeping it clear of encumbrances, so that the rider can move freely from one end of the team to the other. This system, besides its obvious advantages, admits of the use of heavier riders; because the increase in sonsibility caused by the presence of the rider is the same: at any position which it may have; only the

For further information regarding this question we refer to the writer's memoir quoted in footnote to p. 223.

rider in such cases must be counted part and In formulating the relation between a small parcel of the instrument.

In proceeding now to develop the statical theory of the precision-balance we will assume, for a first approximation, that the three pivots are physically and geometrically perfect in themselves, but, for the sake of greater generality, we will not assume that the knives are exactly in their intended positions. Imagine a system of rectangular co-ordinates fixed to the beam so that, while the z-axis ceincides with the axis of rotation, the x-axis goes through the centre of the middle knife, and runs parallel to the line an which joins the two point-pivots. Let the co-ordinates of a and n, of the centre of gravity so of the empty beam, and of a certain

point o to be defined presently, be as follows:

$$\begin{array}{cccc}
x & A & B & S_0 & O \\
x & -V & +V' & O & X_0 \\
y & +h^* & +h^* & s_0 & y_0
\end{array}$$

Let P' denote the tetal charge from A, P" that from B, and we the weight of the empty beam; the joint effect of these three weights is the same as if they were all concentrated in some point oat $x = x_0$ and $y = y_0$. For calculating purposes we may assume gravity in one case I. to act in the direction of the y-axis, and in a second II. in the direction of the x-axis, and it a second (Case I.) . $\mathbf{r}' l'' - \mathbf{r}' l' = (\mathbf{r}' + \mathbf{r}'' + \mathbf{w}_0) x_0$. I. (Case II.) $(\mathbf{r}' + \mathbf{r}'') h + \mathbf{w}_0 s_0 = (\mathbf{r}' + \mathbf{r}'' + \mathbf{w}_0) y_0$. II. In any sensibly constructed balance things

are so arranged that, under all circumstances that come inte practical consideration, the centre of gravity o of the whole system lies outside and below the axis of rotation (i.e. that $y_0 > 0$). Assuming both x_0 and y_0 to have positive values, and the beam to be left to itself in its nermal position, it will turn, and tend to assume that position in which c lies vertically below the axis of rotation. The radius oc then describes an angle equal to that which separates oc from the x-axis, and obviously,

tan.
$$\alpha = \frac{x_0}{y_0} = \frac{\mathbf{p}'' \mathbf{l}'' - \mathbf{p}' \mathbf{l}'}{(\mathbf{r}' + \mathbf{p}'') \hbar \mathbf{w}_0 s_0}$$
. III.
To bring the equation into a handler form

for our purposes, let us separate P' and P" into parts, thus; $\mathbf{r}' = p'_0 + p'$, and $\mathbf{r}'' = p'_0 + p''$ where the p_0 s stand for the weights of the empty

where s has an obvious meaning. We then have for x_0 the equation

 $p''l'' - p'l' = (p' + p'' + w)x_0.$ This equation may be said to state the theory of the ordinary method of weighing. To find the weight p' of a given body we place it on the left pan, and then try heavier and lighter combinations of standards on the right, until we have found out that one (representing p'' grams) which reduces x_0 to nothing, so that the balance is at rest at, or it vibrates about, its

normal position. We then have $p' = \frac{p''l''}{l'}$

Von L

4. 7.

overweight on one side and the corresponding angle of derivation α , we may take $l' = \overline{l'}$ (as it really is very nearly in all well-adjusted balances).

where a means the angle through which the pesition of rest turns in consequence of the addition of a units of weight to the right pan, the charge before leaving been x=p on each

In practice tan. a is measured in degrees of the scale. Supposing a corresponds to n degroes of the seale, and tho index-length is J in degrees, we have

$$\frac{n}{s} = \frac{\Delta l}{ws + 2ph} \quad . \qquad . \qquad . \qquad . \qquad V.$$

The ratio $\frac{n}{\wedge}$ defines the sensibility of the balance; we have for it

$$E = \frac{n}{\Delta} = \frac{l \ J}{Ws + 2ph} \quad . \qquad . \qquad . \qquad VI$$

 $E = \frac{n}{\Delta} = \frac{l}{ws + 2ph}$. . . VI. and for its reciprocal, $\frac{1}{E}$, the weight-value of 1° of the scale,

$$\frac{1}{E} = \frac{ws + 2ph}{l \ J} \qquad . \qquad . \qquad . \qquad . \qquad . \qquad VIII$$

For h=0, the term 2ph vanishes, and the sensibility becomes independent of the charge. In the actual instrument h is a function of the oharge, of the form $h = h_0 + \beta p$, where β is a small constant depending on the coefficient of elasticity and the configuration of the beam. For a given charge, a good maker has no difficulty in bringing h down to less than ± 0.01 mm. The bost instruments are so adjusted that, for a ocrtain medium charge, h=0, so that for p=0 it has a small negative, and from p=maximum charge a small positive, value. The relative change in the sensibility involved in passing from p=0 to p=p, is shown by the equation

$$\frac{\mathbf{E}^{-1} - \mathbf{E_0}^{-1}}{\mathbf{E_0}^{-1}} = \frac{2ph}{\text{Ws}}$$

and consequently is the less (cet. par.), the greater s, i.e. the less the initial sensibility, Ba. It (i.e. the left side of our equation) comes to its minimum (assuming p to represent the maximum charge) if the balance is so adjusted that, for the charge 0.83 p, h=0. Supposing this rule to be generally adopted, the relative inconstancy of the sensibility is independent of the arm-length (see the writer's memoir, p. 318).

No balance is complete without a gravity. bob,' a small butten or sphere of metal attached to a wire which stands vertical on the top of the beam (in the x-axis) so that it can be screwed up and down into any position. Matters are arranged so that when the bob is quite down the sensibility is below the lowest value we care for, while, by screwing up the best to its highest place, we can bring C in the eyen a little to the wrong side of, the axis to bittion. Hence it would appear that, by screwing up the bob sufficiently we can get our balance to turn

For more exact information see the writer's memoir, page 322. + means 'below a.

visibly with say 0.001 milligram or anything less that we might care to name. So indeed it would be if our fundamental assumptions could be-and were-realised. In practice, however, the knife edges are not absolutely straight nor the bearings absolutely plane, and neither are absolutely rigid. Hence the three axes, instead of being always at x = A, o, B respectively, so to say oscillate irregularly, each from $x-\lambda$ to $x + \lambda$, where x is the theoretical x. In going more fully into the matter we see that as a consequence the balance at a given charge (say p left; p right) is in a state of indifferent equilibrium within a small angle $\pm \beta$, which, of course, is the greater the greater is E. But the weightvalue e of this angle is constant, and is governed by some equation like

$$\epsilon = \frac{\lambda}{1} (\mathbf{w} + 2\mathbf{p})$$

where 'A' is meant to lump the joint effect of the three As previously referred to.

may be called the 'unherent error' of the balance. There is obviously no use in screwing up the bob any further than necessary to render this ϵ (i.e. angle β) distinctly visible. It may be inexpedient even to go so far, because, in practice, we never aim at the absolutely true weight, but at a value sure to differ from it by no more than say ± 0.1 mgm. The angle corresponding to this need not be more than distinctly visible. To make the angle greater than necessary needlessly adds to the time of vibration which may already be inconveniently high. Because the time of vibration (t in seconds) is governed by the equation

$$t^2 = \frac{l^2 \{ k w_0 + 2(p_0 + p) \}}{R \{ w_0 s_0 + 2(p_0 + p)h \}}$$
 . VIII.

where R denotes the length of the pendulum beating seconds at the place of observation. kwel2 denotes the momentum inertice of the empty beam in reference to the axis of rotation. The denominator of eq. VIII. suggests the expression given in eq. VII. for the sensibility z. Combining the two we have

$$\ell^2 = \frac{l}{RJ} - \{ (kw_0 + 2p_0) + 2p \} E$$
 . IX

The bob enables us to choose our own E. or our own t (for a named charge), but it does not enable us to choose both. We of course refer to a ready-made balance; in the hands of a mechanician who designs a balance for a stated purpose, I becomes an arbitrary variable, and the equation then assumes something like this

$$t^2 = l(c + kbl + 2p) E$$
 . . . X

where c and b are constants whose meaning is sufficiently apparent. In words: Whatever (reasonable) value for E may have been fixed upon we can bring down t (for say p=0) to any desired figure by making l sufficiently small. But where shall we stop? For high-class balances intended to weigh up to 100 grams, mechanicians used to draw the line at l = 180 to 200 mm. These values (perhaps more by dint of habit than on rational grounds) were retained until about twenty years ago, when P. Bunge, of Hamburg, introduced a new form of the in-

strument, in which the arm-length is reduced to some 60 to 65 mm. Thanks to the general excellence of Bunge's work, these short beams soon became very popular among both chemists and mechanicians; and it therefore is worth while to inquire what their specific advantages as short-beam balances amount to.

For this purpose the writer, some years ago, determined the constants of eq. X. for a very excellent Oertling (hectogram) balance, which he has in his possession, (its l=184 mm.), and, taking it as a general model for an imaginary genus, calculated the values of t for a number of charges and sensibilities, assuming I to be equal to (a) 180 mm. and (b) 60 mm. He found for

I. E = 2 degrees of the scale per 1 mgm.

of over-weight.¹
if
$$l = 180 \text{ mm}$$
. 60 mm.
for $p = 0$; $t = 7'' \cdot 7$ 3".6

The short beam obviously vibrates too fast for high-precision work. To set this right let us screw up the bob on both sides, so as to double the sensibility. We now have

II. E = 4 degrees per mgm. if l =180 mm. 60 mm. for p = 0; $t = 11'' \cdot 0$ $5'''\cdot 2$ as p = 30; t = 14''.87".8 as $\bar{p} = 100$: t =11".6 21".2 2.23 1.93 $t_{100}: t_{0} =$

The times of vibration no doubt assume the more convenient values in the shorter beamed instrument. But what does this amount to practically? In our opinion not to as much as some people seem to think. We are inclined to think that the short beam offers material advantages to those who are accustomed to the deadbrat method of weighing (see below). All those who prefer the method of vibration will on the whole, we think, fare better with the old form of the instrument. But this, to be complete, should be provided with the two following auxiliary contrivances of the writer's invention:

I. An auxiliary small bob2 attached by mere friction to the upper part of the needle, which has the form of a triangular prism and is (virtually or actually) graduated, so that one is able, at a moment's notice, to give to the weight-value of 1° of the scale any convonient pre-determined value, to make it equal to oxactly 2, 1, 0 5, 0 2 &c. mgm. as he may please.

II. A microscopic arrangement 3 for reading the excursions of the needle. A narrow ivory scale, divided into very small degrees, is fixed to the needlo near its lower end, so that a microscope which is fixed slantingly to the stand but passes through the central (fixed) portion of the front pane enables one to read it. The microscope has a vertical who in its focus; this wire appears as a vertical line crossing the image of the scale. Every degree of the micro-scale corresponds to exactly 0.1 degree of the ordinary scale, which latter doos duty as usual. As the microscopo reversos the imago, tho apparent motion of the 'wire' on the micro-scale is in the

^{1.} Let the addition of 1 mgm. causes the needle to oscillate between 0 and $\pm 2^\circ$; 1° in the given instrument 1 mm. very nearly.

1 P.F. K., 1873; C. N. 83, 157.

2 P.A. L. 1874; C. N. 83, 157.

same sense as the real motion of the needle in reference to the ordinary scale, so that there is no fear of blunders through mistaking plus for minus. The writer is indebted to Mr. Oortling for having executed this arrangement for him in a most masterly manner. Though intended originally to be reserved for spocial work, such as weight-tosting &c., it was found so convenient that both the writer and his assistants use it preferably for even their everyday weighings. The specific advantage of the microscope is that it enables one to adjust the 'bob' so as to produce the most convenient time of vibration. The microscope more than makes up for the involved loss of sensibility.

On Weighing.

A precision-balance should stand on an unshakable table, and should not be exposed to the risk of one-sided elevation of temperature. Before being used for a series of weighings it must of courso be set in order, which includes that the case be 'levelled,' so that the plane including the axis of rotation and the zero of the scale is a plumb-plane. The next thing to do (if necessary) is to bring the 'bob' into its proper position; i.e. to place it so that the least difference of weight we care for just becomes visible as an anglo of deviation and no more, because to increase the sensibility beyond what is needful means needlessly to diminish the range of weights determinable by vibration, the constancy of the sensibility, and the rate of vibration. This rate of course must not be allowed to fall below a certain limiting value. In the writer's opinion, t = 5" is about the lowest permissible limit for relatively heavy charges. Next, the balance must be brought 'into equilibrium' at least approximately. For this purpose Oertling's balances carry a vane at the top of the beam, consisting of a little lever hinged to the wire of the bob, which can be turned round, so as to shift the centre of gravity to the right or left.1 A better arrangement is a small horizontal gravity bob at one end of the beam. For simplicity's sake we assume that the balance has been brought into perfect equilibrium, so that the needle in the vibrating instrument moves forwards and backwards between $+n^{\circ}$ and $-n^{\circ}$. To weigh an object (which, to fix ideas, we will assume to be a solid, and non-hygroscopic), the ordinary mode is to place it on the left pan, and thon counterpoise very nearly with standard weights, say p grams, on the right. In order new to determine the small additional weight which is required to establish perfect equilibrium, we may use one or other of two methods. In the

Dead-beat method we simply centinue our trials, until the needle vibrates about the zero as its position of potential rest. It is, however, hardly possible for any thinking person to use this method without at least instinctively combining it with

The method of vibration, which in its most exact form consists in this that we note down (at least mentally) the successive excursions of

A vane with properly graduated limb is as good as a 'rider'; better in fact, inasmuch as it is not liable to drop off and get lost; this innovation was proposed by Hempel, but has not met with much favour as far as we know.

the needle, and from these calculate the position of rest. Supposing we count distances traced by the needle in moving from 0 to the left as positive, and those to the right as negative, and the needle turns successively at n_1 , n_2 , n_3 , n_4 , n_2 , degrees, we have for the position of rest, 4 values,

 $\frac{1}{2}(n_1+n_2)$; $\frac{1}{2}(n_2+n_3)$; $\frac{1}{2}(n_3+n_4)$; $\frac{1}{2}(n_4+n_5)$, and the mean of these four values gives the reading corresponding to the position of rest. But the factor a can be dropped, because we are evidently at liberty to measure in half-degrees. By taking an odd number of readings we climinate the error caused by what the needle loses in passing through its path; for ordinary practice 3 readings are sufficient. It suffices to determine $n_1 = (\text{say}) + 4 \cdot 2$; $n_2 = -1 \cdot 0$; $n_3 = +4 \cdot 0$, and compute 'a' = mean of +3.2 and +3.0 = +3.1. In this case the right pan is too heavy by 3.1 x k mgms., if k is the weight-value of 1° at the respective charge. In a good balance k is almost independent of the charge; the writer's supplementary bob of course enables one to give it a pre-determined value. How k is determined ueed not be explained.

Supposing p grams to have established exact equilibrium, the object weighs

$$x = p \frac{l''}{l'}$$
 grams.

The several weights which enter the calculation of an analysis need only be relatively correct. Hence, if all the weighings involved are made on the same balauce and with the same set of weights, and the objects are always in the left pan, in any such series we may adopt $\frac{l''}{l'}$ grms.

as our unit and say x = p.

We do not consider it necessary to quote examples of cases in which as a matter of principle $\frac{l''}{l'}$ dare not be cancelled; we rather say that in all precision-balances worthy of the name $\frac{l''}{l'}$ is very small, not more than 0.00005 at the most. If the empty balance was in equilibrium at $+a_0$ degrees we must add, if at $-a_0$ degrees we must subtract, a_0k mgms. from p.

Absolute Weighing.

Absolute precision-weighing in the chemical laboratory hardly occurs otherwise than in this sense that we may have to determine the weight of an object in terms of an arbitrary (but for this occasion absolute) standard. For this we have two methods.

I. The Method of Substitution. The object is placed in one pan of the balance, and counterpoised exactly by some suitable tare placed in the other. We then take off the object and put on standard weights until equilibrium is again established. If the method of vibration be used, the immediate result is the proof that the constant tare was balanced by (1) x grams of object plus δ grams, and (2) by $p + \delta_2$ grams of standards. Whence $x = p + \delta_2 - \delta_1$.

II. The Method of Reversion. After having brought the balance very nearly into equilibrium, we ascertain the number of grams which have to be placed in the opposite pan to exactly balance the object, once with the object on the left, and once with the object on the right, side, Assuming, for greater generality, that the right pan was from the first too heavy by δ grams, we

I. $xl' = (p'' + \delta) l''$ by the first trial. 11. $(x + \delta) l'' = p'l'$ by the second trial.

Assuming for a moment that l'=l'' (as we always may if x is small), we obviously have $2x + \delta = p' + p'' + \delta$; or $x = \frac{1}{2}(p' + p'')$.

We will now drop this assumption, but assume that & is so small that the balance cannot distinguish between $\delta l'$ and $\delta l''$; then we may write as

I.
$$(x-\delta) l' = p''l''$$
.
II. $(x+\delta) l'' = p'l'$.
multiplication,

Whence, by multiplication,
$$p'p'' = (x-\delta)(x+\delta) = x^2(1+\frac{\delta}{x})(1-\frac{\delta}{x}) = x^2(1-\frac{\delta^2}{x^2}).$$

It is always possible to make a guess at the maximum value which $\frac{\delta^2}{x^2}$ could possibly have;

supposing $\delta = \pm 0.001$ grm. and x (i.e. p' or p'') = about 10 grms. $\delta^2 \div x^2 = 1 \div 10^9$ and can be neglected. In practice we take care not to allow & to assume a greater relative value, and compute by $x^2 = p'p'$ or $x = \sqrt{p'p'}$, for which expression we may substitute $\frac{1}{2}$ (p' + p''), if p' and p'' differ by less than, say, p mgms.

On Sets of Weights.

A set of weights to be fully on a par with a given balance must be so exactly adjusted that no combination of the several pieces which can ever occur is wrong by more than the inherent orror 'e' (v. supra) of the instrument. This means that chemical weights, to be properly adjusted, require a balance of a very high order. But even the most perfectly adjusted set is of no permanent value unless the substance that it is made of offers a sufficient guarantee for constancy of mass. Of all available materials, rock-crystal comes nearest perfection, but it is difficult to work and bring into a handy shape. Of mctals, Mr. George Matthey's ten per cent. iridio-platinum is the best; it is absolutely proof against even acid fumes, and sufficiently hard to be proof against abrasion by reasonable usage. Next after it comes 'hard' platinum (the slightly iridiferous metal of which crucibles are generally made); pure platinum is too soft. Brass, bronze, German silver, and other cheap motals are mere apologies for what ought to be used; yet these are used (in a sense unavoidably) for making the larger pieces in sets for every-day use. Gilding affords no protection against atmospheric influences, unless the noble metal is laid on thickly; a good lacquer is better than the film of gold which is customarily put on by electrolysis.

In constructing a set of chemical weights, we might choose our own unit, but whatever unit we might fix upon, any other mode of subdivision or multiplication than the decimal mode would be absurd; and there is no reason why we should not adopt some legally fixed and universally obtainable unit as our unit. The gram is used by chemists all over the world, almost to the exclusion of any other unit.

Sets of weights exact enough for all practical purposes can be had in commerce. Whoever may be the maker, a set of weights should not be used without having first been tested and found correct, at least in a relative sense. To show how the errors in a given set can be determined, let us assume for a while our set comprised only the pieces (1), (1), (2), (2), (5), (10) grams, and adopt these bracketed numbers as symbols for the unknown true weights. As a unit for the errors to be determined, we will adopt the 1 milligram as determined by a given rider of 10 mgm. weight; the (1), shall scree as our provisional unit for the values (1) (2) (10). To determine (1) we compare it with (1), by the method of substitution or reversal, and note down the difforence between the two in terms of 'the milligram,' as determined by the method of vibra-tion. We then compare (1)₀ + (1) with (2); then (2) with (2), &c., &c., to establish the following equations:

$$\begin{array}{lll} (1) &= (1)_0 + \delta_1 & \text{mgms.} \\ (2) &= (1)_0 + (1) + \delta_2, & , \\ (2)_1 &= (2) + \delta_2', & , \\ (5) &= (2) + (2)_1 + (1)_0 + \delta_5, & , \end{array}$$

Computed.

To know what the values $n \times (1)_0$ really are in terms of an adopted gram (say the true gram) we must compare one of the pieces, or a combination of some or all, directly with the corresponding standard weight. Supposing this had been done with the 10 gram piece, and this piece had been found free of error, we have $10 \times (1)_0 + \Delta_{10}$ mg. = 10 g. (meaning 10 true grams)

...(1)₀ =
$$\frac{10 \text{ g.} - \Delta_{10} \text{ mg.}}{10}$$
 = 1 g. $-\frac{\Delta_{10} \text{ mgs.}}{10}$ mgs.

and by substituting this value for (1), in the expressions $n \times (1)_0 + \Delta_n$ we obtain the values of all the six pieces in the form

$$(N) = N g. + x mgm.;$$

but our 'mgm.' is strictly speaking an arbitrary unit; we have no right, for instance, to say

$$(5) = 5 \,\mathrm{g.} + \frac{x_5}{1000} \,\mathrm{g.}$$

What the true gram-value of the rider is can only be found by joining on to our gram set a set of deci- and centi-grams comprising that rider, and determining their values by establishing the equation:

$$\begin{array}{l} (\cdot 01) = (\mathrm{rider}) + \delta_1, \\ (\cdot 02) = (\mathrm{rider}) + (\cdot 01) + \delta_2, &c. \ \mathrm{up \ to} \\ (1)_o = (0\cdot 5) + (0\cdot 2) + (0\cdot 2)_1 + (0\cdot 1) &c. \end{array}$$

and thus finding out the value of the rider in ... terms of g. But in practice the rider as a rule does not differ much from 'Olg., and this part of the work is not necessary for the sake of the calculation of the errors, the less so as a great value in any of these would simply condemn that picce.

The above method is always used when we test a set of weights with the view of seeing how it falls in with the rest of the sets in the labo. ratory, which in the aggregate form our set for

general purposes. If the set is meant to be used by itself—if, for instance, we test a set from 50 grms. down to 1 centigr. with the view of using it for our analyses—it is better not to refer to any external standard at all, but to an imaginary nnit so chosen that the sum total of the errors becomes nil, i.e., to choose as nnit $\frac{1}{100}$ of the actual weight of all the '100 grams' which tho 50 gram set represents in toto. If one or more of the pieces come out with relatively large errors, the nnit is re-adjusted so that it suits only the good pieces, the errors are re-calculated, and the two rejected pieces either replaced by new ones, or re-adjusted. According to the writer's experience, we must be satisfied if the errors of the individual pieces are brought down to values varying from very little to about ± 0.04 mgm.

Reduction to the Vacuum

All weighings executed in air are liable to an obvious correction. Supposing an object occupying v c.o. is balanced in air by p grams of standards occupying v o.o.; if the balance were transferred to a vacuum, the side of the greater v (in our case the object side) would become heavier than the other by (v-v) δ grms. where & is the weight of one c.c. of air at the time and place. As p is a close approximation to the true weight, the volume of the object in o.o.s can be put down as $p \div s$, that of the standards of course is $p \div s_v$, where s and s_v are the respective specific gravities which practically need not be reduced to water at 4°. The correction to be applied to p is

$$p\delta\left(\frac{1}{8} - \frac{1}{8_0}\right),$$

 $\delta = 0.46464 \frac{B}{273 + t} \text{ (mgms.)},$

where B is the height of the barometer in mm. reduced to 0°C., and t is the temperature; the constant is calculated from Regnault's weight of 1 litre of air of 0° and 760 mm. at Loudon. For $t=15^{\circ}$, and B=760 mms., $\delta=1.22615$,

which number, at stations where B is habitually near 760, if the highest precision is not aimed at, may often be taken as holding for air gene-

Standard weights for absolute weighings (in trne grams) ought to be adjusted for the vacuum; hence, if the minor weights are of platinum and the larger ones of brass, the brass I grm. should appear lighter than its equivalent in platinum decigrams in air. But sets of this order had better be made of one metal.

For a series of relative weighings, tho buoyancy of the weight-standards in air may be neglected, because we are at liberty to take as our mnit the weight of the 1 grm. piece in air of the average density prevailing during the progress of the experiments. That this unit is estrictly speaking variable is of no practical significance.

The vacuum-correction for any single weighing involved in an analysis amounts as a rule to more than we should care to neglect; yet it may be neglected in most cases, because ths weight to be determined is only one term of a ratio, whose other term is faulty in the same Suppose we have determined two

approximation. If the reciprocals of the specific gravity are s, - and s, -1 respectively, we have

$$x = \frac{p_1}{p_2} \quad \frac{(1 + \delta_1 s_1^{-1})}{(1 + \delta_2 s_2^{-1})};$$
 or as a sufficient approximation

$$x = \frac{p_1}{p_2} \qquad (1 + \delta_1 s_1^{-1} - \delta_2 s_2^{-1}).$$

 $x = \frac{p_1}{p_2} \quad (1 + \delta_1 s_1^{-1} - \delta_2 s_2^{-1}),$ And if s_1 does not differ much from s_2 (as δ_1 and δ_2 are always nearly the same) the bracketed factor may come close to unity although neither of the two terms s-15 could be neglected if it stood by itself. Here, as everywhere in experi-mental science, the golden rulo is neither to strain at the gnat nor to swallow the camel.

Weighing of Gases.

For the weighing of a gas, we have in general three methods.

- (1) If the gas to be weighted is a product formed in a reaction between solids or liquids, we may identify its weight with the loss of weight suffered by the reagents during the reaction.
- (2) We may collect the gas in a evacuated tared balloon, and weigh it like anything else. In this case it is expedient to tare the flask with another flask of the same displacement and nearly the same weight, so that only a few grams need be put on to establish equilibrium. (Regnault's method). The vacuum-correction is then out of court.
- (3) We may measure the gas by volume at a known temperature, t, and pressure, P, and calculate the weight from the velumo.

If the gas to be weighed is a product of a reaction carried out quantitatively, one way of measuring it is to construct the upparatus so that the vessel in which the reaction goes on and tho gas-measurer have a common atmosphere, and to measure the gas-volume as an increase in the total atmosphere of the apparatus (gasvolumetric method). According to Regnault, 1 lit. of oxygen at 0° and 760 mm. of mercury of 0°C., weighs 1.42932 grams. Hence by an easy computation, based on Avogadro's law, we have for the weight of 1 litro of a given species

of gas of the molecular weight M (0 = 16)

$$\mathbf{D} = 0.032089 \frac{\mathbf{P}}{(273 + t)} \times \frac{\mathbf{M}}{2} \text{grams}$$

where r means the dry pressure in nim.

If the gas is meist, the vapour pressure of steam at to must be deducted from the observed pressure to find the D of the formula. constant, strictly speaking, holds only for places where gravity is the same as it is at 45° latitude, and sea-level.

In Paris, | London, | Berlin, | Glasgow, it must be multiplied by 1.000333, | 1.000563, | 1.000663, | 1.000956, .

Indirect Methods of Mass Measurement.

I. Physical Methods.

The nature of these is best explained by a general example. In a given aqueous solution of sulphurie acid, sugar, salt, &c., &c., the specific gravity at to, the refractive index, the weights, p_1 and p_2 , and we want the correct value power of turning the plane of polarised light p of the ratio of which $p_1:p_2$ is only an (if any), &c., bear each a fixed relation to the percentage of substance (or the weight of substance per litre) in the solution, which relation is susceptible of translation into a formula p=f (physical property), or a corrosponding curve, and by means of oither of tabulation. Hence, supposing the function to have been determined by standard experiments, p in a given case can be calculated (virtually or actually) from the value of the respective physical property. In practice we must of course try to establish conditions under which the change in the specific gravity, &c., &c., corresponding te the passing from p to, say, (101) p, assumes a sufficiently great value.

The popular method for determining the strength of aqueous oil of vitriol, &c., by means of a hydrometer may be referred to as an example. The customary method of deducing the percentage of sugar in a syrup from the angle through which a column of given length turns the plane of polarised light is another.

II. Chemical Methods.

These, being all founded upon our knowledge of the quantitative laws of certain reactions, are methods for the indirect weighing of radicles rather than of substances. Scientifically one might arrange them according to their degree of directness. If we do so, the following twe claim precedence before any of the rest.

(1.) The direct gravimetric method. An unknown weight of this or that radicle is determined by separating it out exhaustively, by itself or as part of a compound of known composition, and weighing the product either directly on the balance, or perhaps indirectly by gasometric measurement.

(2.) The method of titration. An unknown weight of radicle is deduced from the quantity of reagent necessary and sufficient to cause it to undergo a certain definite change of combination; the quantity of reagent being ascertained synthetically, i.e. by direct trial.

These two methods we will designate as 'direct' methods in opposition to the following 'indirect' methods.

- (3.) The method of substitution. Instead of determining a radicle \mathbf{r} , we substitute for it an equivalent of some other radicle (or substance) \mathbf{r}' ; we determine \mathbf{r}' by method I. or II., and from it calculate \mathbf{r} . Thus, to determine an unknown weight of free chlorine, \mathbf{r} Cl. mgms., we substitute \mathbf{r} I, mgms. by means of the reaction $\mathbf{Cl}_2 + 2\mathbf{K}\mathbf{I}\mathbf{Aq} = 2\mathbf{K}\mathbf{C}\mathbf{I}\mathbf{Aq} + \mathbf{I}_2\mathbf{Aq}$, and determine the iodine. In some cases we effect a series of substitutions (\mathbf{r}' for \mathbf{r} ; \mathbf{r}'' for \mathbf{r}' , \mathbf{Ac} .), and determine only the ultimate substitute. Thus, to determine \mathbf{r} CrO₃, we substitute first $\mathbf{r} \times 3\mathbf{Cl}$, then for this we substitute $\mathbf{r} \times 3\mathbf{I}$, and by ascertaining the value $\mathbf{r} \times 3\mathbf{I}$ we find $\mathbf{r} \times \mathbf{CrO}_3$.
- (4.) The residue-method. The body containing the radicle is subjected to a definite chemical change by means of a known (excessive) weight of reagent, and the excess of reagent left is determined.
- (5.) Methods founded upon the numerical difference between formulæ-values:—We pass at once to examples:—
- (a.) To analyse a mixture of the compounds AgCl and AgBr, we expose a known weight to the action of dry chlorine until all the AgBr hae

become AgCl, and determine the decrease of weight involved. From the obvious equation of the reaction, we see that every Br.Cl gram of loss of weight corresponds to Br grams of bromine, or AgBr grams of bromide of silver.

(b.) To determine the weight of real snlphuric acid contained in a given quantity of an aqueous acid, we evaporate with a known (excessive) weight of anhydrous carbonate of soda, and weigh the residue (Thorpe). As

x Na₂CO₃ + Na₂CO₃ + H₂SO₄ = CO_2 + H₂O + Na₂O₄ + xNa₂CO₃, every (SO₃ - CO₃) grams of increase of weight indicate SO₃ grams of sulbhur trioxide.

(c.) To analyse a mixture of the sulphates of sodium and lithium; take p grams of the mixture, ppt. all its sulphurio acid with barium chloride, and weigh the barium sulphate.

1 grm. of the sodium salt gives BaSO, Na.SO,

1 grm. of the lithium salt gives $\frac{\text{BaSO}_1}{\text{Id}_2\text{SO}_1} = l$, grms

of barium sulphate. Hence if c grms of pp. were obtained we have (from x grms of sodium sulphate, and y of lithium sulphate)

$$x \times s + y \times l = c$$
$$x + y = p.$$

Whence x and y are easily calculated.

Many other examples might be quoted.

We will now pass to The Operations involved in quantitative determinatione by chemical methods. first let us say a few words about a necessary preliminary to any quantitative analysis, namely, the preparation of the sample. This problem assumes perhaps its most difficult form if the thing to be analysed is a large mass of imperfectly homogeneous matter, say, a cargo of copper orc. In this case the analysis must of course be preceded by the preparation of a sample, which, although it may amount to ouly a few pounds, yet can be assumed with a sufficient degree of probability to have the composition of the whole heap. And supposing even a homogeneous sample to have been prepared for the analyst, a mere assay (of the copper in our case) would be of no use unless accompanied by determinations of the moisture in, a the ore as it lies, and b the small ultimate sample which goes to the balance. An impure specimen of a named chemical species to be rendered fit for the

and allow for them in the calculation.

In now passing to our subject, we will give the first place to the operations involved in those

analysis of the species must first be purified,

unless we prefer to determine the impurities,

I. Assays by igneous operations,

which are so extensively employed in practical metallurgy. These, however, are quite a speciality which is almost entirely confined to metallurgical laboratories. Suffice it, therefore, to say that these assays, as the name indicates, are, at least by original intention, processes of metal-smolting carried out tentatively on a small scale. The metal is separated out as a regulus either of the metal itself or of some definite arsenide, and in either form is weighed directly on the balance-pan.

II. Quantitative expulsion of volatile components by exposure of the substance to regulated temperatures.

Under this heading fall most of our methods for the determination of water, given in combination with wen-volatile residues. Water thus combined, can, as a rule, be driven out with or without the help of a dry atmosphere, by prolonged exposure of the substance to a suitable temperature, and, if other changes are known not to take place, the weight of the water expolled is the loss of weight involved in the drying process.

If the residue, while giving up its water, takes up oxygen or suffers some other change involving change of weight, the water must be expelled in an apparatus so constructed that the steam can be purified (if necessary) and collected without loss by absorption in a weighed U-tube filled with chloride of calcium, or pumice moistened with sulphuric acid, and determined as an increase of weight of the absorption apparatus. We have no means of discriminating experimentally between water present as such (moisture) and water present in chemical combination: nor can we discriminate analytically between the different states of combination which we distinguish in our formulæ. All the analyst can do (after removal of what there may be of palpably free water by mechanical means) is to try, successively, exposure to (1) ordinary 'dry' air; (2) artificially dried air, or a dry vacuum; (3) a graduated series of higher temperatures, such as 100°, 120°, 150°, 200°, in a hot air chamber; (4) a red or perhaps a white heat; and to report the several losses of weight, taking care of course to apply each temperature again and again, until the weight of the residue (or of the calcium chleride tube) has become constant.

From hydrates undecomposable by mere heating, the water must be expelled by suitable reagents. Basic hydrates, like caustic potash, can be dehydrated (quantitatively) by fusion with excess of anhydrous bichrouate of potash; many hydrated acids, by evaporation of their solutions with a known excessive weight of oxide of lead, and weighing the dried residue. The writer directs attention to the applicability of tri-sodie phosphate as a weighable form of Na₂O for the latter purpose.

III. Combustions in glass tubes, v. Analysis, Organic.

IV. Carius' general method of ultimate organic anatysis, v. Analysis, Ordanic.

V. Gas evolutions.

We here refer to a class of methods in which the thing to be determined is measured by the weight of a gas evolved in a wet-way reaction of the substance to be analysed. The gas evolved is weighed as loss, or after absorption

According to the current notions on dissociation, a current of (originally dry) air which has passed over a smillent column of partially dehydrated sut, should take away the free water from a given specimen of motif salt of the same kind at the same temperature. Hence an obvious (sheoretical) method for recognising free water as such.

by a suitable absorbent, or is measured (and thus indirectly weighed) gasometrically (v. supra).

VI. Electrolysis.

Solutions of many heavy metallic salts, when subjected to a galvanio current under suitable conditions, are fully decomposed, in the sense that all the metal separates out as such on the negative electrode. By properly regulating the strength of the current and the composition of the liquid, it is possible, in many cases, to cause the whole of the metallic pp. to assume the form of a coherent, truly metallic, deposit, so that, if a platinum electrode be used, the metal can be determined as an increase of weight of the latter. The method, however, is not as easy as it is obvious; and is in general use only for two metals, namely copper (Luckow's method), and nickel. Classen haz tried, not without success, to extend the method to many other heavy metals; but his processes have failed so far to become popular.

VII. Gravimetric Precipitation.

Our heading refers to the very large number of cases in which we determine a component of a solution by separating it out in an insoluble form, and weighing the pp. or converting it into another body and weighing that. With the preliminary separations that may be necessary we can have nothing to do here; we assume that the ppn. has been effected exhaustively, and that the pp. is (in the sense of the method) free from foreign components. In this case the next thing to be done of course is to separate the pp. completely from the mother-liquor. In some cases (for instance in the case of the haloid salts of silver, and of metallic gold ppd. by ferrons salt) this can be done by decantation; but as a rnle it is necessary to resort to filtration. For this operation the first requisite is a good funnel, and good filter-paper. The funnel should be a smooth cone of exactly 60° aperture, so that a filter folded in quarto fits it exactly. The best filter-paper for general purposes is that Swedish paper known as Muntkell's; only it filters somewhat slowly, and in many cases, therefore, papers of looser texture are preferable. Filters, which, having been washed with hydrofluoric and hydrochloric acids, leave almost no ash on ineineration. In conducting a filtration, the following rules should be observed:

1. Before starting the filtration, allow the pp. to settle completely; then decant off the liquor on to the filter, allowing as little of the pp. as possible to go on to the paper. -2. The same rule holds for the first stages of the washing process; the hulk of the pp. should go on to the filter only after almost all the dissolved matter has been washed away. -3. The washliquor should be employed in small instalments, and each instalment be allowed to drain off, before the next one comes on. -4. The washing must be continued until the purity of the last runnings is proved by direct testing. No calculation of the attenuation reached can be relied on implicitly, although it is valuable for preliminary guidance, and may be the only method

¹ Classen: Quantitative Analyse durch Electrolyse. [2nd ed. Berlin, 1886.]

available; -5. The muer should no the funnel, the filter with pp., must be weighed between a closely; it should be smaller than the funnel, and not much larger than is necessary for the convenient accommodation of the pp.

Many pps. run through the paper as soon as the wash-water becomes nearly pure; bisulphide of tin exhibits this property in a marked degree. Addition of some suitable salt (sal-ammoniac, acctate of ammonia, &c.) to the washwater often helps one over this difficulty.

In the case of slimy or gelatinous pps. (e.g. hydroxides of silicon, aluminium, and chromium) Bunsen's method of quick filtration is employed. It consists in this, that the funnel is made to communicate, by its stem, with a vessel in which a partial vacuum of adequate strength is maintained by means of an aspirator (a Sprengel pump wrought with water, or equivalent arrangement). To protect the filter from being torn by the pressure of the atmosphere, its open end is supported by a small cone of platinum foil, resting on the bottom of the funnel. In regard to the operations subsequent to filtration and washing, pps. may be classified as follows: -

A. Such as stand calcination in a platinnm or porcelain crucible, and when thus treated assume a definite composition. In this case the general modus oparandi is as follows: the pp. is dried in the funuel; it is then detached as completely as possible from the paper, and put into the tared crucible. The filter, with adhering particles of pp. is folded up into a narrow strip, and this is rolled up tightly into a parcel, so that the part stained with the pp. is in the core. A platinum wire is then wound round two or three times, and the parcel is kindled in a gas flame and allowed to burn, the surplus wire eerving as a handle. After the combustion has gone as far as it will spontaneously, the residual charcoal is burned away by applying the outer portion of the flame of a Bunsen. The ash is dropped into the crucible and calcined along with the pp. In some cases, as for instance in that of alumina, it is better not to detach the pp. from the filter, but simply to fold up the pp. in the filter, and heat the whole in a platinum crucible. Any deposit of charcoal formed on the lid or crucible sides is easily removed by heating the respective part while a shield of platinum foil is stretched over the deposit. The charcoal vanishes almost instantaneously. The weight of the filter-ash must of course be ascertained by a blank experiment, and allowed for. The correction (cateris paribus) is proportional to the superficial area of the filter; i.e. ashweight = cr^2 , where c is a constant which can be determined once for all.

It is to be observed, however, that even with the same filter-paper, c depends on the nature of the liquid which passed through the filter. It is less for dilute mineral acid, for instance, than for pure water, or salt solutions followed

by water.

B. Precipitates which do not stand calcination; but assume a definite composition when dried at a suitable lower temperature, say at 100° or 120°C. Such pps. are collected on filters (previously dried at the respective temperatures) and weighed in the filters. As filter-paper is hygroscopic, the empty filter, and couple of closely fitting watch-glasses held

together by a suitable clip.

C. Precipitates which demand some supplementary chemical treatment to become fit for the balance. In regard to these it is difficult to make general statements; suffice it to say that certain metallic sulphides assume a definite composition when strongly heated (repeatedly, and until constant in weight) with sulphur in hydrogen gas. The sulphides of copper, manganese, zinc, lead, may be quoted as examples. The resulting definite sulphides arc Cu2S, MnS, ZnS, PbS, respectively.

OAS ANALYSIS.

A large supply of homogeneous gas may be dealt with, analytically, in a variety of ways. With a small gas-sample given for analysis only one mode of treatment could be-or at any rate ever is -thought of. We must collect our gas over mercury, or some other suitable liquid, and learn what we can concerning it by applying physical or chemical reactions, involving changes of gas-volume; we must measure the gas volumes involved as the only practicable mode of defining the respective masses.

Principles of volumetric gasemetry. measure a given quantity of gas means to determino its volume, v, and its pressure, P, at a definite temperature, t. In any fluid body of known nature the three quantities conjointly define the mass; yet the method is confined to gases, because in these only is the evidence afforded by the three numbers condensible into one numerical statement by more calculation.

Practical gasometry knows of no pressure greater than two atmospheres (indeed pressures above one atmosphere are exceptional); and of

no temperature below 0°C.

Within this range of conditions the law of interdependence between volume, temperature, and pressure, in all gases is in approximate accordance-in the so-called permanent gases it is in perfect accordance -with the equation

$$\frac{\mathbf{v}_{\mathbf{P}}}{\mathbf{T}} = \mathbf{Q}$$
 (1) where \mathbf{T} may be defined as $\mathbf{T} = 273 + t$. \mathbf{Q} is a con-

stant which depends only on the nature of the gas and its mass, and consequently, in reference to any named species, measures the quantity.

Condensible gases and vapours deviate from the law embodied in eq. (1) to a greater or less extent, but always in this sense that the true relations are expressible by an equation of tho form

$$\frac{\mathrm{VP}}{\mathrm{T}} \left(1 + \epsilon \right) = \mathsf{Q} \qquad . \qquad . \qquad . \qquad . \qquad . \qquad (2)$$

where e is an inherently positive number which is a function of r and r, to the effect that, for any given species, e is the less the further the pressure and temperature remove the gas from the state of saturated vapour. Gasometrically speaking e is mainly a function of temperature which runs pretty much like

 $\epsilon = \text{const.} \frac{1}{T}$; not by any means exactly so, but we are safe in saying that for overy gas-species there is a certain temperature To above which this species is, as the phrase goes, a 'perfect gas, in as least this sense that e is less than the unavoidable error involved in the experimental determination of PV+T by the oustomary instruments. In this sense our equation (1) is true for all kinds of gas or vapour without excoption.

The constant q is obviously susceptible of a number of definitions. One definition is to call it the volume which the gas assumes when $T=1^{\circ}$ and P=1 (say 1mm.), or rather the volume which the gas would assume if it were a perfect gas down to $\tau = 1$ or $t = -272^{\circ}$. To eliminate this fiction, let us view v as a function, not of P

and T, but of
$$\frac{T}{p}$$
, thua:

$$\nabla = Q\left(\frac{\mathbf{T}}{P}\right);$$

and, taking 'disgregation' as a name for this ratio T: P, define Q as that volume which the gas assumes whenever the disgregation is unity through r being equal to r numerically. If, for instance.

$$P = Q \frac{T}{v}$$

hence a may be called the particular pressure which the gas assumes whenever T = V; i.e. for $r = 1^{\circ}$ and v = 1 unit; $r = 500^{\circ}$ and v = 500, &c.

For the purely comparative measurement of two or more gas quantities, only one of the three variables need actually be measured.

Assuming the os for the gases I, II, III...to bo Q' Q" Q", &c. wo may (1) keep T and P at constant (though perhaps unknown) values and measure the volumes

$$\frac{\mathbf{v}'}{\mathbf{v}}$$
 $\frac{\mathbf{v}''}{\mathbf{v}}$ $\frac{\mathbf{v}'''}{\mathbf{v}}$ &o. which are $\frac{\mathbf{q}'\mathbf{T}}{\mathbf{P}}$ $\frac{\mathbf{q}'''\mathbf{T}}{\mathbf{P}}$ &o.

The constant factor disappears in the ratios. This used to be, at least by intention, the method of comparative gasometry.

(2) We may keep T and v constant and measure the pressures P' P" P" which are

$$Q' = \frac{T}{T} \qquad Q'' = \frac{T}{T} \qquad Q''' = \frac{T}{T} & dec.$$

 $q'\frac{T}{v} - q''\frac{T}{v} - q'''\frac{T}{v} & c.$ and consequently again measure the Qs (Regnault's method).

(3) We may allow T and P to vary, but keep their ratio, the disgregation, constant, and measure the volumes, i.e. substitute for the QS

$$(Q' \quad Q'' \quad Q'''...) \times a \text{ constant } \frac{T}{P}$$

(Doyère's method).

But q has an important chemical significance. A glanco at eq. (1) shows that the specific gravity of a gas, meaning the number of timee its weight is greater than that of the same volume of some standard gas of the same diegregation, ie independent of T and P. As stated hy Avogadro, and since proved by numerous experiments, we have for any set of pas-species e'', e''' ... = M' : M'' : M'''.....

where m is the mol. weight.

e = const. M

Hence supposing, at a given disgregation unit volume of standard gas weighs a unite, ther nuit vol. of another gas of the mol. w., x weighe $n \frac{M}{M_0}$, where M_0 refers to the standard gas Hence unit-volume of any gas, if measured at that disgregation, contains $\frac{R}{M_a} \times M$ units of weight of its substance; hence oqual volumes of any two contain the same number of molecules, where 'molecule' may have the usual meaning given to this term. Hence our constant q, or any of its substitutes as given above under (1) (2) and (3), in a relative sense counts the molecules of the

respective gas. Eq. (1) tells us nothing about the relation between the volumo v of a gas-mixture, and the volumes v' v'' v'''...of its components; but we know, by direct experience, that v = v' + v'' + v'''...; hence Avogadro's law holds for mixed as well as for homogeneous gases; and, independently of it, we have

$$q' + q'' + q''' \dots = Q$$
. . . . (4)
and at any constant value of

$$y+T$$
 $p'+p''+p'''...=p$ (5)
(where the small letters refer to the components and the large ones to the mixture). And so quite generally

Hence our customary mode of stating the composition of a gas-mixture is susceptible of three readings. Instead of saying (1) 100 volumes of air contain 21 vol. of oxygen and 79 of nitrogen; we may say (2) the partial pressure of the oxygen is 21 p.c. and that of the nitrogen 79 p.o. of the total pressure of the air; or (3) every $n \times 100$ mols, of air contain $n \times 21$ mols, of oxygen and $n \times 79$ of nitrogen.

In the more easily condensible gases, the number & (which might be called the measure of gascoue imperfection) assumes appreciable values at the ordinary temperature; yet in the ordinary practice of gas analysis even these gases are eustomarily being measured at, or near, the temperature of the laboratory. To give an idea of the possible value of the error thus neglected we will take up the case of carbonic acid, which, of ordinarily occurring gases, is perhaps the most imperfect.

According to Amagat, carbonic acid, from 0°C. npwards, expands at a greater rate than air, np to about 200°, whence onward it behaves like a perfect gas in ref. ence to expansion caused by changes of temperature or pressure. At 760° mm. its expansion from 0° to 200° is in the ratio \$ of 1:1.74065. Hence supposing we find for a quantity of carbonio acid $v = v_0$, for r = 273 and r = 760, we have for the constant q : -

(1) By the ordinary routine mode of calculation, i.e. by eq. (1); $q' = \frac{v_0 \times 760}{278}$

' It is worth while to note that this all-important proosition has never been looked into in the Regnaultfashion.

"Calculated by the writer from the coefficients of empansion stated by Amagat for 50°, 100°, 180°, and 200°.

(2) For the true q; $q_0 = \frac{v_0 \times 1.74065 \times 760}{1.74065 \times 760}$

Whence $q_0 = 1.0046 \, q'$; or in the sense of our equation (2), for $\tau = 273$ and r = 760 nm.

$$\Delta Q_0 = \frac{v_0 \times 760}{277} (1 + \epsilon)$$
; and $\epsilon = 0.0016$. (7)

This number, or say 0.005, might perhaps be put down as the maximum value which e may assume in the enstomary mode of measuring gases proper, were it not for the following consideration. As a rule the gas to be measured is contaminated with vapour of water, and it is the Q of the dry gas that is wanted. One mode of obtaining it is to remove the water by chemical absorbents and to measure the dry gas; but this is a tedious process; hence we prefer, in practice, to saturate the gas completely with water, to measure it in this condition, and, before calculating by eq. (1), to correct the observed pressure by deducting the maximum steam-pressure at the respective temperature, as determined by Magnus and by Reguault for the vacuum, assuming the corrected value $\mathbf{r}_0 = \mathbf{r} - \mathbf{\pi}$ to represent the pressure which the gas would exhibit at the same vol. and temperature if it were dry. As shown by Regnault, this is not quite exactly the case, yet if # is small, i.e. if the temperature is tow, the error may be neglected. A low temperature, it is true, means a relatively great e, but π certainly, and the error in π probably, increases (with t) much faster than ϵ decreases.

Both the authorities named give their ws in terms of the pressure of a column of mereury of 0°C, whose height equals 1 mm. Hence to bo able to use their numbers directly we must provide our eudiometers and barometer with true mm.-scales. And we ought to reduce all mercury columns (measured as pressures) to 0°C. This, however, is necessary only in the case of absolute measurements, i.e. if we measure a gas as a step towards calculating its weight; for relative measurements we may choose our units for v, T, and r, at pleasure, hence the absolute magnitude of our 'mm.' is of no consequence. Nor is it necessary to reduce the # to what our mm. is at the respective temperature, because the correction is practically irrelevant.

Gases like hydrochloric acid, ammonia, sulphur dioxide, &c., must be measured dry-

for an obvious reason.

(a) Proximate. Gas-Analysis. For the proximate analysis of a gas-nuxture we have only one direct method. After having measured off a convenient sample, we withdraw the several components (singly or in groups), by the successive application of appropriate chemical absorbents, as pressureless solids or liquids, and, after each absorption, we measure the gas-residue left. Supposing the sample measures v units at T and P, and the same, minus component I, measures v' units at r' and r'; we have for the sample $\mathbf{q} = \frac{\mathbf{v}\mathbf{p}}{\mathbf{r}}$, for the residuc $\mathbf{q}' = \frac{\mathbf{v}'\mathbf{p}'}{\mathbf{r}'}$; hence for the

percentage of 1; $x = \frac{Q - Q_1}{Q} \times 100$.

To show the possibilities of the method, we

enumerate the most important reagents and state the powers of each as an absorbent. (1) Water (as such or as Na, SO, 10H, O)

absorbs HCl, HBr, HI, very promptly.

(2) Solid DRY caustic potash absorbs water very completely; acid gases generally more or less slowly.

(3) Solid Moist caustic potash absorbs all acid gases (CO2, SO2, II2S, HCl, &c.) very readily.

(4) Caustic potash solution acts like (3) and (1).

(5) Dilute sulphuric acid absorbs all alkaline gases (NII3, CII3NH2 &c.); besides acting as

(6) Oil of vitriol (II₂SO₄ + ¹/₁₂II₂O) absorbs (a) water, alcohol, ether, methyl-oxide, very readily; (b) propylene and higher homologues, with a fair degree of promptitude. CaH4 is absorbed only on long-continued shaking (Berthelot).

(7) Sulphuric anhydride in H.SO, absorbs C.H. in addition to the gases named in (6).

(8) Bromine (over water in diffused daylight) acts pretty much like (7); the excess of Br vapours left is removed by means of KHOAq. (9) Pyrogallic acid in caustic potash ley absorbs oxygen abundantly and promptly (Liebig), besides acting like (4).

(10) Cuprous chloride in hydrochloric acid absorbs oxygen; also CO, C2H2, C4H4 (Bertholet).

Spoils the mercury.

(11) Same reagent in aqueous ammonia acts like (10), and besides absorbs certain other gases, e.g. all the olefines (Berthelot).

(12) Ferrous sulphate in concentrated solution absorbs nitric oxide; but hardly in the chomical sense, as the compound has a measurable dissociation-pressure.

(13) Binoxide of manganese, as compressed powder, is used by Bunsen for absorbing H₂S and SO, -Solution of CrO, or of KMnO, acts similarly and more promptly.

(14) Chromous sulphate in NH3 and NH1Cl solution absorbs O,NO,C2H2,C31I4, but does not

act on CO, C.H., or C.H. (Berthelot).

That all gas mixtures cannot be analysed by means of these 14 reagents is obvious. Unfortunately they are all group-reagents, and a group when once absorbed is not susceptible (practically) of further gasometric analysis. One or other of the absorbed components may be determinable otherwise-thus for instance H.S (absorbed in KHO) by titration with iodinebut these are rare exceptions. For the analysis of a gas-mixture which, with regard to chemical absorbents, behaves as a whole, only two methods are at our disposal; one is to determino the ultimate composition of the gas (if possible), and from the results to try and arrive at the proximate composition; the other is to examine the gas by means of physical absorbents. But to obtain definite results with these we must follow the lead of Bansen, and both contrive their application and interpret the results, in the light of the laws of gas-absorption.

Analysis by physical absorbents.

Imagine v volumes of a mixture of the unitary gases I., II., III., . . . to be shut up

NO is absorbable also by the conjoint action of O and KHO solution, as KNO, and KNO.

¹ I.e. we may, if we choose, measure our is with a Fahrenhelt thermometer and take T as being T=4594+6 (in F. degrees).

in a close vessel over h volumes of water or alcohol, an impervious diaphragm separating the two. As soon as the diaphragm is removed, the gas and liquid exchange molecules, and this goes on for ever; but if a constant temperature t is maintained, a point is reached, cooner or later, at which the changes of composition, exactly compensate each other, so that matters are the same as if the exchange had come to a etop. This point of dynamical equilibrium is reached almost instantaneously on violent shaking. The final result is that the gas-space v is saturated with the vapeur of the liquid, while a quantity q of each of the components of the gas is held in solution by the h volumes of liquid. This quantity q at a given temperature is in (more or less exact) accordance with the cquation -

$$q = h\beta\pi$$
 (8)

where π means the partial pressure of the respective component in the residue, and β is a constant, which may be defined as being the value which q assumes when h=1 and $\pi=1$ mm. q and β are, of course, of the came denomination; if q means mgms. β means mgms. likewise. But we will assume q to be measured by volume at 0° °C. (or T = 273° °C.) and p=1 mm., and on the basis of this assumption (with Bunsen) call β the 'co-efficient of absorption.'

Our equation has been tested experimentally only with water and, in a more limited sense, alcohol, as a solvent; and in reference to either, it may be assumed to hold, at pressures up to about 1 atm., and temperatures from 0° to about 30° C., for all gaeee which, under the circumstances, do not act chemically on, or dissolve very abundantly in, the respective liquid. With a given gas-species, β , in general, increases when the temperature falls, or when alcohol is substituted for water. It has, in general, different values for different species of gas. Hence we at once see our way towards distinguishing a unitary gas from a mixture. Take, for instance, the case of marsh-gas CH, as against a mixture of equal volumes of II. and C2H6 = CII4 per 1 volume. With alcohol as an absorbent, the \$ of C.H. is far greater than that for H2. Hence, if the mixture be dissolved partially by alcohol, the residue will contain lese earbon per unit volume than {C, and similarly in eimilar cases.

The relation between the composition of the mixture operated upon and that of the unabsorbed residue is easily formulated. Let m', m'', &c., stand for the quantities of the several components present in unit volume of the original gas, and let n', n'', &c., have a similar meaning in reference to the residue; let r stand for the (dry) pressure of the original gas, and p for that of the residue, then we have for any one of the components $q = l \beta(pn)$; and for the unabsorbed part of that component

$$r = (np)v \frac{273}{273 + t} = (np)''v_e'',$$

and $q+r=pn(v_o+\beta h)$; but $q+r=rnv_o$, hence $pnv_o=pn(v_o+\beta h)$, which enables us to calculate the 'n' of a named component from its 'm.' For further developments we refer to Dittmar's 'Exercises in Quantitative Analysis,' esction on gas analysis (Glasgow, W. Hodge & Co.). Without mathematics it is clear that the quantity, λ ,

of total gas absorbed, reduced to p=1 and h=1,

$$\lambda = \frac{v_0(P-p)}{hp}.$$

In the case of a unitary species λ is the coefficient of absorption, and is consequently constant, while, in the case of a mixture it varies (in general) with h.v., i.e. with varying quantities of water for the same quantity of gas started with. Hence an obvious second method for testing a gas for chemical oneness.

Of general methods of gas analysis, only one remains to be considered. We refer to

Method of Combustion.—A method of ultimate analysis which presumes that the gas to be analysed is in, or by addition of hydrogen or of oxygen or of either plue fulminating gas, can be brought into, such a condition, that the mixture, when fired with an electric spark is resolved entirely into (in general) carbonic acid, nitrogen, and water, and excess of either hydregen or exygen as the case may be. The method consists in this that a measured velume of the given gas is exploded, and the gas quantities involved are measured as far as necessary te enable one to calculate the elementary composition of the gas under operatien, the results being regarded enstomarily in volumes (reduced to some tacitly assumed constant disgregation) of the respective elementary substances. For uniformity's sake this system. is extended even to the carbon, one volume of earbon being used as a phrase for the quantity of carbon centained in twe velumes of carbonic anhydride. This mode of reporting comes to the same as stating the quantities of hydrogen, oxygen, &c., as multiples of the molecular weights H₂, O₂, N₂, and of the double atom C₂ of carbon. In the ecquel we sometimes nso H.,O., N., CO., CO, as symbols for '1 volume.' When in a calculation we have to refer to a certain (reduced) volume of carbonic acid we designate it by the letter K; in a similar sense s refers to oxygen; w to water vapour; \$\mathstreeta\$ to nitrogen ('n' is reserved for the atom); o to contraction. The following examples explain the method:

I. The gas is a mixture of hydrogen and hydrocarbons; i.e. 1 vol. = aC_wβH_.. We determine the following gas-quantities:

- (0) The volume of the sample, as . v
 (1) , , plus
 - added oxygen, as . . . ▼₁

 And after firing
- (2) The volume of the total product measured cold, as
- (3) The velume of the residue left after removal of the carbonic anhydride, as

The quantity of earbonic anhydride produced in the combustion is ' $\kappa' = v_2 - v_3$, whence $\alpha = \frac{1}{2} \frac{v_2 - v_3}{v}$.

The hydrogen is calculated from the 'contraction,' meaning the difference 'o'= v_1-v_2 , thus: $v_1=v+s$, where s stands for the added oxygen.

 $v_2 = \kappa + \text{oxygen left unburnt, which is } s - \kappa - x$,

¹ The mixture H₂+½O₆ obtained in the electrolysis of water.

where a means the oxygen which converted the hydrogen into water. Hence c = V1 - V2= $r + s - [\kappa + (s - \kappa - x)]$ or $c = \nabla + x$; hence t=c-v, and $\beta=\frac{2(c-v)}{c}$

The sum $a + \beta$ is, of course, always greater than unity unless $\alpha = 0$.

We will assume now

II. That the gas contains (in the v units taken for analysis) z volumes of free oxygen and y volumes of free nitrogen beside v, of hydrocarbons; both s and y being unknown. Here we at once see that the measurements of ∇_1 , ∇_2 , ∇_3 , de net enable us to calculate z or y. But we cannot even calculate the volume x of oxygen which combined with the hydrogen in the combustion; because from case I. we see that $x = c - v_0$, and v_0 is unknown. Nor does a direct determination of the oxygen-residue s, in v, help us, because s, is a function of c, independent of g and s. We have, in fact :-

$$s_r = z + s - x - K;$$

$$0 = V - (z + y) + x;$$

$$s_r + 0 = s - K + V - y;$$

 $\mathbf{s}_r = -\mathbf{o} + \mathbf{s} - \mathbf{K} + \nabla - \mathbf{y}.$ and The determination could only confirm this calculation. If s is known to be = 0, or s and yconjointly are known to be so much air, the problem becomes easy of solution.

III. The gas is aC...\$H...\gammaO...\delta N...=1 volume; states of combination unknown. If we add to the values (for v of substance) of k and c that of the nitrogen in the ultimate residue (lct its quantity be = Ω) we have α and δ at once. But, (even supposing we did not care for γ), to determine & we must measure the quantity, w, of steam produced in the combustion. Frem

w we have $\beta = \frac{1}{v}$ w; and from this, and the contraction c, we can calculate \gamma thus; let so denote the quantity of oxygen which, conjointly with the oxygen in the substance, is just sufficient te burn the substance into CO2 H2O, and N2, and let sr be the surplus added, so that s, + s, = s; we have

$$\begin{aligned} & \nabla_1 = \nabla + S_0 + S_1 \\ & \nabla_2 = \oint 2 + K + S_e \\ & C = \nabla + S_0 - \oint 2 - K; \\ & \nabla_1 - \nabla_3 = \nabla + S_0 - \oint 2. \end{aligned}$$

Now, it was obviously the oxygen sum s₀ + vγ which produced the H₂O and CO₂; hence,

$$S_0 + \nabla \gamma = K + \frac{1}{2}W;$$

$$\gamma = \frac{1}{V} \left(K + \frac{1}{2}W - S_0\right).$$

Whenever, in a gas of unknown constitution, oxygen may be present, the determination of w becomes indispensable, because without it the water possibly present in a gas would escape ns altegether; we could not, for instance, discriminate between cthylene and oxide of methyl.

The case which we have just been discussing includes that of the analysis of any gas \(\gamma O_r \delta N_2 \) which is combustible by means of hydrogen. Because the added hydrogen, for calculating purposes, may be included in the 'v' of our formulæ, to be ultimately allowed for. In practice, however, the variety of proximate compositions included in the formula $\gamma O_T \delta N_2$ is very small, so that, in the case of such a gas,

we had better at once calculate the proximate components (N2, N2O, O2 dec.) directly from the data of the combustion.

IV. Let us now see how far the method of combustion goes as an indirect method of proximate analysis. Let ns assume that we have to deal with a gas of the nature pre-supposed in ease III., and that the quantities, k, c, w, A, have been determined, and none of them found =0. We also assume that we know the formulæ of all the several species I., II.,, which can possibly be present. To find the quantities of these contained in unit-quantity of the given gas (x' for I.; x" for II.) we might begin by calculating the elementary composition of our gas, i.e. the coefficients in the average formula $\alpha C_2 \cdot \beta H_2 \cdot \gamma O_2 \cdot \delta N_2 = 1$ vol., and then express these algebraically in terms of the special values a', a", B', B''....., &c., appertaining to the components I., II, &c., thus,

$$a = a'x' + a''x'' + a''x'', &c.
b = \beta'x' + \beta''x'' + \beta''x'', &c.
\gamma = \gamma'x' + \gamma''x'' + \gamma''x'', &c.
\delta = \delta'x' + \delta''x'' + \delta''x''', &c.
1 = x' + x'' + x''', &c.
V.$$

In practice, of ceurse, we need not calculate a, \$\beta\$, &c., but may at once form equations between $\frac{1}{v} = k$; $\frac{1}{v} = c$; $\frac{1}{v} = n$, &c., and the special

values k'k"..., c'c'..., n'n"..., thus-

and solve these equations; but the former set shews more clearly hew far the method goos as a method of proximate analysis.

From either set we at once see that if the number of potential components dees not exceed five, we can in general calculate the quantity of each in unit quantity of gas, i.e. x' x"...x". In general we say, because obviously if one or more of the co-efficients α , β ... is = 0, so many equations collapse; in the case, for instance, of $\gamma = 0$ and $\delta = 0$, only three equations are left. And (to adhere to the example) if it should happen that all the values of B are the same function of the respective values a, then equation II., or, if you prefer it, equation I., is lost, and only the case of two compenents is susceptible of a selution. A similar result occurs if all the components should happen to contain the same number of hydrogen-atoms (or the same number of carben-atoms) per melecule. Supposing, fer instance, all the components were of the general formula Call, then & would by necessity be = 3, and equation II. would be resolved into 3 = 3x' + 3x'' + 3x'''... which is a more repetition of equation V. And similarly, if all the components were di-carbon gases, equation I. would become useless.

The general rule is, first of all to find out how many of the quantities k, c, n, w... in addition to our knowledge of the constitution of the gas, we should need to calculate the co-efficients α, β... of the average formula. Surposing 4, 8, 2 suffice, then (in general) 3, 2, I, (but not any 3, 2, 1), equations of the second set, taken along with equation V ..., will suffice to find the unknown quantities w, w... sought, provided their number does not exceed 4, 8, 2.

For examples see the writer's Tables to facilitate chemical calculations (Williams & Norgate).

The following table gives the values of c, k, s, w for several gases.

I .- Combustible by Oxygen.

	<u> </u>				
	6	k	20	10	n
Hydrogen, H	1.5	0	0.5	1.	0.
Carbonio oxide, CO	0.5	1	0.5	0.	0.
Methyl-aldehyde, CH.O	1·	1	1.	1.	0.
Ammonia, NH,	1.25	0	0.75	1.5	0.5
Methylamine, CH,N .	1.75	1	2.25	2.5	0.5
Cyanogen, N.C.	0.	2	2.	0.	1.
Hydrocyanic acid, NCH	0.75	1	1.25	0.5	0.5
Marsh gas, CH	2.	1	2.	2.	0.
Acetylene, C.H.	1.5	2	2.5	1.	Ö٠
Ethyleno, C ₂ H.	2.	2	3.	2.	٥٠
Ethanc, C.H.	2.5	`2	3.5	g.	0.
Propylene, C.H.	2.5	. 3	4.5	ğ.	0.
Propane, C.H.	3.	' 3	5.	4.	ō.
Oxide of mothyl, C.H.O.	2.	. 2	3.	3.	0
Benzene, CaHa	2.5	6	7.5	3.	0.
1 vol. = $C_{\alpha}H_{\beta}$	$1+\frac{\beta}{4}$	a	$\alpha + \frac{3}{4}$	0.5β	0.
				. '	•

II .-- Combustible by Hydrogen.*

			c	h	w	n
Nitrous oxide, N ₂ O Nitric oxide, NO† .	:	:	1· 1·5	1.	1· 1·	1.

The Practice of Gas Analysis.

In this section we take cognisance only of the chemical methods, and in regard to these confine ourselves in the main to those apparatuses in which mercury serves as a trapping fluid.

Taking ordinary laboratory appliances for granted, all that gas analysis demands of special apparatus is: a barometer, a pneumatic trough with transparent sides, and a series of glass tubes, closed at one end and open at the other, and provided, virtually, with two scales, of which one divides the gas capacity, and the other the axis, into units of sufficient smallness. One or more of those tubes must be provided near the closed cud with a couple of fused-in platinum wires so that a combustible gas-mixture in it may be exploded by means of an electric spark. The possibility of obtaining exact results by means of these simple contrivauces is proved by the fact that all the great gasometric work of Cavendish and Gay-Lussac, which laid the foundations for our present chemistry, was done with apparatus like those referred to, or even with apparatus of a lower order of complexity. Of course to obtain exact results we must be alivo to all the numerous sources of error involved, and eliminate them as far as possible experimentally or otherwise. It is one of the

• h=hydregen necessary for combustion.

† Nitrio oxide cannot be burned with H, alone; it requires addition of a certain proportion of N,O; and even then the combustion is irregular (Bunsen, Uas. Meth. Bad Ed. pp. 98, 96).

merits of Bunsen to have done this for us, and to have thus brought the old method of gas analysis into a form which, on the score of preoision at least, leaves nothing to be desired.

Bunsen's Apparatus and Methods. The first requisite of exact gas analysis, Bunsen says, is a special room in which the temperaturo is subject to only slight, and to no sudden, variations. The ideal gas-room forms part of a substantial building; it is not warmed artificially nor is it contiguous to any other room thus heated; and its windows face the North, to keep out the sun. In such a room the temperature during a working day remains constant as a rule to within 1°C. although the variations of temperature of the outside air may amount to as much as 12°C. A characteristic of Bunsen's method is that the chemical treatment of a gas is effected in the tube in which it has been measured; but he uses two kinds of tubes, one for the absorptions, the other (cudiometers) for the combustions. Both are about 20 mm. wide (inside measurement; in narrower tubes the capillarity assumes measurable values) and 2 mm. or so strong in the body, which strength suffices even for the eudiometers. The absorption tubes are about 250 mm. long, and are provided with sponts, so that a gas contained in one can be transferred to another tube by laying down the absorption tube in the trough. In the caso of the cudiometers a length of 500-600 mm. suffices for all ordinary purposes. The platinum wires are fused in somewhere near the closed end, and are bent so that the two ends stand opposite cach other at a distance of about 2 mm. Every gas tube is provided with an ctched-in millimetre-scale, and the gas-volumes corresponding to the several marks are determined by calibration, so that each tube is a laboratory, a volumeter, and a manometer, in one. The scale is figured from the closed end downwards. calibrate a tube it is fixed, open end upwards, in a vertical position; successive, exactly equal. quantities of mercury are introduced, cach corresponding to some 20 mm. of scale, and after each such addition the exact position of the top of the meniscus in reference to the scale is observed by means of a horizontal telescope standing at a distance of 1-2 metres, and the readings are taken down, care being taken, bofore each reading, to remove any air-bells that may be imprisoned between the mercury and the sides of the tube, by means of a long stick of whalebone. The measuring off of the standard volume of mercury is effected by means of a short etout test-tube, ground exactly flat at its lipless rim, and provided with a lid of ground plate-glass. It is filled from a pipette-like reservoir provided with a long narrow outlet tubo and a stop-oock at the top end of this tube. If care bo taken so to operate that the mercury, while it fills the measure, forms one continuous mass, the forms tion of air-bells ie easily avoided. The measure. while being filled, is held in a wooden olip (not directly in the hand, which would cause the mercury to expand) while the lid is slung to the thumb of the same hand. The measure is filled to overflowing, the excess of mercury is removed by putting on the lid, and the mercury is poured into the tube.

The meroury-measure is assumed to hold

'v' volumes of mercury, v being so chesen that, for differences of capacity at least, the numerical valus of the volume corresponds as nearly as possible with the respective seale readings, so that, for small differences, every I mm. of difference of level can be assumed to correspond to unit-volume (i.e. to $\triangle v = 1$). Supposing after addition of k measures full of mercury the meniscus stands at a mm., the volume of the body of quicksilver now in the tube is kv units by definition; but the gasvolume corresponding to R is greater than kv, by the volume x of the slaped space between the meniscus as it is when the gas is being measured, and the meniscus as it was in the calibration. To determine x, we pour some corrosive sublimate solution on the meniscus (after having read off the number R in calibration) which causes the menisons to flatten out into a plane, and we read the position of this plane which stands say at R-5 mm. Counting from seme horizontal reference-plane 00 upwards, the volume of the mercury and the total space from 00 to the horizontal plane through a are constant. The volume Ax has become visible as a cylinder of the height δ millimetres, and consequently of the capacity of & 'units.' Hence the gas velume corresponding to point R is $kv + 2\delta$. From the values $kv + 2\delta$, and the corresponding readings a' R" R" &c., it is easy (though tedious) to calculate a calibration table which gives all the gas-volumes from mm. to mm. directly. In reading off with a good telescope one soon learns to divide every individual degree into tenths by the eye; the (Δv) s corresponding to them are found by interpolation from the tabular entries. Should the tube be used for measuring over water, we remove the meniscus-correction by subtracting 28 from the registered volume, and thus obtain as good an approximation to the gas-volume over water as is called for in such a case.

During the course of the calibration the temperature of the mercury must be kept as nearly as possible constant, or else the values recorded for the lower marks may be very appreciably incorrect. It is well to record the mean temperature t₀ during the period of calibration, and to determine the weight in grams of a measurefull (v'volumes') of mercury at t², in order to be prepared for reductions of gas-volume to gasweight. One gram of mercury at 0°C, occupies 007355 c.c. (log. 2.866589), and the volume at t²C, is 0.07355 (1+0.0001814 t) c.c.

To prepare a cudiometer for receiving a gas we first make it rigorously elean, and next, if the gas is meant to be measured 'moist,' attach a small drop of water to the closed end, which during the operation of filling with mercury gets flattened out and spread over the inside, and so offers a large surface to the gas. The mercury is introduced through a long funueltube (provided with a stop-cock at the bottom of the funnel) which goes to the bottom of the endiometer. By means of this arrangement it is easy, after the introduction of the first thimblefull of metal, to let the mcrcury in eudiometer and funnel form one unbroken mass, and thus to avoid formation of air-bells at the sides of the tabe.1

In regard to the sollection and preservation of gas

Assuming the gas to have been introduced, and the tube to have been fixed in a vertical position, we begin by preparing for the reading of the level of the trough by inserting a paper soreen, provided with a shaped perforation, between the moreury and the front (glass) wall of the trough, which gives a fairly distinct image of the line of intersection between scals and trough-level plane; we then suspend the thermometer somewhere close to the tube and next leave the room for a time to allow the gas to assume the temperature of the air. On returning we read off:

1. The position n of the meniscus in the tube.

2. The level of the mercury in the trough, Ber

3. The temperature, to.

4. The barometer; let its height bo=mm. This reading comes last because the barometer requires to be tapped before being read and this cannot be done from a distance.

In the vast majority of cases the temperatures t't"t"... for the soveral gases to be compared do not differ much from their mean: lichec, even if they differ considerably from the temperature which prevailed in the calibration, the value furnished by the calibration table for R can be put down as the correct relativo volume of the gas measured; and the pressure of any mercury-column measured may be identified with its nominal height in mins. as read. Hence we have for the pressure of the dry gas at the observed volume P= $B + R - (a_0 + \pi)$ where π is the maximum pressure of steam at to, and for the gas-quantity (the volume reduced to unit disgregation) $v_0 = \frac{v_P}{273 + t}$ (see theoretical part).

Bunson prefers reducing to 0°C. and 1000 mm. pressure by the formula

$$v_0 = \frac{v_F}{1000(1 + 0.003665t)}$$

which, if a table of the logarithms of all the values $(1+\alpha t)$ is at hand, is as short a method as the one recommended by us.

Corrections of tuhe-capacities and mercuryheights for variations of temperature occur only in the rare case when one of the gases concerned in the analysis was measured at an artificially established high temperaturs t. In this case the value v furnished by the calibration table for the reading a must be corrected thus:—

(True capacity down to h) = $v[1+\lambda(t-t_0)]$ where λ stands for the coefficient of the cubical expansion of glass, and may be put down at $27^{\circ}6 \times 10^{\circ}$. And for the observed height h of a mercury column measured at a high temperature t we must substitute the height h_0 of the equivalent column of mercury of t_0 degrees. t_0 stands in both cases for the average temperature that provailed during the determinations made in the ordinary manner. Obviously $h_0 = \frac{h(1+kt_0)}{1+kt}$ and with sufficient exactitude.

$$h_0 = h \left[1 - k \left(t - t_0 \right) \right]$$

samples, and the mode of introducing a sample into the cullometer, we refer to Burson's Gasomertiche Methoden, sesond edition, Braunschweig, 1877. The first edition, 1857, was transinted into English by Roscoe, and published by Walton and Maberley, London. k = 00018.—Strictly speaking the nominal value i L of a piece of millimetre-ecale as measured at

t should be corrected thus:

(True length at t) = $L(1 + (t - t_0)9.2 \times 10^{-6})$. (It is easier to remember that 1000 mm. expand by 0.92 mm. per 100° of increase of temperature.) But our work must be very exact to be worth this correction. It is more relevant to state that whenever we wish to make use of Regnault's determinations of absolute gasdensities we must measure by his unit of (temperature and) pressure, and consequently reduce our mercury-columns to true min. of mercury of 0°C. Regnault's densities D, on the other hand, ought to be reduced to the gravity of the place of observation; this correction, however, may as a rulo be neglected.

For the execution of an absorption the most obvious method is to shake the gas with the respective reagent in the liquid form, and to measure the gas-residue as it stands over the layer of liquid reagent. But this method is in general attended with a number of obvious grave errors, and, besides, does not readily adapt itself to the successive application of different reagents. To overcome these difficulties Bunsen, as a general rule, uses all the absorbents in the form of solid or semi-solid balls, fixed each to the end of a platinum wire. Caustic potash, chloride of calcium, &c., are east in a bulletmould around the coiled-up end of the wire. To bring sulphuric acid, alkaline pyrogallatesolution, and other intrinsically liquid reagents into a quasi-solid form, a ball of some suitable porous material-battery charcoal for vitriol; papier-maché for pyrogallate, &c .- is fixed to the end of the wire and the ball is then soaked in the respective liquid. In this manner it is quite possible to accomplish an absorption even with oil of vitriol, without soiling the tube or the mercury to an inconvenient degree. Reagent vapours left after an absorption, or foreign vapours produced by the reagent-e.g. the SO, and SO2 which are always left after an absorption of olefines by funning vitriol-must of course be removed by suitable reagents (SO3 and SO, hy a soft potash ball) before the residue is measured. As small remnants of, for instance, KHO, remain unavoidably in the tube, the residues must in general be measured dry, because the pressure of water in the pressure of moist KHŌ is incalculable.

The weak point in Bunsen's method is that it is tedious, and that it does not enable one to see the end of an absorption otherwise than by the repetition of the process with a fresh reagent ball. Bunsen himself has indeed come to effect carbonic acid absorption, by shaking the gas with solution of caustic soda, and measuring tho gas-residue over the layer of reagent. To be able to correct for the pressure of this layer and for the vapour-pressure of the feagent, he employs it in the form of a standardised solution containing exactly 7 p.c. of NaOII, which has a practically constant specific gravity. He has also determined the course of the pressure-curve by standard experiments; the results are embodied in a table appended to his Gasometrische Methoden, second edition.

In this connection we must refer to an ingenious method devised by Russell (C. J. [2]

6). He introduces the reagents as solutions by means of a graduated syringe; and after they have done their work, removes them by means of a ball of cotton-wool, previously rendered airfree by kneading it under mercury. To remove what adheres to the tube and mercury he rinses the inside with some injected water and removes this by a fresh cotton-wool plug.

In the analysis of a gas by combustion & necessary preliminary step is to remove (and determine) what there may be of SO, CO, NH, and similar gases, by suitable absorbents. Part of the residue is transferred to the eudiometer and measured. Let its volume (reduced to, say, unit disgregation) be equal to v units. The necessary quantity of oxygen or hydrogen is now added and its quantity is determined by measuring the mixture (let its red. volume be v'). The mixture is now rendered explosive, if necessary, by adding the requisite proportion of fulminating gas, the whole is well mixed and prepared for explosion by pressing the open end of the endiometer firmly against an india-rubber pad lying on the bottom of the trough. The upper surface of the pad must have been rendered air-free by rubbing it over with a few drops of corrosive sublimate and mercury. After these preliminaries the combustion is effected by passing an electric spark through the mixture. After the combustion, the cudiometer is carefully lifted from its cushion, so that the mercury enters slowly and without drawing in air. Tho gas, after having been allowed to cool down to the temperature of the room, is measured, to determine its reduced volume v". From the data mine resolutioned so far, we ... unit of original gas; $c = \frac{1}{v} \ (v' - v'').$ obtained so far, we have for the contraction per

After this determination comes, if necessary, that of the water produced, which of course is practicable only if the original gas and the added oxygen were used in the state of perfect dryness and any added fulminating gas measured exactly. To determine the water-of which part in general separates out in the liquid form -the endiometer is lifted out of the trough by means of a small heaker, and with it, as its temporary trough, placed within a glass cylinder through which a current of steam can be passed to raise the temperature of the whole to something like 100°C. The exact temperature t'' is noted down. If care be taken to arrange matters so that the pressure of the gas mixture produced is not more than 0.5-0.6 atmospheres the steam may be practically regarded as a perfect gas, so that the measurement of the mixture enables on to calculate its quantity. If the red. volume of the mixture be v", we have for the steam per unit of original gas;

$$w = \frac{1}{v} (v''' - v'').$$

In this measurement the corrections for the expansion of the glass and mercury, which were referred to above, necessarily come in.

The determination of the carbonic anhydride produced is effected by caustic potash. In an aliquot part of the residue, the surplus-oxygen (or hydrogen if we have to deal with a gas combustible by hydrogen) is determined, if neces

Oxygen can be determined by explosion with excess of hydrogen (its quantity is i of the contraction), or it may be determined by absorption with pyrogallate; hydrogen is determined by explosion with oxcess of oxygen, 3 of the contraction is the volume of the hydrogen. The nitrogen is found by difference. The method of combustion—as a method of ultimate analysis at least-is susceptible of a high degree of precision, which, however, is attained only if wo take carc to avoid its numerous sources of error.

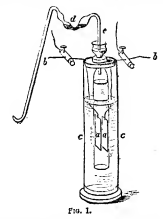
I. The reagents used must be absolutely pure, which of course includes absence of air; hence in any case the gas-evolution apparatus employed should be no larger than is absolutely necessary, so that the air space is reduced to its

minimum.

Pure oxygen is easily made. A few grams of pure potassium chlorate are introduced into a little bulb blown to the end of a glass tube, and the latter is then drawn out and bent into the form of a gas-delivery tube. The rest needs no

explanation.

Pure fulminating gas is best produced electrolytically from 10 per cent. pure sulphurio acid. The two elements are sure to be produced in the exact ratio of H2: 202, but whether the gas as it comes off really has this composition depends on the observance of certain conditions which cannot be formulated better than by a description of Bunsen's apparatus (Fig. 1).



The decomposition cell consists of a cylindrical bottle provided with fnsed-in platinum electrodes aa, and terminating in a funnel; it is filled with the acid up to about this of its capacity. The end e of the washing-balbs and deliverytube is ground into the neck of the funnel; a few drops of acid poured over the joint make it absolutely tight. The bulbs d are charged with a few drops of oil of vitriol to dry the gas evolved. The bottle is suspended within a bath of water c c (or alcohol to avoid its freezing in winter-time). To produce a current of fulminating gas, the wire ends bb are connected with the poles of a battery of four 'Grove' or 'Bunsen' cells, and the gas evolved during the first five minutes is allowed to escape in order to expel the air, and to establish absorptio if it be nroduced—at. say, 1 atm. pressure—by

metric equilibrium between the gas above, and the gas held in solution by, the soid. As oxygen has a greater coefficient of absorption (8') than hydrogen (β'') , the first portions of gas that come off contain an excess of hydrogen. Besides, the ratio &': B" varies with the temperature; for this roason, and also to avoid undue heating of the conducting-wires, the bath is usod.

Imagino the apparatus to be so modified that the oxygen electrode is immersed in a mass of liquid zinc-amalgam, which takes up the oxygen as quickly as it is liberated from water, and you have Bunsen's apparatus for producing pure hydrogen. But a sufficiently pure gas for most purposes can be obtained in the ordinary manner, namely, by the action of 10 p.c. (pure) sulphuric acid on pure zinc, in the presence of platinum, within a small, narrow-necked, flask. The hydrogen thus evolved is filtered through a short narrow tube full of fragments of caustic potash to remove traces of sulphuretted hydrogen and moisture.

II. The second point to be attended to is that the quantity of oxygen (or hydrogen) added to the gas to be burnt must be in excess over tho calculated quantity (a large excess is not The mixture must be perfectly necessary). homogeneous before the spark is sent through

III. The gaseous mixture must be brought to a proper state of attenuation. Let us assume that the gas to be burnt is a pure specimen of H, CO, Cli, or some other gas, $C_{\lambda}\Pi_{\mu}$. A glance at the formula shows how many volumes of oxygen we have to add to produce what we may call the respective fulminating gas. Thus the equation $C_2H_4 + 3O_2 = 2CO_2 + 2H_2O$, tells us that every one vol. of ethylene needs 3 vols. of oxygen. Any fulminating gas will explode when the spark is sent through it at the ordinary pressure, but the force of the explosion is in general more than the best cudiometer will To avoid such aecidents, we must attenuate the gas by addition of diluents (such as surplus oxygen or air), or by mere expansion, or in both ways. In practice we must go even beyond the safety point, because in most cases nitrogen is present even in the original gas, and a considerable quantity of this nitrogen may be converted into nitric acid if the temperature of the flame is too high. But we must take eare on the other hand not to attenuate too largely, or else the mixture may miss fire, or, what is worse, suffer only partial combustion. The effect of an explosion-in the chemical, physical, and mechanical, sense-is determined by many independentvariables, which, if arranged in the order of their importance, would begin with the chemical constitution of the gas to be burnt, and end with the relative narrowness of the audiometer. But given a certain eudiometer, and suppose it to be charged with a certain fulminating gas which contains, let us say, unit vol. of the respective fuel,' measured at the ordinary temperature and the pressure of one atmosphere, the attenuation of this gas to a certain eudiometer space, equal to a units of vol., will render the explosion both safe and effective. A of course, has one value if the attenuation be produced by mere expansion (mero reduction of pressure), another valus

addition of air, a third, fourth, &c., in intermediate casea; each case fortunately admits of a liberal toleration, $\pm (\Delta A)$. The A for a given species of fuel can of course be determined only by experience; eupposing it has been ascertained for H, CO, CH, and the value for CH, ie A, we might suppose that the proper A for C₂H₄ or C₂H₆ would be about 2A₀, that for a C₃-gas about 8A, &o.; but unfortunately the supposition is not borne out by experience; C₂H₂, for instance, explodee far more violently than C₂H₃, although it contains less hydrogen per molecule. But to pass to experience. According to Bunsen and Kolbe, the explosion of ordinary fulminating gas $(H_2 + \frac{1}{2}O_2)$ in admixture with air takes its normal course at from 500 to 600 mm, total pressure, if the percentage of the explosive gas lies between 20.8 and 39.1. According to our calculation from the data of the five experiments recorded by Bunsen, this comes to the same as saying, if the partial pressure of the fulminating gae liee between 108 and 230 mm.; or if a, roferred to the hydrogen, is between 4.9 and 10.5. If A>10.5, the gas fails to burn; if A<4.9, nitric acid is produced. In the combustion of a given quantity of oxygen by added hydrogen, we may use 3-10 volumes of the latter, per 1 vol. of oxygen, if we start with almost pure oxygen. In the analysis of ordinary air, 0.5-1 vol. of hydrogen per 1 vol. of air works well (Bunsen). Whenever hydrogen is used as a reagent, the chance of nitrogen being drawn into the combustion is relatively small, so that we have greater latitude on this score, in choosing our conditious. If the oxygen to be determined is accompanied by an unknown proportion of nitrogen, we first try two volumes of hydrogen for one of total gas; if the mixture fails to explode properly we add the requisite proportion of fulminating gas, i.e. so much of the latter that it forms about 40 p.o., but no more, of the whole, and explode again; this time presumably with success (Buusen). In the case of marsh gas, Bunsen directs us to add 8-12 volumes of air besides the necessary 2 volumes of oxygeu, which, assuming the mixture before the explosion to be at 600 mm., makes our a equal to 14 to 19. For C2H4, his directions are somewhat obscure, but in a testanalysis quoted by him, the pressure of the mixture as exploded was 516 mms., and it contained 0.04868 of its vol. of C₂II₁. Hence $\lambda = 28.6$; and the partial pressure of the explosive gas $(C_2H_4 + 3\tilde{O}_2)$ was 106 mm. The addition of large volumes of air to tho

The addition of large volumes of air to the gas to be analysed does not of course add to the precision of the work generally, and in the best case will reuder the determination of the nitrogen in the ultimate product somewhat uncertain.

Thomas (C. J. 35, 213) was the first to substitute mere expansion for dilution; the (Frankland) apparatus he used enabled him to do this without trouble. Lothar Meyer and Senbert (C. J. 45, 581) have lately taken up the same method and rendered it available for Bunsen's apparatus by the invention of an auxiliary apparatus in which a kind of moreurial air-pump, constructed on the Geissler principle, serves to establish any desired pressure at the same time in the eudometer and in a moist-vacuum barometer, so that the difference of level between the meniscuses of the two at once gives the pressure of

the dry gas. By means of this apparatus, they ascertained, for each of a series of gases combustible by oxygen, the minimum pressure at which the undiluted fulminating gas is exploded by an electric spark, and also a range of preseuree at which the explosion is both safe and effectual. The following table summarisee what for us are tho main results. To explain the headings let us give the reading of the table for CH, in full. Imagine a given quantity of marsh gae mixed with a little more than two volumes of oxygen; this mixture will explode normally if its pressure is reduced to P = 140 mm. by mere diminution of pressure, the partial pressure of the CH, itself will now be at 47 mm., and its attenuation (as defined above) at $\lambda = 10$, that is to say, every 16 units of vol. of the expanded mixture contains 1 vol. of CH, measured at 760 mm.

	P	. <i>p</i>	
Fuel	in milli	imetres	A
CH,	140	47	16
C,H,	70-80, say	75 19	40.5
$\mathbf{C}_{2}\mathbf{H}_{2}$	40-50, say	45 13	5 9· 1
C_3H_5	80	14.5	52·2
$\mathbf{C}_{\mathbf{s}}^{\mathbf{H}}\mathbf{H}_{\mathbf{s}}^{\mathbf{s}}$	80	13.3	57·0
co	243-219	162 - 146	4.7-5.2
H_2	176_127	117-85	6.5-9
	(Partial 1		
$[\mathbf{H_2}]$	176_127	117-85	4.9-10.5]

[By Bunsen and Kolbo's experiments (vide supra); air added as diluent; total pressure in the mixture as exploded, 520-590 mms.]

With Meyer and Scubert's, or some other equivalent, apparatus at hand, the order of operations with a gas of unknown composition is as follows:—After having added a sufficient volume of expension, we next expand so largely as to be certainly on the right side of the safety line, and apply the spark; if no explosion occura we repeat the trial at successively greater pressures. Should the greatest available pressure fail to produce inflammability, we add a suitable proportion of ordinary fulminating gas $(\mathbf{H}_2 + \frac{1}{2}\mathbf{O}_2)$ as above explained, &c., &c.

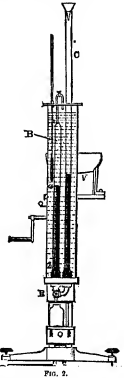
(H₂ + ½O₂) as above explained, &c., &c.

The Bunsenian mode of gas-aualysis, while perfection in regard to precision and eleganco, is very wasteful of time, for obvious reasons, which any reader who has followed us so far will easily discorn. The desire to do away with this evil has led to the construction of quito a series of more or less complicated gas apparatus. The more important of these are described in the following paragraphs. To avoid repetitions, let us state beforehaud that all the apparatus to be noticed agree in the following points:—

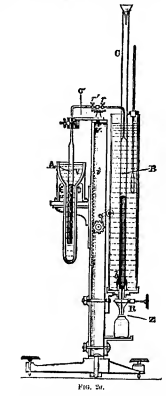
1. For accelerating the absorptions the reagente are all use a sa liquids, and the absorptions are carried out in a special piece of apparatus (laborat ire); the residual gas is then transferred to the measurer, where it is saturated with vapour of water, and measured.

The measurer is immersed in a water-bath to bring the gas contained in it to a definite constant temperature, without much loss of time.

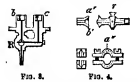
 The mode of measurement is so contrived that the calculation of the gae-quantitiee (the qs) becomes very easy or even unnecessary. Regnant and Reiset (A. Ch. [8] 26, 883), while engaged in their great research on respiration, felt the want of a quick-working apparatus for the numerous gas-analyses involved, and at last adopted the combination represented in figs. 2 and 2a. The vertical tube A conjointly with the moveable trough v constitutes the



laboratory; the measurer consists of a long U-tube, the limbs of which are of glass, while the bend consists of an iron or steel tube, terminating in two sockets b and c (see auxiliary figure 3), in which the two glass tubes B and c are fixed by means of a resinous cement. A two-way cock R below n (fig. 3) enables one to effect the necessary connections. Tubo B is provided with a couple of fused-in platinum wires near its top, so that it can be used for the combustions as well as for the measurements of the gases. Tube c conjointly with n serves as an open manometer. The capillary ends of A and B are cemented, each into the socket of a capillary steel stop-eock, and the ends of the two steel-fittings which face each other are shaped so as to constitute the two halves of a Regnaul'coupling, so that the two tubes can be united hermetically, or can be separated, at a moment's notice. The construction of a Regnault's coupling is seen from fig. 4. To unite a and b, the convex end of r (fig. 4) is smeared over with melted india-rubber, pressed against the concave part a' b', and the two are then bound together by means of the olip a''. As the conical groove in a" has a slightly less angult aperture than the sharp welt which it got over, if the two halves of a" are screwed again each other, they exert a powerful pressure, an make the joint absolutely tight. The volumete n, in the original apparatus, had only or mark, somewhere about the middle; but the



inventors subsequently added two more, on close to the upper end, and one near the lower for the measurement of exceptionally small, o large, quantities of gas. The manometer c i not graduated, as the apparatus is intended t be used with a cathotometer; where this costly



instrument is not at hand, tube o must be provided with a millimetro scalo.

To prepare the apparatus for use it is placed on a substantial support not liable to inconvenient vibration, and the three levelling scrow of the stand are adjusted so that the tube and o stand vertical. To determine the relative gas-volumes corresponding to the three marks, the volumeter is filled with mercury, through σ , and after the air-bells have been removed by the well-known artifices, the weights of mercury w_1 , w_0 , w_2 , which the tube holds from its exit end at r (fig. 2a) to the highest, middle, and lower, mark, respectively, are determined. For comparative measurements the volumes are put down as $\frac{w_1}{w_0}, \frac{w_0}{w_0} = 1$, and $\frac{w_2}{w_0}$,

respectively. In the absence of a cathetometer the level points of the three marks on the scale c must be determined with the help of an ordinary gas-room telescope. Lastly, a drop of water is introduced into B and spread over its surface. To analyse, say, a mixture of carbon dioxide, oxygen, and nitrogen, a sample of the gas is collected over mercury in a (perhaps with the help of an auxiliary-trough); tube A is coupled on to n (which is supposed to be quite full of mercury), and the gas is snoked into this tubo by letting mercury run out at z. B having been closed by shutting the cock r, communication is made with c, and mercury is run out until the meniscus in n stands at say exactly the middle mark; the final adjustment is made with the tolescope when the temperature of the gas has certainly become equal to that of the When the final reading is made, n must of course communicate with c only. The reading of the height h of the mercury column in c, counting from the respective mark up or down as the case may be, and the reading of the barometer, complete the measurements. Supposing h to be positive, and the barometer to stand at n, the gas-quantity measured is

$$Q = \frac{(V_0 = 1) \times (h + B - \pi)}{273 + t}$$

To absorb the carbon dioxide, the laboratory tube (which was left full of mercury) is charged with a little caustic potash solution, and the gas is blown into it from B. By letting the gas travel forwards and backwards between A and B a number of times, the absorption can be completed in a short time. The residual gas is then sucked back into B, care being taken to shut the oock r'as soon as the potash solution comes to some mark o, in the capillary part of a. The thread of gas from σ to r which is thus lost is of no consequence, as it amounts to only 1000 of The mixture of nitrogen and oxygen is measured as before. The rest requires no explanation. If all the several gases are measured at the same temperature and volume, the (dry) pressure P', P", P", of course may be taken as representing their os (red. vols.).

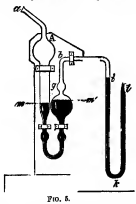
Frankland and Ward, in 1853, introduced an ingenious modification of Regnault's apparatus, which differs from the original model chiefly in this, that the volumeter bears ten marks, so adjusted that the respective gas-volumes are to one another as 1:2:3 10 exactly, and that in addition to Regnault's open tube c (Figs. 2, 2a), there is a third tube, p, which terminates above in a stoppered funnel or stop-cock. Tube p stands in the same water-bath with an and c; when used it contains only mercury and a little water, and thus assumes the character of a 'moist' barometer, which serves to directly measure the dry pressure of the gas shut up in b. Tube o (in F. and W.'s apparatus) serves only

for the introduction of the mercury. The levels of the ten volumeter marks, in reference to the scale on the barometer, are of course determined once for all, hence the measurement of a gas, supposing its volume to have been adjusted to one of the ten marks on the volumeter, involves only one reading, namely that of the height of the mercury column in the barometer, which balances the (dry) pressure of the gas. Another advantage of F. and W.'s apparatus is, that for each gas measurement it gives one the choice among at least some three of the ten standard volumes, and thus enables one to reduce the error by an obvious method of repetition. Unfortunately, however, the barometer rather aggravates what in the original apparatus is a sufficient trouble, namely, the liability of the apparatus to get out of order. However carefully it may have been constructed, the joints between the glass tubes and their sockets are sure to become leaky, and the capillaries between the laboratory and the volumeter are oxasperatingly fragile.

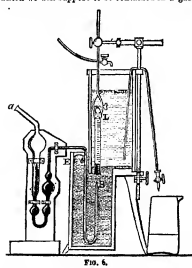
MoLeod [1869] (C. J. [2] 7, 314), and Thomas [1879] (C. J. 35, 218) endeavoured to remedy these evils, and to effect other improvements. For details see the papers referred to.

Infinitely handier than Regnault's unwieldy machine, though not quite equal to it in potential precision, is

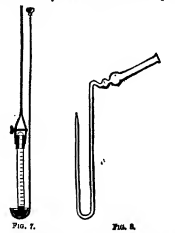
Doyère's Apparatus.—(First notice dates from 1818. Full description in A. Ch. [3] 28, 1.) The essence of Doyère's system is that the measurement of the gases is effected in a plain graduated eudiometer, while a series of Ettling's gas pipettes serves for the chomical treatment of the gases, and their transference from vessel to vessel. The Ettling gas-pipette is depicted in fig. 6, and a glance at the figure suffices to



show, in a general way at least, how the instrument is used for the transference of a gas from one tube to another; nor is it necessary to formulate the conditions or limits of its availability. The measurer (fig. 6) when in use is suspended over a pneumatic trough, deep enough to admit of the total immersion of the measurer, and is surrounded by a mass of water contained in a cistern whose sides are of plate-glass, while the mercury of the trough forms its bottom. To prepare the measurer for the reception of a gas, it is cleaned, slightly moistened inside, transferred to the trough by means of the portable mercury trap (fig. 7) fixed in the clip L, and filled with mercury by sucking out the air, by means of the U-shapod tube (fig. 8). The gas, which we will suppose to be contained in a gas-

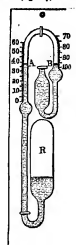


pipette, is then blown in, to be measured at a certain fixed disgregation, which is kept rigorously constant for the set of gas-quantities to be compared. A glance at fig. 6 at once suggests a mode of fulfilling this condition. But this mode is not Doyero's. He allows the tempora-



ture of the bath and the barometer to take care of themselves, but before each measurement he so adjusts the height of the water in the bath that the volume of a certain fixed quantity of air, shut up over water at a place within the water of the bath, assumes a certain fixed value,

v. The standard body of air is contained in the 'Regulateur' (fig. 9), a kind of air-ther-



F1G. 9.

mometer which is fixed against a glass-plate, and, by it, susponded at a certain (hy intention constant) height over the mercury-level of the trough. The water of the bath goes to some point B in the ascending branch of the capillary U-tube; BA is a thread of air; from A downwards there is a continuous mass of water, over which the standard body of air is shut up at B. Before each gas-measurement, the height

of the water in the trough is so regulated (by means of taps) that meniscus a stands at some determined point of the scale, and the air which serves as regulator is consequently at some fixed volume v_0 . This being done, the eudiometer is raised or lowered, until the height of the column of mercury suspended in it is at some fixed value, h_0 . As a result, the gas is now practically at least, at a fixed disgregation.

Proof. The pressure of the gas exceeds that of the air of the regulator by $\Lambda + p_1 + p_6$, where Λ stands for the height of u 'over' Λ (we refer to the regulator), and p_1 for the height from the level in the trough to that in R of the regulator—both reduced to mercury. With a properly chosen h_0 , the value $e=\Delta + p_2 - h_0$, if not nil, is at least small, and nearly constant. Now supposing we have, for two successively measured quantities of gas, I and II: I. For the reculator-sir V, V': V', V': V'': V''.

For the rogulator-air v., r'; 1'. v.; r''; p''.
For the gas . . . v'; r''; (p''+c) v''; r''; (p''+c).
As the regulator-air is at the constant volume, v., we

$$\frac{T'}{p'} = \frac{T''}{p''}.$$
The 'reduced volumos' (the qs) of the two gases are
$$Q' = \frac{V'(P'+c)}{T'} \text{ and } Q'' = \frac{V''(1''+c)}{T''}.$$
and, as e

$$\frac{P'}{P'} \text{ or } P''$$

 $\frac{Q^{r}}{Q^{r}} = \frac{Q^{r}}{Q^{r}} \left\{ 1 + \frac{\theta}{P^{r}} \left(\frac{P^{r} - P}{P^{r}} \right) \right\}$ As both factors in the second term with the bracket are very small, we have practically,

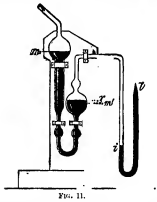
Q" = \vec{v}''

As the measurer is necessarily very small, the adjustment of h_0 must be made, and the

gas-volumes read, with more than ordinary exactitude. Doyère accordingly provides a small short-vision telescope, which has a glass micrometer-scale (fig. 10) in its focus. The



telescope is attached to a three-legged stand (which rests on a horizontal glass-plate fixed on the table close to the trough), in such a way that in all the necessary shiftings the optical axis remains parallel to, or when necessary, in, the same horizontal plane. To adjust ha the telescope is so focussed that it gives a distinct image of the mercury meniscus in the trough, which image is then made to coincide with line C-O' (or BB' if the telescope is an astronomical one). The eudiometer is then lifted or lowered nntil the image of the top of its meniscus touches the central line A-A', which assigns to he a definite, though unknown, value. This adjustment being made, the telescope is drawn backwards a little on the glass-plate to afford a good image of the eudiometer-scale, and to enable one to read the volume of the gas. The



miorometer-scale serves to sub-divide the individual divisions on the eudiometer, which at does with an amply sufficient degree of precision. Before reading h_0 the eudiometer must be tapped

to bring the mercurial meniscus into its normal shape.

Assuming now that a gas, measured as described, contained carbon dioxido and air, and that we wished to determine the carbon dioxide by absorption with caustic potash. We begin by charging a gas pipette with mercury to about the extent shown in fig. 5. We then take the pipette to an auxiliary trough, immerse its U in the well, and, after having blown out the air, suck in the requisite quantity of caustic potash solution from a test-tube inverted over the trough, taking care not to let any more mercury follow than is necessary (practically) to trap

the contents by a thread of mercury il. We then transfer the pipette to the measurer containing the gas (as indicated in fig. 6), press down the measurer over the outer branch of the U, and transfer the gas from the measurer to the pipette, by sucking at a, until drops of mercury are seen to fall into the working bulb, but no longer. Things are now in the condition depicted in fig. 11, and all that remains to be done is to agitate the contents gently so as to insure absorption of the CO2, and then to return what is left of the gas to the measurer. This, however, is a delicate operation, which in the hands of a beginner is not unlikely to fail. The first step is to replace the pipette under the measurer, to lower the latter sufficiently (v. infra), and to blow into the pipette so as inst to dislodge the mercury thread i l. Supposing the pipette contains no more surplus mercury over and above that which was in it at the beginning, then as long as the meniscus in the eudiometer is below or at a level with that of the mercury in the trough, as it is underneath the bath, only part of the gas will pass out of the pipette into the eudiometer. The second step is to lift the pipetto, so that its outflow end, B (fig. 6) or l (fig. 11), becomes visible within the gas space of the measurer. As long as it is there, and the pipette is kept vertical, whether the gas flows out of B, or in at B, or remains at rest, depends mainly on the pressure of the gas in the eudiometer, and consequently on the altitude of the latter. But this altitude we have under absolute control. Hence what we have to do is carefully and slowly to lift the eudiometer until the thread of liquid reagent which makes its appearance as soon as the bulk of the gas is out, has come to, say, 2 mm. from the outflow end. We then stop sucking, pnt the pipette down on the table (which of ceurse at once seals the end B with mercury), suck at a until we see mercury dropping into the pipette, take the pipette out of the mercury, and put it on the table to have it at hand for a repetition of the absorption.

The sequence of operations described is not quite so easy in practice as it looks on paper, because success depends largely on the permanence of the position of the pipette in reference to the plumb-line. Tilting over the pipette in the direction of the U means adding to the pressure of the gas inside; and vice verst.

For the explosions, Doyere provides a special stout pipette, with fused-in platinum wires, &c.; but the method of combustion finds little favour in his eyes, because his apparatus does not

readily fall in with its requirements.

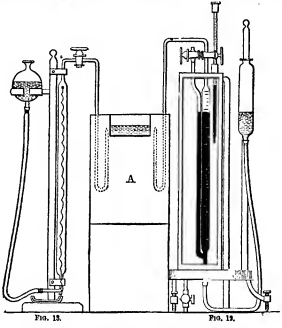
In conclusion the writer may be permitted shortly to describe an apparatus of his own invention, which, thanks to the valuable assistance of Mr. Lennox, he was enabled to construct on his own premises, and which has since done him good service.

Dittmar's apparatus, like Doyère's, is based upon the Ettling gas-pipette. Apart from the necessary two troughs, it consists of the follow-

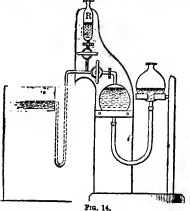
ing three independent parts.

The measurer (fig. 12) is a combination of a wide with a narrow glass-tube, after the manner of Gay-Lussac's burette. The wide tube communicates by its lower contracted end

with a long capillary tube of india-rubber, and | volumes are counted from the point of the through it with a Geissler mercury-reservoir. junction, because, after the introduction of a At their upper ends both tubes are provided gas, the narrow canal firmly retains its thread with Geissler stopcocks; to the exit-end of the of mercury. The measurer holds a fixed posiwide tube is soldered the capillary U tube, I tion on the right side of a pneumatic trough, A,



characteristic of Ettling's pipette. The wide tube bears a mm. scale; the gas-volumes ecrresponding to the several marks are determined by gravimetric calibration, at a rigorously



constant temperature, maintained by means of the water-bath. The narrow bit of tube between the top of the measurer and its stopcock is a capillary of the same bore as the U; it joins

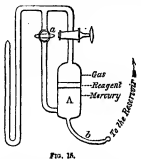
provided with two wells, one for the U of the measurer, the other for that of the exploder. In regard to the exploder (fig. 13), we have nothing to add to what is clearly seen from the figure, except the statement that the exploder in its present form is wider than the figure represents it to be, so wide, indeed, as to enable one to expand a gas considerably before exploding it.

The absorber in its original form is represented in fig. 14. For the interpretation of this fig. it suffices to say that R is a small mercuryreservoir which enables one to sweep out the thread of gas left in the capillary after the liquid reagent has been allowed to travel up to the safe side of the point of junction between the horizontal part of the capillary delivory tube and the stem of the reservoir. An improved form of the absorber (devised by Mr. Lennox) is represented in fig. 15.

To prepare the measurer for the reception of a gas, it is completely filled with mercury from the reservoir, the stopcock of the side tube is turned off as soon as all the air is driven out of it, and a drop of water is introduced into the main tube at a suitable stage.

The gas to be measured must be contained in a tube short enough to be within the range of the U: from this tube the gas is sucked into the measurer with the help of the reservoir, which on quite abruptly to the wide tube, and the is then adjusted so that the gas-pressure inside

is about one atmosphere. The stopcock at the side tube is then opened, and the height of the reservoir is re-adjusted so that the manisci in the narrow and wide tubs are in the same horizontal plane. A horizontal wire in the telescope facilitates this adjustment materially, but is not indispensable. The gas ie now at the preseure $B + \bar{b} - \pi$, where B is the height of the baromoter, # the pressure of the vapour of water, and b the excess of the capillary depression in the narrow eide tube as compared with that in the wider branch. The temperature is of course that of the water-bath. As both P and Toscillate with a series of measurements only



within small amplitudes, it is expedient to reduce, not (for instance) to unit disgregation, but to come mean pressure and temperature (if there has been any variation in either or both) by means of some suitable formula, such

$$\mathbf{v}_0 = \mathbf{v} \left\{ \mathbf{1} + \frac{(\Delta \mathbf{P})}{\mathbf{P}_0} - \frac{\Delta \mathbf{T}}{\mathbf{T}_0} \right\}$$

 $v_0 = v \left\{ \begin{array}{l} 1 + \frac{\left(\Delta \, F\right)}{P_0} - \frac{\Delta \, T}{T_0} \right\} \\ \text{whero } P_0 \text{ and } T_0 \text{ stand for tho standard values,} \end{array}$ and the observed values are assumed to be greater than these by (AP) and (AT) respectively. If a table of the reciprocals of the practically occurring rs and Ts is at hand, the calculation becomes so very easy that it is not worth while to set up a disgregation indicator.

Technical Gas-Analysis.

To meet the demands of chemical industry there has been invented a variety of methods for the rapid, though perhaps only approximate, analysis of certain classes of gas-mixtures. The methods all agree in this, that the use of mercury is dispensed with, the gases being measured over water, or even perhaps over the respective absorbent solutions. The Bunte Gas-burctte may be quoted as a typical example of this class of apparatus. Imagine a long cylindrical pipette graduated for gas-volumes and provided with a stop-cock at each end, and combinable with a reservoir by means of a long india-rubber tube. To analyse, say, a chimney-gas, the burette is filled with the gas by displacement, and the reservoir, after having been filled with water, is attached below. By placing the reservoir at a certain convenient altitude, and temporarily opsning the upper cock, a certain volume of the gas is shut off at the pressurs of one atmosphers. In order new to determine the carbonic soid, we suck out the water by an (easily imagined) auxiliary apparatus, and replace it by a solution of caustic potash, which is shaken with the gas. The caustic potash is then sucked out, water ie let in, the original pressure is re-establiehed, and the recidus is measured. In a similar manner, the oxygen ie determined by absorption with alkaline pyrogallate.1

ANALYSIS BY THE METHON OF TITRATION.

This branch of analysis comprisee the applications of what was described in a previous section as the titrimerric method of indirect weighing. The method in any of its present forms is applicable only to such reactions as processd readily in aqueous solutions; the reagents, accordingly, are always used in the form of standard solutions (liqueurs titrées), i.e. solutions the strengths of which are known in reference to the process under consideration.

The amount of standard solution required in a titiation may be measured either by weight or by volume; in either case the measurement of the solution is only an indirect mode of weighing the active agent contained in it. The gravimetric method is certainly susceptible of the higher degree of precision; yet the volumetrio method is universally preferred, because it is by far the more handy and expeditious of the two, and, if properly conducted, (with very few exceptions) does aniple justice to even the bost titrimetric processes.

The invention of volumetric analysis must be credited to Gay-Lussac. Long before him, it is true, Stirling enunciated the principle of the method, and Vauquelin and Descroizille used it for assaying commercial alkalis; but to Gay-Lussac undoubtedly belongs the credit of being the first to bring the method into an exact form, and to work out all its technicalities in the most masterly manner. Volumetric analysis was slow in progressing. Gay-Lussac's more immediate successors, misled by his success in regard to silver, directed their attention almost exclusively to the translation of established gravimetrio into volumetric methods; failing to see (what ie now so obvious) that the number of reactions to which both methode are applicable must necessarily be very limited.

Very little real progress was made until 1856, when Bunsen, by introducing a new idea, gave a fresh impetus to investigation. Starting from the well-known reaction which takes place when iodine solution is dropped into aqusous sulphurous acid (and which Langloie had already utilised for the detormination of this substance), he established the conditione under which the process takes the precise course indi-cated by the equation; and on the basis thus gained he developed exact methods, not only for the direct determination of these two bodies, but also for the indirect determination, by means of the same two solutions, of a whole

¹ Professor Winkler, of Freiberg, has made a special study of this branch of gas-analysis, and has written two excellent books on the subject; one of these has been translated into English by Professor Lunge. To these books and another we refer for further information. (1) Dr. Clemens Winkler, Antelump, sur chemischen Analyse der Industrie-Gase (Freiberg, Engelnardt, 1876). (3) An ahriiged edition of the same by the author. Translated by Lunge (Van Voorst, London). Also, New Methoden sur Analyse der Gase, von Walter Hempel (Brannschweig, Vieweg n. Bohn, 1880).

ceries of oxidising agents, for which an equivalent of iodine can be substituted by the purely qualitative execution of certain reactions.

By this memorable research volumetrie anslysis found its true ephere of action, as an invaluable means for the determination of generic radicles, such as the active oxygen in peroxides, the loosely held chlorine in perchlorides, the replaceable hydrogen in acids, the oxygen or chlorine-equivalent of reducing agents; for a hoet of determinations, in short, which practically lie outside the range of the gravimetric method. Where the two methods compete in the colution of the same problem, volumetrio analysis generally offers the advantages of greater promptitude and facility of execution; it, indeed, stops where with gravimetric analysis the most difficult part of the work would begin. advantage, however, is not an absolutely clear gain. The volumetric method, so to say, does not look at the body to be determined, but, in a somewhat blindfolded way, only measures one of its ohemical properties, which in no case appertains to that body only; hence errors are more likely to be overlooked, and are far more difficult of subsequent correction, in volumetric, than in gravimetric, analysis; for gravimetric analysis furnishes the thing to be weighed in the form of a definite compound, which can be examined for its purity, and, if necessary, be purified before it is weighed. A small amount of iron, nickel, zine, &c., which has escaped precipitation may be searched for in the filtrate and recovered; any deficit or excess obtained in a titration is thrown away with the rest of the products.

We have no space for a full history of our subject; yet we must not forget to give credit to the late F. Mohr for having contributed largely to the modern development of volumetric analysis, by his criticisms of old, and his invention of new, methods; by the construction of useful apparatus; and last, not least, by the compilation (for the first time) of an original and com-

prehensive handbook 1 on the subject.

In now passing to the systematic exposition of our subject, we will begin with a few remarks on the

Graduated Glass Measures

which serve for the preparation of the standard solutione, and the necessary measurements of liquids generally. Volumetric analysis of course involvés only comparative measurements; we indeed never measure a etandard solution otherwise than in reference to itself; hence the unit of volume may not only be chosen at pleasure, but need not bear any known relation to the nuit of weight. But the only correct mode of gauging a liquid measure is to determine the weight of water it holds (or delivers); hence for those who are in the habit of using the gram as their unit of weight the most convenient unit of volume is the volume at (let us say) 15°C. of that mass of water whose uncorrected weight in air is one gram. We might herewith adopt this unit and call it the 'fluid gram.' In doing so we should not be guilty of any innovation. The customary unit with most chemists and instrument-makers,

Mohr, Lehrbuch der chemisch-analytischen Türirme-thode. The first edition dates from 1887; the fourth and hast from 1885.

it is true, is the cubic continetre, but it is this only nominally; the actual unit in almost all commercial 'cubic centimetre' measures comes nearer to onr fluid gram than to the nominal unit. From what we have said, the reader will nnderstand that if in the sequel we speak of cubic centimetres, or litres, these terms mean only unit-volume, and 1000 unit-volumes, respectively, unless it is clear from the context that we mean to approximately define an absolute quantity, or to refer to the well-known relation between the litre and the kilogram.

As all aqueous liquids wet glass, the mark on a litre flask, &c., can be correct only in reference to a epecified mode of reading. The best mode is this. Place the vessel so that its axis is vertical, and look at the meniscus horizontally with one eye. The meniscus then appears as a flat crescent-shaped strip. The lower boundary of this strip is taken as the line of reference, and the real, or imaginary, mark on the graduation with which it coincides (visionally) is taken down as the reading of the liquid. The line referred to gains in sharpness of definition if it is observed in transmitted light, and a strip of black paper is fixed to the back of the measure about 2-3 mms. below the line. With only one of the customary standard solutions, namely the almost opaque solution of permanganate of potash used for iron titrations, this mode of reading does not work. In the case of this liquid we must take the upper boundary of the meniscus as our reference mark; this upper boundary assumee its maximum definition if viewed in reflected light, and with a white background (a picce of paper) immediately behind it. Any reading made in this exceptional manner ie of course subject to an obvious correction, the amount of which is ascertained by measuring the height of the meniscus of a transparent solution in the same vessel. In most cases, however, the volume to be determined is the difference between two consecutive readings, so that the correction in question becomes unnecessary.

In a vessel which serves for measuring out a certain volume, the small quantity of liquid which permanently adheres to the glass must be allowed for by the maker, i.e. the seale must be constructed so as to include what would otherwise be a necessary correction. In the case of graduated pipettes this can of course be done only on the basis of a conventional mode of emptying out, which, when once fixed upon, must be rigorously adhered to. The thermic expansion of glass may unhesitatingly be neglected in the graduation of a titrimetrio measure. A glass flask which holds 1000 c.c. at 15°C., at 15 ± 10° holds 1000 c.o. ± 0.27 c.o.; i.e. only 1100th more or less. The thermic expansion of the solutions measured is far more considerable, and cannot in all casss be neglected. We shall come back to this point in the next section.

The chemiet now-a-days has no occasion to graduate his own burettes, litre-flasks, &c., but he should never use a set of instrumentsalthough they come from the most famous maker—without having first tested them. The following is the method to be pursued. Passing from measure to measure, and with each measure from mark to mark, measure in or out the several marked off volumes of pure water of

known temperature, and determine their weights in grams. In the case of apparatns with a running-on scale, only every 10th or 20th mark need be checked in this manner, unless there are visible irregularities in the graduation. After having thus gone through the whole system, reduce all the water-weights to the same temperature, eay to 15°C. (i.e. from the observed weight of water of to calculate the weight of water of 15° which fills the same space); divide each corrocted weight by the corresponding nominal volume, to find the weight-value of the actual unit-and draw your conclusions. Supposing the coveral units agree fairly, select a suitable average value (net necessarily the mean, because the numbers are not all of the same weight mathematically) as the unit, and calculate the volumes corresponding to the several marks in terms of thie adopted unit. The results ought, by theory, to agree with the respective nominal values, but in practice, of course, we cannot expect absolute coincidence. In a burette, for instance, which gives $\frac{1}{2}$ o.o.s directly, we must tolerate ± 0.1 c.c., and with the lower marks, even a little more. Whether the actual unit is, or is not, equal to the nominal is of no consequence; yet, if it is not, it is obviously advisable to note down its value -in fluid grams or c.o.s -for future reference.

To facilitate the calculations involved in such work as the graduation of instruments, the writer many years ago calculated the following table:—

A mass of water, which, in air of $t^{\circ}C$, and 760 mm. pressure, balances a brass kilogram weight, at $t^{\circ}C$. occupies (1000 + x) fluid grams = (1000 + y) true cubic centimetres.

	to.	x	3/ a	to	\boldsymbol{x}	y
+	00	-0.64	+ 1.25	+ 15°	+0.00	+1.89
	4	0.78	1.12	16	0.15	2.04
	8	0.68	1.21	17	0.30	2.20
	9	0.63	1.27	18	0.47	2.37
	10	0.56	1.34	19	0.66	2.55
	11	0.47	1.43	20	0.85	2.74
	12	0.38	1.52	21	1.06	2.95
	13	0.27	1.63	22	1.28	3.17
	14	0.14	1.76	23	1.51	3.39
	15	0.00	1.89	24	1.75	3.63

Preparation of Standard Solutions.

In fixing upon the degree of concentration for a epecified standard solution, we may allow ourselves coneiderable latitude. As a rule the nature of the volumetrio process for which the solution is to be used may be left out of account; all we need look to is that the probable inherent error of the method corresponds to a distinctly vicible difference of level in the burette; say to 0.1 or 0.2 c.c. according to the size of the 'c.c.' With methods possessing a very high degree of inherent precision this rule would lead to an inconvoniently dilute reagent. In such (rure) cases we holp ourselves by supplementing a moderately dilute etandard solution with a decimal solution, meaning a colution prepared by diluting the reagent proper with water to 10 times its volume. The decimal colution serves only to finish the titration which has already been almost completed by means of the standard solution proper. This system of course is a mere delusion unless the stronger solution be

measured with at least 10 times the precision attained with the decimal one.

The strength of any given solution should be so defined as to reduce the subsequent calculations to their highest degree of eimplicity. Supposing, for instance, we have to deal with a standard sulphuric acid intended for the measurement of alkalis; evidently it would not be expedient to note down the number, n, of grams (or mgms.) of H2SO4 or SO3 which is contained in each litre (or o.e.) of the roagent. As the number when calculated into KHO, NaHO, Na₂CO₃, &c. has always to be divided by $SO_4H_2(=98)$ or by $\frac{1}{2}H_2SO_4(=49)$, it is obviously better to calculate the value $\frac{n}{49} = \tau$ (as a decimal) and put down this T as the strength of the solution. Were the solution intended exclusively for the determination of soda, to be reported as Na2O, it would be still better to calculate the value of $\frac{n \times \text{Na}_2\text{O}}{\text{H}_2\text{SO}_4} = \tau'$, and note down the weight r' (of sodium monoxido) as the

strength of the reagent. For the etandardisation of a epecified solution we have in general the choice between two methods; (1) a quantitatively exact synthesis; (2) an approximate synthesis, fellowed by an exact analysis. The first method may assume one of two forms; we either weigh out exactly eo much of the pure reagent, dissolve it in water, and diluto to, say, 1 litre; or else (if the pure reagent is not itself at hand) we analyse. say, a strong solution of the pure or impure reagent, by means of some very exact method, and synthesise directly on the basis of this determination. In regard to the second method. the first step of course is to procure an approximately correct solution. For example, let ns assume we wish to prepare a standard hydrochloric acid containing exactly HCl = 36.5 grams per litre. An apology for such a solution might be obtained from the ordinary (pure) laboratory acid, by determining its epecific gravity, deducing the percentage, and synthesicing on the basis thue gained. In all cuch cases it is expedient to so allow for the uncertainty in the assumed percentage that the solution obtained is sure to be stronger than intended. In accordance with the rule, let us assume the actual strength, as found by analysie, to be p, instead of the intended strength p_0 , and let $p > p_0$. Obviously our 'v' litree of reagent must be diluted to $\mathbf{v} = \frac{vp}{p_0}$ to bring the strength down from p to p_0 ; the liquid must not be diluted with $v(\frac{p}{n}-1)=1$ litres of water, because the two liquids when mixed would contract, and a little more than w litre, say $(1+\epsilon)$ w litre, of water would be necessary to bring up the volume to the intended value, v. In practice, however, large volumes (euch as we assume our v and v to be) cannot be measured with adequate precision, so that the second (theoretically faulty) method is generally the better of the two. It certainly is the better if the required correction is only emall; if for instance $(\frac{p}{p_0}-1)$ is something like 0.03 or less. In such a case, if only w as calculated is measured accurately, the corrected solution will be as near the intended strength, p_0 , as the given solution was near its strength p.

As an example, let us take $\frac{p}{p_e} - 1 = 0.03$; v = 10 litres; $\delta v = \pm 0.5$ litre (which is a liberal allowance); and we have

$$\frac{\delta p_0}{p_0} = \frac{\delta p}{p} \pm 0.0015.$$

Even in such a case it is only prudent again to analyse the corrected solution, to see that no blunder has been made. Supposing (to return to the example) the number p had been the result of three well-agreeing analyses, the intended value for p, had been 37-00, and the analysis of the corrected solution had given for p, the value 36.84; the most probable value for the actual strength would be $(3 \times 37.0 + 36.84) + 4$.

Thrning back, let us now assumo $p_o > p$. In this case our v litres of solution should be reduced to $\frac{p}{p_o}v$ litres, by elimination of $(1+\epsilon)$ $(1-\frac{p}{p_o})v = (1+\epsilon)w$ litres of water. Even where evaporation would be permissible, it is better to compensate for the surplus water by addition of the substance which served to make the solution. Supposing we had used s grams of substance for every 1 litre of reagent produced. Clearly $\frac{p}{p_o}s = s_o$ grams is what ought to have

been taken. One way then is to prepare some (say $\frac{1}{2}$ litre) of the solution by means of the corrected method and to determine its specific gravity, π_0 , in order to be able to reduce to weight; thus:

Weight of substance in it . $= 1000 \, \pi_0 \, \mathrm{grms}$. Hence weight of the water . $= 1000 \, \pi_0 - s_0$

Or every gram of water requires *6 $^*=\frac{s_0}{1000}\frac{s_0-s_0}{\pi_0-s_0}$ grms. of substance to be converted into solution of the intended strength, p_0 ; hence our $v\times (1-\epsilon)$ $w\times$ kilos. of water require $v\times (1+\epsilon)$ $wc\times$ kilos. of substance. All that we need for the calculation of $(1+\epsilon)w$ is the specific gravity π of the uncorrected solution. Obviously

 $(1+\epsilon)w=1000(\pi-\frac{p}{p_0}\pi_0)$ grams. In practice, however, it is scarcely advisable to go to all this trouble. It is easy by some short cut (based on the above) to name a number of grams of substance, which if added to one litro of solu-

tion would bring up the strength to a little above p_{\bullet} . Suppose the increase of volume involved in adding these grams of substance is less than, say, 0.1, 0.2..., say y litre. Then, to set things right, we calculate the correct mass of substance

for 1+y litres, which is $(1+y) \frac{p_0}{p} s$, weigh out

what this mass is more than the s grams present, in each litre, add this to each litre of solution as given, and dilute to 1+y litre by addition of water. If p differs much from p_w it is expedient to slightly over-correct the solution, to determine the exact value, p', which the solution now has, and (if $p' > p_o$ as intended) to correct the strength, by dilution, as explained above. If p is only a little less than the in-

tended value p_{\bullet} , we may salely assume the surplns water per litre to be $1-\frac{p}{2}$ litre, and add the exact weight, x, of substance, which by calenlation converts the small quantity of water into correct solution. The result (in the absence of blunders) will be quite correct oven if v was only approximately measured, because a very small volume of water added or withdrawn from the total of v litres would make the solution absolutely correct (apart from the error in p of course). One point remains to be considered. Supposing the strength of a solution at t_0 degrees is p_0 , what is the strength p_1 at t_1 degrees? It would not do to calculate the correction from the expansion of pure water from t_0 to t_1 , because all standard solutions expand more largely than pure water does. A correct method is to determine the specific gravity (say the weight contained in a narrow-necked 100 c.o. flask) of the solution at t_0 degrees and at t_1 degrees. Supposing the weight of it is π at t_1 and π_0 at \hat{t}_0 , we have $p_1 = \frac{\pi_1}{\pi_0} p_0$.

This correction of course is indicated only in the case of very exact methods. But in their case it is best altogether to eliminate the uncertainties of volume-measurement hy effecting the final standardisation by volume and by weight at the same time; by determining for instance at the same time the weight in grams and the volume in c.o.s of the quantity of standard nitrate of silver which is required for the exact ppn. of (say) $\frac{\text{KCl}}{100}$ grms. of pure chloride of potassium. This need not hinder one in so adjusting the solution that the quantity referred to may for all ordinary purposes be assumed to be equal to 100 c.o. For the purpose of a highly exact determination, the bulk of reagent (e.g. AgNO₃ solution) required, after having been measured out, is weighed into the (chloride) solution to be analysed, and the small excess of substance or reagent left is determined by volumetric titra-

tion with decimal solutions.\footnote{Images} The adjustment of an analytically standardised solution to an exact pro-determined strength is advisable only if the solution is permanent, and is meant to be used very frequently, otherwise it is better to note down the strength as it is, and calculate from it. A solution known to contain \$1.023 \times IICl grains per litre is almost as convenient as one containing \$1 \times ICl exactly. Because for one or two analyses we can well afford to calculate, say, the product \$1.023 \times \footnote{Na_0}0\$, and for a very long series of such determinations the value may be calculated once for all and noted on the label.

On the Theory of Titration.

Let A and B be two chemical species, which, when their solutions are mixed together, combine with, or decompose, each other in some definite manner. Is the reaction available for the mutual volumetric measurement of A and B, or (let us rather say) for the measurement of A

Compare Dittmar's Memoir on the Composition of Ocean Water, 'Challenger' Memoirs, page 4; site his Exercises in Quancitative Analysis, section on Sea-scaler. by 2? It may be if, under easily realisable conditions, it procesds rapidly, and, if it is possible under these conditions to recognise the point of saturation with sufficient sharpness, i.e. the point from which onwards an additional drop of B-solution does not produce a recognisable change. In some cases the point of saturation defines itself naturally by coinciding with some sudden visible change, e.g. a change of colour. It does so, for instance, if the reaction is a double decomposition, a+b=c+d (where a, b, c, d, stand for definite relative quantities of the reagents a or B, and the products o and D respectively), and if A (or B) is intensely coloured, while n, c, and D (or A, c, and D) are relatively colourless, or at least do not hinder the observation of the colour of the last remnant of A, or a slight excess of B.

Examples: 1, Oxidation of ferrous salt (A), by permanganate (B), with formation of ferric salt (c), and manganous salt (D).—2. Decolourisation of the intensely bluo solution of cupricammonium salt (A), by the reducing action of (standard) cyanide of potassium (B), with formation of colourless double cyanide of copper and alkali metal (c), and cyanate and other salts

of alkali (D).

Sometimes when such colour-changes do not occur, they may be produced by addition to tho solution to be titrated of a suitable indicator. Thus: 1. Litmus solution may serve as an indicator in the volumetric neutralisation of acid by alkali (or vice versd) .- 2. Iron-alum may serve as an indicator in the determination of silver (salt) by added sulphocyanide of ammonium, the red colour of Fe(NCS), becomes permanent only when all the silver has been ppd. as AgNCS, and a slight excess of sulphocyanido has been added. The indicator in this case would evidently be of no use if it were not the case that Fe(NCS)_s, which is produced locally from the first, is decomposed as readily and in the same way by silver salt as the alkalino sulphocyanide is. A similar remark applies to indicators generally. If an indicator, while otherwise trustworthy, fails only to fulfil the condition stated above, it may still be available in the sense that, instead of adding it to the 'A'-solution from the first, we may apply a little of it to drops of the mixture taken out at suitable stages in the process of the reaction. Thus, for instance, in this titration of phosphato (A), by uranic acetate (e), prussiate of potash may serve as a drop-reagent, because, although unavailable as an indicator proper, if added to a drop of the mixture it produces the red-brown colour of ferrocyanide of uranium only if the uranium is present as (an excess of) acetate; the uranic phosphate is not decomposed by the prussiate. The action of an indicator nced not necessarily consist of a colour-reaction; a ppn. if sufficiently delicate, is as good in principle, though not as a rule in practice because the locally formed characteristic pp. will not disappear so readily on stirring up as the colour of a dissolved product would.

If the reaction is a steadily progressing ppn.
of the essential radicle a in A by n, the end of
the reaction of course coincides with the completion of the ppn., i.e. the point when (supposing n to be added in successive drops) the last

remnant of a has just been thrown down by the nth drop of s, so that the (n+1)th drop fails to give a turbidity. For such reactions we need no indicator or drop-test, although such may be very convenient.

very convenient. As soon as we have found somo means for recognising the end-point in our reaction with sufficient sharpness, we can decide the question as to its availability by preparing standard solutions of A and n respectively, and determining the ratio a: b corresponding to the end-point under a sufficient variety of conditions. In a first series we work with the plain solution, but take caro in one set of trials to begin with a and drop in n until the reaction is apparently completed; and in another set of trials we begin with B, pour in a slight excess of A, and then finish with B; this is done in order to see whether the ratio a:b is independent of the mode of mixing. In a second series, we add known, but varying, proportious of water. In a third series we add more or less (but always a known weight (x) of this or that body x which in the practical application of the method would be likely (if not sure) to be present, &c. From Series I. and II. we easily calculate the small excess of reagent B which must be added, per F c.c. of total mixture at the end, to produce a visible ond-reaction. We then calculate for each trial the value $x = \frac{v_b - \beta F}{}$, and see whether there is a practically sufficient and up as hopcless; it may still remain worth while to see whether agreement between theory and

available area of experimental conditions within which the ratio a; b has a constant value. Or, what comes to the sams thing, we take the mean of all the RS (let it be $= \kappa_0$), and see whether the values of v_0 as calculated by the equation $v_b = \kappa_o v_a + \beta F$ agree sufficiently with the directly observed values. Should this not be the case, the process need not necessarily be given practice cannot be established by adding a term cx to the right side of the equation, where x stands for the weight of some subsidiary component x, and c is a positive or negative constant, whose value must of eourse be experimentally ascertained. In such eases, how-ever, it is better to leave the chemical significance of κ_0 , β , and c, entirely on one side, and to calculate them as so many empirical coefficients from the sum-total of the results. A formula thus obtained is of course of no practical value unless β and c are so small that an approximate determination of F and X suffices for an exact calculation of the respective terms. As an illustration we may quote Liebig's method for the determination of urea (A), in presence of chloride of sodium (x), by means of standard mercuric nitrato (v) as a pptnt. of the urea, and carbonate of soda as a drop-test for excess of pptnt. The exact volume v, of mercuric nitrate solution (i.e. weight of HgO) to reach the endpoint for a given weight (a) of urca, varies with the dilution, r, and the weight x of salt present; but ve is in sufficient accordance with equation $v_b = ak + \beta r + cx$, whose constants have been determined (virtually) by Liebig.

Nothing said so far is based on the presumption

1 Or at least is supposed to be.

that the exact chemical theory for the reaction between B and A is known. There are indeed a number of nseful volumetric processes which ars based upon unexplained, or only half-explained, chemical reactions. Fehling's method for the determination of glucose affords an illustration. If a dilute solution of glucose is dropped into a hot, strongly alkaline, solution of tartrate of copper (CuO) and potash, the CuO is reduced to (a pp. of) Cu₂O, the blue colour of the solution disappears, and the sugar suffers some unknown kind of oxidation. Yet the ratio between (say) dextroso oxidised and copper-oxide reduced, under specified conditions, is fairly constant, and the reaction is accordingly available for a fairly exact method for the determination of dextrose.

The well-known process of Clark for the determination of the hardness of a water by means of standard soap might be quoted as another example. But such purely empirical processes, however useful they may be for this or that practical purpose, are of little importance as auxiliaries of oxact analysis, which demands of a titration-process in the first instance that in any given case the question of its applicability can be decided a priori with at least a high degree of (if not with perfect) certainty. And this is possible only if tho process is based on a definite chemical equation which gives a qualitatively and quantitatively exact account of what is going on.

From the fact, however, that some equation, a+b=c+d, is in itself a correct theory of the action of a on n as resulting in the products c and D, it does not follow that the equation is a sufficient theory of the corresponding process of titration. Because experience shows that. in general, ready-mado c and b when mixed together produce a and B in accordance with the inverse equation c + d = a + b. Hence supposing we start with a parts of a and add more and more of n, the end reaction is reached only when a part (say qa) of A is transformed at the expense of qb parts of n, while (1-q) times (a+b) are still present in their original condition. Generally, q is a continuous function of the experimental variants (state of dilution, temperature, &c.), and the translatability of the reaction into a titrimetric process depends on the possibility of finding a sufficient area of conditions within which q is, at least prastacelly, equal to unity.

If one of the products (c and D) separates out as an absolutely insoluble pp., or escapes as a gas, the reverse reaction does not occur, and q becomes equal to unity; the apparent end-point is the real end-point of the reaction. Hence we should think that ppns. (we mean cases where that radicle in & which is really the thing to be determined, by uniting with the essential radicle in B, separates out as a pp.), should be pre-eminently suitable for volumetric application. Experience, however, shows that the reverse is true. Because in the majority of cases the pp. carries down more or less of one or other of the other reagents or products, and so disturbs the quantitative relations. Very often also a considerable excess of pptnt. is required to produce complete ppn, within a reasonalor time. Both difficulties (for example) pre-

sent themselves in the case of that reaction, BaX + SO,R₂=R₂X + BaSO₄, which is so largely used for the gravimetric determination of SO₄ and of Ba. The irregularities referred to 'can be set right (more or less easily) in the gravimetrio application of the reaction; to the volumetric application they are absolutely fatal. The number of ppns., indeed, which afford a basis for correct volumetric processes is extremely limited.

Cortain classes of double decompositions and oxidations, in which reverse reactions are prevonted by the great inherent stability of one of the products, are admirably adapted to volumetric processes. To give examples:

Any strong soid, XH, when mixed progressively with a solution of some strong base of the type ROH (ex. KHO, NaOH, Ba(OH),), is nltimately converted into normal ealt, XR, with formation of that highly stable substance water. The general reaction is XH + ROH = RX + HHO. and the end-point can in all cases be sharply deflued by means of a few drops of neutral litmus-solution as an indicator. Hence any acid (or rather the 'H' in any acid) may be accurately measured by means of a standard solution of, for instance, caustic potash; and any of the bodies ROH (or rather their 'OH') by means of a standard solution of (say) hydro-chloric acid. The latter method applies almost directly to the (soluble) carbonates, sulphides, cyanides, &c., of the alkali metals. All the carbonates &c. referred to can be measured indirectly by the combined application of the two standard solutions: we add first an excess of standard acid, and heat to expel the volatile acid (CO2,H2S, &c.), then colour with litmus, and titrate back with standard alkali, till the point of neutrality is exactly reached. By substituting aurine (in alcoholic solution) for litmus, the method becomes available also for magnesia (Tornö); and by using nitric acid as the standard XH, we can determine even oxide

of silver (Dittmar). What we said of carbonates, &c., in reference to the metallic radicles, R, holds for the ammonium salts of our acids, XH. To determine, for iustance, HCl or H2SO4, in the presence of ammonia (as the only base), we need only add a known execss of standard alkali, expel the liberated ammonia by evaporation, then add litmus, super-saturate by standard acid, hoil off the carbonic acid, titrate back with standard alkali until the point of neutrality is exactly reached, and balance the equivalents of base and acid used as reagents, against each other; the balance of base-conivalents measures the acid given for determination.

That this method of acidimetry applies also to cases where the base can be separated out by excess of standard alkali, as an acid-free pp., is obvious. Oxide of copper (given as CuSO, or other cupric salt) fulfils this condition in the sense at least that the acid ppd. at first as part of a basic salt, can be re-extracted by boiling with excess of alkali.2

And other irregularities such as for instance the variability of the ratio of sulphur to copper in ppd. sulphide of cupper.

"We will avail ourselves of this opportunity for referring to the process of fractional filtration as useful in

To pass to another example. There is a series of reducing agents a, the solutions of which, when mixed with a solution of iodine in iodide of potassium, are oxidised into products a, while the corresponding quantity of free iodine passes into iodide.

$$\left. \begin{array}{l} \textit{For example:} --\\ \textbf{I. } SO_2 + \textbf{H}_2O + \textbf{I}_2 = 2\textbf{H} \textbf{I} + SO_4\\ \textbf{II. } 2\textbf{S}_2O_3\textbf{R}_2 + \textbf{I}_2 = 2\textbf{R} \textbf{I} + \textbf{S}_2O_3\textbf{R}_2\\ \textbf{III. } A\textbf{S}_2O_3 + 2\textbf{H}_1O + 2\textbf{I}_2 = 4\textbf{H} \textbf{I} + A\textbf{S}_2O_3\\ \textbf{IV. } S\textbf{S}_2O_3 + 2\textbf{II}_1O + 2\textbf{I}_2 = 4\textbf{H} \textbf{I} + A\textbf{S}_2O_3\\ \textbf{V. } S\textbf{n}C\textbf{I}_2 + 2\textbf{H}C\textbf{I} + \textbf{I}_2 = 2\textbf{H} \textbf{I} + \textbf{S}\textbf{n}C\textbf{I}_4\\ \textbf{V. } \end{array} \right\} \begin{array}{l} \text{in solution.} \end{array}$$

Each of these reactions takes its normal course only nnder certain conditions, which, however, in cases I. to IV. at least, are easily established. All go on readily in the cold; and with all, starch solution is a safe and delicate indicator of excess of free iodine. Hence, to determine any of our reducers, a, we bring it into solution in the proper manner, add starch solution, and then drop in iodine solution from the burette until the blue colour of iodide of starch, which appears locally from the first, bocomes permanent on stirring. Supposing t c.o. of iodine solution to have been used, and one c.c. of it to contain $\tau \times 127$ mgms, of free iodine, the weight of R is $t \tau$ ($\frac{1}{2}SO_2$, S_2O_3 , $\frac{1}{2}As_2O_3$, 1Sb2O3, 2SnCl2), as the case may be. By theory, any one of our reducers might serve as a reagent for the measurement of free iodine; in practice sulphurous acid and alkaline thiosulphate work

According to Bunson, sulphurous acid acts normally on iodine, if it is diffused through at least 3,000 times its weight of (air-free) water. For the determination of free iodine he uses an aqueous sulphurous acid diluted to the extent stated, in combination with a standardised solution of iodine. The sulphurous acid is measured out by means of a glass-stoppered oylinder (or a narrow necked flask with one mark on the neck) holding some 100-200 c.c. To determine an unknown weight (x mgms.) of free iodine given as solution in HI or KI solution, we add the least number n of measures of the sulphurous acid water which suffices to decolourise the solution, then starch solution, and lastly, from the burette, standard iodine, until the blue colour becomes pormanent after addition of, let us say, t e.e. On the other hand, we ascertain the number, t_n , of e.e.s of standard iodine required for 1 measure-full of the sulphurous acid. Obviously, $nt_0 \tau \times 127 = x + t \tau \times 127$ (mgms. of I_2). Whence $x = (nt_0 - t) \tau \times 127$.

(mgms of L_2). Whence $x = (n_0 - 1) \tau \times 12t$.

So far Bunsen had done no more than translate an old process for determining SO₂ into a precise method for determining iodiue. His great merit was to see that, given a method for determining free iodine, we have an indirect mothod for the determination of any of the large number of oxidising agents for which a definite proportion of iodine can be substituted by the purely qualitative execution of suitable renotions. cases like that referred to. Instead of filtering off the CuO pp., we allow the nixture to cool, dinte to a known rolume, v.o., filter through a dxy filter, and fresaure off a known aliquot part of the filtrate, v c.o., for the titration. If v is sufficiently large, the volume of the CuO need not be taken into account; supposing for instance v = 500 c., and the pp, of CuO amonnts to 1 gm., the error introduced by neglecting its volume amounts certainly to no more than about 0.5 c.c., or 0.001 of the whole.

Thus, for instance, we may determine frebromine, iodate RIO₃, bromate RBrO₄, hypochlorite RClO, ozone O₃, by letting the respective substance act on excess of iodide of potassium solution, aciditying with hydrochloric acid, and then titrating the iodine liberated as above oxplained.

From the respective equations, we see that Br₂, or Cl₂, or RClO, or O₂, liberates I₂; and that RIO₃ or RBrO₃ liberates 3I₂.

The same principle obviously applies to all those peroxides which, when distilled with excess of hydrochloric acid, liberate a definite proportion of ohlorine. As examples: $MnO.O_x$ (when distilled with HCl) yields $x \times Cl$, of free chlorine, which when passed into iodide of potassium solution liberates $x \times I_y$ of iodine. Hence for overy

one I mgm. obtained, there was $\frac{1}{2\pi} \times (MnO.O_x)$

mgms. of that peroxide of manganeso. And similarly (to quote another ease which is known to work) every CrO_3 mgm. of chronic trioxide, liberating $3 \times \operatorname{Cl}_1$ ultimately yields $3 \times \operatorname{I}$ mgms. of iodino; or, in this case, every $1 \times \operatorname{I}$ mgms. corresponds to $\frac{1}{2}\operatorname{CrO}_3$, or to $\frac{1}{3} \times \operatorname{K}_1\operatorname{CrO}_2$, if the CrO_3 was present in this form. It is as well to mention that what the method in any case really determines is, not the respective species, but he I_2 -yielding radicle; the active oxygen in the $\operatorname{MnO.O}_2$, or the $\operatorname{Cr.O}_2$ -O₃ or RClO_3 ; the O_2 in the KlO_3 ; the one O in O₃, &c.

The applicability of the general method, however, goes further. As ferrous chloride is readily converted into ferric salt by free chlorine, we can determine an unknown weight of ferrosum (ferrous iron) (given as FeCl., FeO, FeSO, &c.) by distilling the respective substance with a weighed excess of potassium dichromate and hydrochloric acid, and collecting the chlorine in iodide of potassium, &c. Supposing we used $k \times K, C, C, O$, mgms. of this salt, the chlorine furnished by it is $6 \times k \times Cl$ mgms., and, if the iodine obtained at the end was $(nt_0-t) \times \tau \times I$ mgms., then $6k \times Cl \cdot (nt_0-t) \tau Cl$ must have been used by the FeCl., and consequently, $\{6k-(nt_0-t)\tau\} \times (Fe=56 \text{ mgms.})$ of ferrosum must have been present in the substance analysed.

Strictly speaking, all volumetric methods are empirical methods, in this sense, that the andamental ohemical equation is only an approximate theory of the process. Hence, unless we are sure that the error in the equation, considered as a theory of titration, is less than the unavoid. able error involved in the operations, to attain the highest possible degree of precision wo must standardis, our measuring reagent (if possible) by means of a known weight of the very thing (or radicle) to be determined, and both in the standardisation and the analyses we must maintain as nearly as possible the same conditions. To illustrato this, let us assumo we had to analyse a series of alkaline carbonates by means of a standard hydrochlorio acid. Ppn. of a known volume of the reagent by nitrate of silver, and weighing the pp. of AgCl (or the corresponding process of titration) would no doubt give the most exact result for the number of mgms. of HCl contained in 1 c.e. of the reagent. Yet it is better in our caso to standardise the

acid by means of a known weight of pure carbonate of soda, although this method, as one for the datermination of HOl, could not for a moment be compared with either silver process in point of inherent precision.

In now passing from generalities to the consideration of individual methods, we shall confine ourselves in the main to those methods which are applicable to whole classes of bodies. Under the head of each we shall briefly state what applies to it as a general method. For special applications of these methods, as for special methods generally, also in regard to technicalities, we must refer to the special hand-books.

I. Methods based on double decompositions.

Theoretically these processes are founded on equations of the form

ax + by = ab + xy

A, B, O, D,

where a and b are the constant radicles characteristic of the process. Here we have to distinguish between two cases :- I. o and D remain dissolved. Only a very few processes fall into this group. As an example, we may quote Liebig's process for the titration of NCH by neutral nitrate of silver. Large excess of potash is added, the liquid is diluted largely, and, after addition of a little NaCl as indicator, standard AgNO, (neutral) is dropped in until the cloud of AgCl becomes permanent, showing that the reaction 2KNC+AgNO₃=KAg(NC)₂+KNO₃, has been just completed. II. The characteristic product o = ab comes down as a pp. Of thesa nnmcrous processes, only those need now be noticed in which, on account of the absence of an end-reaction, and of a suitable indicator, the and-point cannot be recognised otherwise than by proving quite directly that the ppn. has just reached ite end. If the pp. settles readily, this can ba dona with comparative ease-in an obvious manner; but casily settling pps. are exceptional. It is more generally practicable to get the pp. to settle so far that it is possible to draw off a few drops of the clear top-stratum, and to examine them on a watch-glass by addition of a drop of B, or of a solution of A, or of any delicate reagent for a or B. If this method does not work, the only course left is, from time to time, to take out a little of tha mixture, filter it through paper, and examino the filtrate. One way of doing this is to put a drop of tha finid on a small double filter-paper, and to examine the lower filter by means of some reagent which strikes an intense colour with A or B as tho case may be. But such colour-fests are not always available, so that ordinary filtration must generally be resorted to. Each such filtration of course means a loss of a, and consequently ought to be done with a measured aliquot part of the whole, to enable one to allow for the loss by calculation. This, however, is apt to lead to errors; in practice it is better in the first trial to neglect the error, and in a second and third practically

to avoid it by aroung only a rew times near the end of tha process, when tha amount of upped. A has become very small. In any cass it is convenient to have a standard solution of some reagent, ax', by means of which to retrace ona's steps if an excess of pptnt. has been addad. This auxiliary solution is best adjusted so that it pps. exactly its own volume of n. Tha msthod of procedurs then assumes this form:—We add s, finally in small instalments of, say, 4c.c. each, until by the last instalment the end-point has been over-stepped; we then go back with a-solution, adding it in instalments of 2 o.c., until this reagent is in excess; wa then again apply s in portions of 1 o.o., &c., until we come to know that, say, v.o., of s is too little, while v+0.2 o.o. is an excess; or that (v+0.1 e.o.) ±0.1 c.c. may be adopted as the final result.

Of the vast number of precipitation-analyses which have been invented, only those founded npon mutual decomposition of solutions of Silver salts and haloids occupy the rank of precise methods. If (dissolved) chloride and (dissolved) silver-salt meet in a neutral or acid solution the whola of the potential chloride of silver is formed, and comes down as a pp. as

demanded by the equation,

RCl + AgNO₃ = AgCl + RNO₃. Upon this, and the fact that the AgCl (if sufficiently abundant) readily unites on shaking into a quickly settling pp., Gay-Lussac long ago founded his famous process for the determination of silver by standard NaCl solution, which process is directly translatable into an equally exact process for the determination of chloride by standard silver. The equation, however, is not an absolutely correct theory of either process. Gay-Lussac observed that if the silver nitrate and the sodium chloride are exactly balanced against each other, the clarified mixture gives a distinct cloud with either reagent. Hence to exactly complete the ppn. of (say) AgNO, mgms. by salt, we must add, not NaCl, but a trifle more, call it $(1+\alpha)$ NaCl mgms. And similarly, the complete ppn. of NaCl mgms. demands $(1+\beta)$ AgNO₃ mgms. The exact values of a and B vary with the experimental conditions, and are not susceptible of separate determination. Hence to determine an unknown weight, $x \times Ag$ mgms. of silver (if we do not care to neglect the correcting factors), all we can do is: (1) to add standard chloride solution-at last in very small instalments, corresponding to say 0.02 mgms. of silver each-nntil tho ppn. is exactly completed by, say, n x RCl mgms. as calculated from tha strength of the solution, and the quantity used. We then (2) titrate back with (very diluto) standard silver until the last drop no longer gives a cloud of AgOl, which will take, say, exAg mgms. Tha mixtura now is (practically) in the samo condition as if no silver had been added but the chloride diminished by $\epsilon \times RCl$ mgms.

Obviously the truth lies between x=n and

$$x=n-\epsilon$$
, and we may say $x=(n-\frac{\epsilon}{2})\pm\frac{\epsilon}{2}$.

Or, to put it somewhat differently; we have $2x=2n+(\beta-\alpha)n-\epsilon$. I. and $0=n(\beta+\alpha)-\epsilon$ I. If we know that $\alpha=\beta$, we should have $x=n-\frac{\epsilon}{2}$ exactly.

¹ Mohr's Lehrbuch der chemische-analytischen Türirmethode; 2, Fleistuer, Die Türirmethode; 3, Fleischer, Die Türirmethode, Finglish edition Volumetrie Analysi, translated by M. M. Pattison Mair; 4, Sutton, Volumetrie Analysi; 8, Freenius, Quentitates Austysie.

According to Mulder, if the silver, calculated as metal, amounts to about 1 gram, and is diffused throughout some 120 to 150 o.c. of mixture $(\alpha + \beta) = 0.001$, about. The explanation given in regard to ohlorides holds substantially for bromides, iodides, cyanides (NCR); eulphocyanidee (perhaps also for cobaltocyanides, and some other metallo-cyanides); in the caso of bromides, however, the numbers a and \$ are practically equal to nil (Stas), AgBr being even more insoluble in HNO, and KNO, &c. solutione than AgCl is; hence we may surmise that tho $(\alpha + \beta)$ for iodide is etill nearer to nothing. The cases of NC.R and NCS.R have not been investigated in this sense. Presumably the $(\alpha + \beta)$ for cyanide is greater than, and that for (NCS)R is about equal to, the value for chlorides.

Given (let us say) an alloy for which the percentage of silver is approximately known (say to ±0.5 p.c.); the exact determination of the noble metal by titration with standard chloride (c.g. NaCl) solution offere no difficulty; but with an alloy &c. of utterly unknown composition the procees even in practised hande is apt to be tedious. Practical assayers, indeed, never apply Gay-Lussac's method without having first made a preliminary assay by the old method of oupellation. Volhard, some years ago, invented a modification of Gay-Lussac's method, which, with a emall number of samples at least, is quicker even than cupellation, and, in all casee, is more accurate. He dissolvee a known weight (equal to presumably 0.5 grm. of silver) in nitric acid, dilutes moderately, boils off all the N2O, adds 5 c.c. of saturated iron-alum solution, and then drops in standard sulphocyanide of potaesium (or ammonium) until the red colour of Fe(NCS), becomes permanent. (The large quantity of indicator prescribed is necessary, or electhe end-reaction lacks delicacy.) For the doter-mination of chlorine (given as RCl), Volhard pps. the chlorine first by an oxcess of etandard silver, he then adde iron-alum, and (without removing the AgCl 2) titrates back with KNCSAg until the end-point is reached.

A very handy (but less exact and lese widely applicable) method, introduced by Mohr, may here be referred to. He brings the chloride into neutral or very feebly alkaline solution, and, after adding a few drope of yellow chromate of potassium, titrates with neutral eilver nitrate until the red colour of the Ag, CrO, becomee permanent. The method, if used as an empirical one, gives very good results.

STANDARD SUBSTANCES AND SOLUTIONS.

1. Standard silver .- Bost prepared by Stas's2 process (precipitation of a cupriferous ammoniacal colution of nitrate by added alkaline ammonium sulphito). The ppd. metal, after having been washed, first with ammonia in the

Our impression is that Mulder over-estimated the

¹ Our impression is that Mulder over-estimated the value.
² The writer finds that high precision can be reached only by removing the AgOl pp. before titrating hack with KNCS. (See Dittmar's Report on the Composition of Ocean Water, p. 4. ['Ogallenger' Memoirs,] Also his kerveises to Quantitative Analysis, section on Sen-vater.)
² Recherches sur les reupports réciproques des poids atomiques (Bruxelles, 1880); and Nouvelles Recherches, de. (Einzelles, 1855), or German translation of both works Aronstein (Leipzig, 1867).

broken up in a mortar into granules, again heated, and preserved in this form. There is no need of going to the trouble of fusing the metal. provided it is proved to be free from every traca of chloride.

presence of air, then with water, is heated to red.

ness, when it becomes semi-compact. It is then

2. Standard chloride. - Pure chloride of eodium is universally recommended. The writor prefers pure KCl prepared by strongly heating re-crystallieed perchlorato. The perchlorate is dooxygenated as far as convenient in a platinum basin, and then fully by fusion in a platinum crucible. The fused calt is quita neutral; yet for very precise work it is porhaps better to diecolvo the fused ealt in water, add hydrochloric acid, evaporate to dryness (in platinum), and keep the residue at a temporature just ehort of the fusing point until the weight is constant.

3. Standard solutions of 1 and 2.—Both can be standardised synthetically, on the basis of Stas' atomic weights; for general purposes Ag and KCl grms. per litre are convenient 10 10 strengths. For exact work the eclutions ara combined with centinormal colutions, containing $\frac{Ag}{100}$ and $\frac{KCl}{100}$ grms. per litre.

4. Pure bromide of potassium, and standard solution ($\frac{\text{KBr}}{10}$ grms. per litre) of the same for very precise determinations of eilver. Regarding the preparation of the pure salt, see Stas's Memoir.

5. Standard sulphocyanide. - About NCS.NH, grme. of the pure (chlorine-free) ammonium ealt is dissolved to 1 litre, and the exact strength is determined empirically by means of a known weight of silver dissolved as nitrato.

II. Methods based on saturations, that ie, reactions of the type

XH + ROH = HOH + XR:

regarding these, we have little to add to what was given under Theory of Titration (q. v.). For XH = NO₃H, \$SO, H₂, ClH, HClO₁, and other strong acids (including oxalic and formic) on the one hand, and R = K,Na, generally, and for $R = \frac{1}{2}$ (Ba, Sr, Ca) as long as no insoluble salt is produced, on the other, the equation is a strictly correct theory of the process. For phosphoria acid, HX mnst be taken as representing the H₂(HPO₄), RHO being an alkali, but even then the results are not very constant. For weaker acids, such as acctio, butyric, &c., the method is purely empirical. An approximation to a standard acetic acid is obtained by measuring off a known volume of standard sulphurio acid, and adding say two equivalents of perfectly neutral acetate of soda. In determinations of ammonia it is as well not to assume that NH,OH is an absolutely exact equivalent for KOH or NaOH.

STANDARD SUBSTANCES AND SOLUTIONS.

1. Pure carbonate of sodium, as a general standard alkali. Prepared from pure bicarbonate (recrystallised as such) by strongly heating in platinum. The salt must not be fused for dehydramest, or ease is loses carbonic said. To obtain a really pure, and especially a potashrice, salt, the best method is to add pure oxalic acid to a decided excess of solution of the purest obtainable carbonate of soda, to collect the ppof C₁O₂Na₂ formed, to wash it by displacement and to reduce it to Na₂CO₃ by heating strongly (W. D.).

2. Oxalic acid, C₂H₂O₄ + 2H₂O₅, recommended by Mohr as a general standard acid. See that the preparation is free from fixed matter (a.g. potassium salts). If not, recrystallise it from hot 10 p.c. HCl, repeatedly, and lastly from water (Stolba). The earcfully air-dried crystals have the correct composition. We prefer a hydrochloric acid, standardised by silver, for general

purposes.

3. Solution of standard acid.—Sulphuric works best for alkalis; hydrochloric is preferable for general purposes. The latter may be standardised by means of silver; either acid by means of a known weight of carbonate of soda, with standard alkali as an auxiliary reagent. Thorpe recommends for the standardisation of SO₄H₂, to add a known (excessive) weight of Na₂CO₃, to evaporate to dryness, heat, and weigh the residue. As every Na₂CO₃ grms. in passinginto Na₂SO₄, gains (SO₄—CO₃)grms., every

1 grm. of gain of weight corresponds to $\frac{SU_3}{SO_4-CO_3}$ grms. of sulphuric anhydride. (I have tested this method, and found it to give very good re-

sults.-W. D.)

- 4. Solution of standard alkali.—Caustie potash or caustic soda for general purposes. The reagent must be as free as possible from carbonate. The preparation known as potash purified by alcohol almost fulfils this condition. The best method is to causticise an almost carbonic acid free (dilute) ley with a stight excess of baryta in a nickel vessel; allow to settle, and preserve in a bottle provided with a protection-tube filled with granulated soda-lime, or baryta, Ba(OH)₂. The trace of dissolved BaO disposes of traces of CO₂ that find their way into the reagent while being preserved. 4ROH grms. per litro is a convenient strength. It is standardised empirically against measured standard acid.
- 5. Standard baryla water containing about Ba(OH), grms, per litre is used for special purposes, e.g. determination of free or liberated CO. A stronger reagent is apt to deposit crystals in cold weather. It is standardised empirically against standard hydrochloric acid. In the absence of sulphates, baryta water is the best standard alkali for all purposes.

III. Methods based upon processes of oxidation and reduction.

(As illustrated in *Theory* of *Titration*, by reforence to Bunsen's methods.)

I. Iodinc (solution of I in KI) as oxidant is available for the measurement of the following reducers:

1. Dissolved sulphurous acid acts normally under the conditions stated under Theory of Titration.

Ppn. with Ball, and weighing the BaSO, is not a very exact method for the standardisation of a sulphuric acid.

2. Dissolved alkali thiosulphate (in the absence of excess of alkali; even carbonate is not permissible) acts normally at any state of concentration which one could reasonably employ. In the presence of acid the reaction takes its normal course only if the solution is sufficiently diluted, and the H2S2O, has no time to decompose spontaneously. In practice, howover, this spontaneous decomposition is out of court, because, in all cases in which free acid is present, it forms part of the iodine solution, and the thiosulphate plays the part of reagont, so that the S2O3H2 liberated passes at onco into S₄O₆H₂, which is sufficiently stable. Free sulphuric acid in any quantity must be avoided (v. supra); free hydrochloric acid in moderate quantity does no harm, if the given iodino solution is diluted to about $\frac{1}{4}$ of the strength of the customary standard solution.

3. Alkaline arsenite. The reaction proceeds (not as promptly as those of 1 and 2, but) in a fair degree normally, provided there is onough of pure carbonate or bicarbonate of alkali to keep up an alkaline reaction to the end (Mohr). The best auxiliary alkali to add is sesqui-carbonate of ammonia; it does not decolourise iodide of starch, to anything like the (slight) extent to which Na₂CO₃ does (Mohr; later communication).

4. Alkaline antimonite, or rather Sb₂O₃ given as tartar emetic or in similar forms, is oxidised by iodine just as As₂O₃ is (Mohr).

Results fair.

5. Stannous chloride. The execution of the process (SnCl₂+2HCl+I₂=2HI+SnCl₄) offers no difficulty, and added starch solution defines the end-point sharply; but the results are very variable and inexact.

6. Sulphuretted hydrogen H_2S (in much air free water) with iodine feacts substantially thus: $-\mathbf{I}_2 + \mathbf{H}_2S = 2\mathbf{HI} + \mathbf{S}$. Results are only approximate, yet the method is of some value for determining small quantities of \mathbf{H}_2S in much water.

II. Iodine in combination with reducers for general purposes.

Only the combinations I₂ and H₂SO₃, and I₄ and Na₂S₂O₃ are used now-a-days. Discussion anticipated in section on Theory of Titration.

III. Permanganate of potassium, as an oxidant.

measures the following reducers directly, and in all eases the intense colour of the reagent marks the end-point with great sharpness.

1. Ferrosum. A dilute, strongly acid, solution of ferrous sulphato, when titrated with solution of permanganate, is promptly oxidised into ferrie salt with formation of Mn0-salt from the

reagent (Marguerite).

Conditions of success:—a, large dilution; 1 litre of solution should contain at most 1 gram of metallic iron; b, a sufficiency of free sulphuric acid, more than the equation demands, or else MnO₂ may separate out as a pp.; c, absence of hydrochloric acid (and chlorides generally), or clso part of the reagent is reduced by it with formation of Cl₂. According to Zimmerman, however, this by-reaction can be prevented by addition of manganous sulphate to the ferrosum solution. 4 grams of the sait MnSO₄+4H₂O₅

auffice per 60 c.c. of 20 p.c. HCl used for dissolving the respective iron compound.

Iron given as ferric salt may be reduced to ferrous salt, by (1) treatment with H.S.; (2) prolonged treatment in a warm solution with Na, SO, and HCl (works better with chlorido than with sulphate solution) -in either case the excess of reducer must of course be expolled by ebullition -(3) zine and acid; handy, but not so trust-

worthy as (1) or (2).

2. Oxalic acid. A strongly sulphurie solu-tion of this acid is oxidised by the reagent into oarbon diexide and water (Hempel). The reaction at first proceeds very sluggishly, but then more and more promptly, as the quantity of MnSO, produced increases. Hence the expediency of adding MnSO, from the first (De Koninck). Whether hydrochloric acid interferes with this reaction as with the preceding ono (whether, for instance, it is permissible to dissolvo oxalate of lime given for the determination of its exalic acid in hydrochloric acid before titrating) has not yet been determined.

3. Arsenious acid. Arsenious acid in strongly hydrochlorio solutions is oxidised by permanganate into arsenic acid, but part of the manganese

separates out as MnO2 (Kessler).

4. Antimonious acid as SbCl, in a solution which contains not less than 1-2 volumes of hydrochloric acid for 5 of water, is readily and completely oxidised into Sh.O.. The reaction is available quantitatively. [(3) and (4), Kossler, J. 1863. 683)].

5. Sulphurous acid is readily oxidised into sulphuric; not investigated quantitatively, as far

as we know.

6. Peroxide of hydrogen. In the presence of water and dilute sulphurie acid, the mutual

 $5H_2O.O + Mn_2O_7 = 5H_2O + 2MnO + 5O_2$ proceeds normally and promptly.

7. Nitrous acid (liberated from nitrito by H₂SO₄ in very diluto solutions) is oxidised by permanganate te nitric acid. Results, under certain conditions, fair.

8. Cu2O (ppd.) dissolved in acid iron alum. is oxidised readily, and fairly normally, to 2CuO.

IV. Combined application of permanganate and reducing agents.

A. Ferrosum as reducer.

The higher oxides of manganese, when digested with HCl or dilute H.SO, and excess of ferrous salt, are readily dissolved as MnO-salt, with formation of a quantity of ferricum proportional to the loosely held oxygen in the peroxide. In the absence of atmospheric oxygen, i.e. in an atmosphero of CO, the reaction takes its normal course, and becomes available for an obvious remainder-method for the determination of such oxygen. No doubt available for other peroxides.

Upon the ready action of alkaline permanganate on the sulphides, sulphites, thiosulphates, iodides, arsenites, formates, of K or Na, with formation of sulphate, iodate, arsenate, carbonate, respectively (and hydrated binoxide of manganese), Péan de St. Gillos (A. Ch. [3] 55, 374), has founded a general method for the determination of the respective acids by means of a standard solution of permanganato, and an

auxiliary solution of ferrous sulphate. After having carried out the required oxidation by means of excess of permanganate and a sufficicnoy of caustic potash, the mixture is acidified, the MnO, and surplns Mn₂O, reduced by addition of, first, acid, and then excess of ferrous solution, and finally the surplus ferrosum is titrated by addition of more of the permanganate solution.

B. Oxalic Acid as reducer.

Any higher oxide of manganese, MnO.Ox, when digested with oxcess of oxalic acid and sulphuric acid, is dissolved as MnSO, with formation of CO, from the reagent. The oxalio acid is used as a standard solution, and what remains over after the oxidation is determined volumetrically by permanganate. (Calcium oxalate may separate out as a pp.)

V. Chromic acid (in practice a solution of K2Cr2O7) as oxidant

is available for the direct titration of the following reducers:

1. Ferrosum. Ferrons sulphate or chloride, in the presence of free acid, is readily and completely oxidised by added bichromate solution. The latter may be standardised synthetically (or analytically by means of a known weight of dissolved ferrosum). The end point is recognised by means of ferricyanide as a drop-test. The results are in exact accordance with the chemical equation, oven in the case of hydrochlorio solutions (Penny; Schabus). An unknown weight of CrO₃ can be determined with equal exactitude by adding a known excessivo weight of ferrosum (as sulphate) to the previously acidified solution, and titrating back with bichromate solution.

The combination K_2Cr_2O , and ferrous salt is equivalent to that of Mn₂O₂K₂O and the same reducer. It is available likewise for the determination of As₂O₃ and Sb₂O₃ in strongly hydro-ohloric solutions. The solution is mixed with a (measured) excess of bichromate solution, and the mixture allowed to stand to give the oxidation time for completion; a known excessive quantity of ferrosum is then added, and its excess is titrated by means of bichromate (Kessler).

2. Sulphurous acid, Sulphuretted hydrogen, Stannous chloride, in mineral acid solutions, are readily exidised by CrO, into SO, S+H2O, SnCl,, respectively, and in all cases iedide of potassium and starch afford a sensitive indicator of excess of exidant, which sharply defines the apparent end-point of the process. unfortunately, the corresponding ratio of the reagents in no case coincides with that demanded by the respective equations, ner is it even constant in itself. I' aries according to the degres of dilution, the proportion of free soid, the quantity of absorbed air in the reagents, &o., in a manner which defies all calculation (Kessler, Mohr, Casselman).

VI. Ferrio ohloride, in combination with Stannous ohloride.

The oxidation of an acid solution of SnCl, by added ferric chloride proceeds very readily when the liquid is hot, and in fair accordance with the equation:

SnCl2 + Fe2Cl4 = SnCl4 + 2FeCl2; but the dissolved air of the reageuts is drawn

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into the oxidation, and the results are consequently irregular. If however (according to Fresonius) we start with a hot, strongly acid, solution of ferrio chloride, and at a nearly boiling temperature drop in stannous ehloride, tho process proceeds exactly as described by the equation, and the disappearance of the last trace of the yellow colour of the ferrio salt defines the cudpoint very sharply. In case of doubt, add a slight excess of SnCl₂, allow to cool, add starch, and titrate with iedine solution to determine the oxeess of SnCl2, and allow for it. According to the writer's experience the whole of these operations must be done in an atmosphere of CO2, else the results are not exact. Fresenius utilises the process for the determination of nitrie acid. The nitrate to be analysed is allowed to react with an excess of ferrous sulphate, strongly acidified by IICl, in an atmosphere of H or CO₂ first cold, then hot; the NO is boiled off, and the ferricum produced is determined by means of standard SnCl,. The ferrioum present as an impurity in the ferrous salt is determined by a blank experiment, and is allowed for: $6 \times Fe$ of ferricum produced, indicate $1 \times N_*O_*$ of nitric anhydride.

STANDARD SUBSTANCES AND SOLUTIONS, FOR THE PROCESSES OF OXINATION REFERRED TO.

1. Pure iodine is best made by Stas's method. Ordinary iodine is disselved in the minimum of a highly concentrated solution of iodide of potassium, and re-ppd. as far as pessible by dilution with water. The pp. is washed, dried first on a porous tile, then over CaN O. The dry product is distilled (or the small quantity needed for an experiment sublimed extempore between watch-glasses), the first instalments of vapour being rejected on account of possible contamination with water. From such iedine

2. A standard iodine solution might easily bo made by exact synthesis. But it is more economical and less troublesome to prepare this solution by approximate synthesis from ordinary good iodinc (5 grams of 12 dissolved in 10 grams of 1K and 10-20 c.c. of water in a mortar, and diluted to 1 litre, gives a selution of convenient strength); and to determine the exact strength by comparison with a known weight of pure iodine, by means of a thiosulphate solution of arbitrary strength. Supposing p mgms. of pure iodino weighted directly, and dissolved in IK solution, require t_0 c.e. of thissulphate for their decolonrisation, while n c.c. of the iodine-solution require t c.c.; then 1 c.c. of thiosulphate is equivalent on the one hand to

 $\frac{p}{t_0}$ mgms. of iodine, and on the other to $\frac{n}{t}$ c.c. of iodino-solution. Hence 1 c.o. of the latter contains $\left(\frac{pt}{t_g n \times 127}\right) \times I(=127)$ mgms. of pure

3. Thiosulphate solution. - Mado by dissolving 10 grms. of the pure salt, Na,S,O,.5H,O, in water, to 1 litre. It decolourises about its own volume of the above iodine-solution. The strength is determined empirically by means of the latter.

4. Arsenious acid.—Pure As2O, is to be had in commerce; but the best qualities even are liable to be contaminated with Sb₂O₂, powder is not hygroscopic.

 Arsenite solution.—4.95 grms. = As₂O₃ powdered As₂O₃ along with 11 grms. of p Na₂CO₃ (=30 grms. of orystals, Na₂CO₃ +10H are dissolved in a slanting litre flask in wat over a water-bath; after cooling, the liquid diluted to 1 litre. 1 c.o. = 0.1 x 'I,' 'Cl,' d mgms. [This solution, as a reagent, may ser for the direct titration of dissolved hyp chlorite — $2RClO + As_2O_3 = 2RCl + As_2O_6$. end-point is recognised by means of iodide potassium and starch paper; a drop of th mixture when placed on the paper produces blue stain only as long as the RClO is in exces

6. Standard ferrosum.—Fine pianoforte wir is sure not te contain more than 0.4 p.c. c impurities, and consequently may be assume to represent 0.998 ± 0.002 times its weight o real iron. A known weight is dissolved in HC or dilute H2SO,, in the absence of air, &c. More convenient is

Ferrous sulphate. - FeSO, 7H2O, ppd. from pure, hot, concentrated, acid solution, by alcohol. The ppd. salt is washed with alcohol, dried on bibulous paper, and finally by exposure to the 'dry' air of a room. The dried salt is sifted to remove lumps, again exposed to the air for a while, and bottled up for use. The exact percentage of iron is determined by strongly heating a known weight in a platinum erucible at the end in the presence of airuntil the weight is constant, and weighing the Fe₂O₃. This old preparation of Otte, according to the writer's experience, has a higher degree of stability in air than Mohr's salt $(\text{Fe}(\text{NH}_4)_2\text{S}_2\text{O}_6.6\text{H}_2\text{O})$

7. Standard oxalic acid.—The crystallised, normal, ammonium salt is the best standard oxalate for processes in which it serves as a reducing agent. If air-dry, it has exactly the composition $C_2O_4(NH_4)_2 + H_2O = 142$.

8. Standard permanganate of potassium .-An almost pure salt is to be had in commerce. Yet it is not pure enough to serve as a standard substance in itself. A convenient solution is obtained by disselving a little mere than 3-16 grms. (=5-6 grms. of ferrosum) in water, in a mortar, and diluting to 1 litre. The solution is standardised by means of a known weight of ferrosum or oxalate of anmenium, according to the object which it is meant to serve.

9. Bichromate of potassium.—K,Cr,O,. The pure salt is not difficult to obtain; but it is not casy to prove that it contains exactly 2CrO, for 1K.O. Besides, the uncertainty of the atomic weight of chromium is a difficulty. To prepare the salt for use, it is powdered and dehydrated by keeping it near its fusing-point for a time in a platinum basin. It is then fused at the lowest temperature, and allowed to solidify, when it broaks up spontaneously into small granular fragments, and thus assumes a convenient form for weighing.

10. Standard solution of bickromate of potassium.—A convenient concentration is $\frac{K_2Cr_2O_2}{60}$

4.92 gms. per litre. It may be standardised

synthetically; but for the reasons stated it is on the whole preferable to standardise the solution analytically by means of a known

weight of dissolved ferrosnm.

11. Standard ferric chloride. — Puro forric oxido is prepared by strongly heating forrous oxalate. It is dissolved, by prolonged digestion, in fuming HCl, and the solution is diluted to the right volume. Fe₂O₃ = 8 grms. per litre is a

convenient strength.

12. Stannous chloride (for No. 11). Pure granulated tin (approximately weighed) is boiled with pure HCl until a sufficiency of the metal is dissolved. The residual metal is weighed, to ascertain how much has passed into solution. For every 3 grms. of dissolved tin, the solution is diluted—with air-free water—to 1 litre. The solution decolorises about \(^1_2\) its volume of the iron solution, which latter serves for its standardisation. This solution is so prone to oxidise in the air that it must be restandardised expressly for each analysis.

W. D.

ANALYSIS, ORGANIC. Ultimate organic analysis is the determination of the elements present in an organic substance. Proximate organic analysis is the determination of the compounds present in a nixture, or of the

radicles present in a compound.

ULTIMATE ANALYSIS. Qualitative.

Carbon. If a substance blackens when it is probably contains carbon, in which case the black residue may be burnt away by heating to redness in air. As substance that does not blacken may nevertheless contain earbon. A more general method of detecting carbon is first to warm the substance gently with dilute sulphurio acid, in order to expel CO₂ that may be present as carbonate, and then to add several volumes of strong H₂SO₄ and some K₂Cr₂O₇; when the mixture is heated any organic substance will be oxidised, and the escaping CO₂ will give a pp. with lime-water.

Hydrogen. The substance is mixed in a tube with dry CuO or PbCrO, and heated to redness; water comes off and condenses in a cold tube. Very small quantities of water may be detected by passing the gases through a glass tube lined with P₂O₃, which will deliquesee. In these experiments carbon may be detected by passing the escaping gases into lime-water.

Nitrogen. The substance is heated with soda-lime and the nitrogen may then be given off as NH₄ and detected by its smell, action on red litmus, or famos with HCl. The soda-lime must be strongly heated before use, antil it no longer gives off NH₂. This test will not succeed with compounds rich in exygen. A more delicate test consists in heating the substance with potassium in a test-tube drawn out to a point. After deflagration, the mass is dissolved in water and examined for cyanide (Lassaigne, A 46, 367). This test is no applicable to diazo-compounds (Graebe, B. 17, 1178).

Chlorine. The chlorine is eliminated in the form of a chloride, the presence of which is detected by AgNO₂. The conversion into eliloride

11

can be effected: (a) by boiling with furning HNO₂; in the case of volatile substances, the operation must be performed in a scaled tube: (b) by boiling with alcoholic potash: (c) by mixing with quicklime and heating to redness: (d) by heating with H_sSO₄ and MnO₂.

Bromine and iodine may be detected by the same notheds.

Halogens may also be detected by fixing a lump of CuO to a platinum wire, dipping it into the substance, and heating first in the inner and then in the outer part of a Bunsen flame: a green coloni in the outer part of a Bunsen flame; a green

colour indicates halogens (Beilstein, B. 5, 520). Sulphur is detected by strongly heating the substance with a mixture of sodic carbonate and sodic nitrato, or sodic carbonate and potassio chlorate; and testing the product for sulphate. Or the substance may be fused with sodium free from sulphur in a test-tube, and the product examined for sulphide (Schönn, Fr. 8, 52, 399). Some compounds, such as albumen, give a black pp. of PbS when boiled with a solution of PbO in NaOHAq. Boiling HgCl₂ or ammoniacal AgNO₃ often give a black pp. of metallic sulphide.

Phosphorus may be detected by fusing with Na₂CO₂ mixed with Na_{NO2}, and examining the product for phosphate. Or the carbonised substance may be heated with magnesium powder; the product, containing magnesium phosphide, is luminous in the dark, and when moistened with water will give off PH₂ (Schönn, Fr. 8, 55).

Quantitative.

Substances containing no elements beside

carbon, hydrogen, and oxygen.

The substances are subjected to Combustion as proposed by Liebig (P.21, 1), hydrogon being weighted as H_2O and carbon as CO_p . The operation is performed in a closed or in an open tube.

Closed tube, combustion.

A tube of hard glass (diameter .5 inch) is drawn out as represented, the length being about 18 inches. It is thoroughly cleaned by washing with funing HNO, water, alcohol, and ether; and is then dried. Coarsely powdered oxide of copper, which has been prepared by oxidising tho metal, not by strongly heating the nitrate, and has been dried at a red heat, is poured in as far as a; a mixture of the weighed substance with finely powdered dry CuO is then p surou in, it fills up the space from a to b; the vessel (mortar or glass tube) in which the mixture has been effected is then rinsed with more finely divided CuO, and these rinsings are poured into tho tube and take up the space b to c; finally some coarse CuO is poured in, taking the space c to d. The tube is thon gently tapped to onsure free passage for gas from end to end. A tube containing CaCl2 is fixed by means of a cork to the open end of the combustion tube, and a bulb-apparatus (Liebig's or Geissler's) containing caustie potash (1 pt. KOH to 2 pts. H₂O) is attached to this, and in accurate experiments a drying tube containing CaCl, or solid KOH is placed beyond the potash-bulbs.

The oxide of copper at d is first heated to redness, and then the tube is heated at the other end; the gas-burners of the furnace are then

gradually turned on, at either end, so that a regular stream of bubbles passes into the potash bulbs. When the entire tube has reached a dull red heat, the potash selution will begin to be sucked back, owing to absorption of CO2; at this moment the point of the tube is broken off, and air is sucked through the entire apparatus in order that the gases still contained in the tube

of gas. At the end of one experiment the tube is quite ready for a second.

Liquids of high boiling point are analysed in the same way as solids, except that they are weighed in short open tubes; volatile liquids such as ether are best put into a bulb or V-tube, which is inserted between the oxygen apparatus and the combustion tube; the pro-



may be drawn into the weighed bulbs; in this operation a long glass tube, open at both ends, may be placed over the broken point of the tube to prevent furnace gases being sucked in. The calcium chloride tube and potash bulbs are weighed when cool: 10 of the increase of weight of the former is hydrogen, in of the increase of weight of the latter is carbon.

Open tube, combustion.

It is in every way better to make the combustion in an 'open tube,' that is a tube through which oxygen is continually passing.

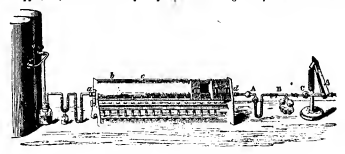
The greater part of such a tube is filled with

portion between oxygen and vapour of the substance depends upon the temperature of the bulb tube and should be so regulated that the oxygen should be always in considerable excess, otherwise an explosion might occur.

The potash-bulbs may be replaced by a U-tube containing soda-lime; in this ease the escaping gas must be allowed to bubble through H2SO, in order that the rate at which it is com-

ing off may be noted.

Minute quantities of carbonic acid are absorbed by CuO and even by PbCrO, and retained at a red heat. Hence in the determination of minute quantities of carbon (as in the residue oxide of copper, cd; this is followed by an open from drinking water) these substances should



space of about 2 inches; then cemes a percelain be previously ignited in a current of air (Dittmar or platinum boat, b, centaining the weighed substance; beyond (between b and a) it is advisable to have a spiral of oxidised copper. The boat and its contents are not inscrted until tho whole tube has been red hot for some time, during which a current of dry oxygen, free from carbonic acid, has been passing through it; it is of course necessary to allow the end a c of the tube to ceol down before inserting the substance, otherwise this would be volatilised too rapidly. While the tube is cooling, the calcium chloride tube A, potash bulbs n, and the drying tube c are attached. The tube, which is still red-hot frem c to d, is now heated at a and the burners are lighted one after another until the whole tube is red hot. A slow current of oxygen is passed in at a during the combustion. The combustion is continued until oxygen escaping from c will rekindle a glowing match. Before weighing, the oxygen in the tubes A, C, but especially n, must be displaced by air; in a properly conducted experiment e will not gain more than '01 g., a greater increase indicates spurting of the potash due to a too rapid current

a. Robinson, C. N. 36, 26).

Minute quantities of carbon may also be estimated by burning in an open combustion tube in a current of oxygen, in the usual way, and absorbing the CO, in baryta water. The BaCO, is filtered off, converted into sulphate and weighed (Dupré a. Hake, C. J. 35, 159). Other methods are described below.

Combustion with Chromio Acid.

Carbon may be determined by heating the substance with CrO, and H2SO, and measuring the mixture of CO and CO, given off (Cross a. Bevan, C. N. 52, 207).

Substances containing Nitrogen. Determination of Carbon and Hydrogen.

If the substance contains nitrogen, nitrous fumes might be evolved, and there would be absorbed in the weighed tubes. To prevent this, a layer of metallic copper is put in the front of the tube, near d, and kept red-hot: it reduces oxides of nitrogen to nitrogen. This copper is best abtained by beating a roll of wire gauze in a Bunsen flame, and reducing the oxidised surface in a current of hydrogen; it should then be allowed to cool in a current of CO₂ as it would absorb hydrogen if left to cool in that gas. The copper spiral may also be roduced by heating it in the mixture of CO and CO₂ obtained by warming oxalic acid with H₂SO₄ (C. E. Grovee, C. J. 37, 505).

Binoxide of manganeso mixed with potassic chromate may be used instead of a reduced copper spiral in combuction of nitrogenous subetances. The mixture is made by stirring precipitated binoxido of manganese with a saturated solution of potassio chromato containing a little bichromate; the paste is dried and heated somowhat strongly. The combustion is performed with plumbic chromate (or copper oxide) in the usual way, about 5 inches of the chromate mixture being put in the front part of the tube to absorb the nitrous fumes. In performing a combustion, the whole tube is strongly heated, while pure air is passed through it, then the absorbent mixture is allowed to cool to 200°-250° and kept at that temperature during the combustion (Porkin, C. J. 37, 457).

Estimation of Nitrogen.

Will and Varrentrapp (A. 39, 257) mix the substance with soda line, that has recently been strongly heated, and put the mixture into a short combustion-tube drawn out to a point at one end. The operation is conducted exactly as in combustion in a closed tube (v. supra), the escaping gases being passed into a bulb-apparatus to absorb ammonia. The bulbs contain hydrochloric acid, the NH₄ being weighed as (NH₃) ItCl₄; or, better, standard hydrochloric or oxalic acid, the amount of NH₄ being then determined by subsequent titration.

If the soda-lime contain nitrate it will evolve NH₃ even when heated with sugar (Schulze a. Kreussler, Fr. 12, 362). If in preparing the soda-lime a little Na₂S₂O₃ be added before evaporating and strongly heating, small quantities of nitrates and nitrites will be reduced and eliminated as NH₃ (Dittmar, priv. com.). Substances rich in nitrogen should be mixed with sugar after weighing.

Unfortunately many organic compounds do not yield all their nitrogen in the form of ammonia when ignited with soda-lime; such are nitroso-, nitro-, azo- and diazo-, compounds, and even some proteids (Ritthausen, Fr. 17, 501; Kreussler, J. 1884, 1608); in this case the method of analysis proposed by Dumas is generally used. Modifications of the soda-lime process intended to overcome this difficulty have, however, been proposed. Ruffle (C. J. 39, 87) mixes the substance (1 g.) with sulphur (75 g.) and finely powdered wood charcoal (75 g.). Scda-lime prepared from NaOli (160g.), water and CaO (56 g.) is dried and mixed with Na₂S₂O₃(21 g.). thirds of the tube is filled with this mixture, containing the substance to be analysed; the remaining third is ordinary soda-lime, which prevents evolution of H.S. The thiosulphate reduces nitro compounds.

Arnold (B. 18, 806) prefers a mixture of sods-lime, codium formate and Na₂S₂O₃.

Dumas (A. Ch. [2] 53, 171) heats the eub-

stance with oxide of copper and measures the escaping nitrogen. A combustion-tube closed at one end has first some bicarbonate of soda. or, much better, magnesito, put into it; this is followed by pure oxide of copper, a mixture of oxide of copper and the weighted substance, pure oxide of copper, and finally a bright copper spiral -just as in an ordinary combustion; the end of the tube is closed by a cork through which passes a delivery tube dipping under mercury. Before beginning the combustion all the air must be driven out of the tube by carbonic acid; this is effected by heating the magnesite; the combuetion is then proceeded with in the ordinary way, and the gaseous products are collected in a graduated tube standing over mercury and containing 50 c.e. of a solution of caustic potach (equal weights of potash and water). The products of combustion are water, earbonic acid, and nitrogen; the two former are stopped by the potash, so that the gas that collecte ie pure nitrogen; at the end of the experiment the combustion tube still contains nitrogen which must be expelled by heating the magnesite a second time. The cudiometer and its contents is then transferred to a vessel containing air-free water, which takes the place of the mercury and potash. The volume of the nitrogen corrected for pressure and temperature enables one to calculate its weight.

As there is some danger that the magnesite may be all used up in the preliminary expulsion of air from the tube, a convenient modification of this process consists in expelling the air by hydrogen; the hydrogen is got rid of by igniting a little of the copper oxide in the front part of the tube; a complete vacuum is thus formed, and the mercury rises in the delivery tube to the height of the barometer. The combustion is proceeded with in the usual way, and the residual nitrogen expelled by heating the magnesite.

Bicarbonato of soda, MnCO,, or a mixture of Na₂CO₃ and K₂Cr₂O₇ may be used instead of magnesite as a source of CO. In order that the tube may be used several times without turning out all the copper oxide, C. E. Groves (C. J. 37, 501) places the substance intended to evolve CO, in a separate tube, 7 inches long, which is attached by a short glass connecting-tube to the end of the combustion-tube, which is in this ease open at both ends. A fresh carbonic aoid tube is used for each experiment. The CO, may also be obtained from marble and HClay H.SO, but it is then liable to contain air nnlese the apparatus be first exhausted by an air-pump (Bernthsen, Fr. 21, 63) or heated to boiling (Hufschmidt, P. 18, 1441). The nitrogen is frequently contaminated with NO. Frankland s. Armstrong (C. J. 21, 77), after reading off the nitrogeu, pass up a little oxygen, and, when the resulting NO₂ has been absorbed, they remove the excess of oxygen by potassium pyrogallate. The mean between the volumes of gas before and after this operation is the true volume of nitrogen (Thudichum a. Wanklyn, C. J. 22, 293). Apparatus for collecting and measuring the nitrogen have been devised by Zulkowsky (A. 182, 296), Schwarz (B. 13, 771), Ludwig, (B. 13, 885), H. Schiff (B. 13, 885), C. E. Grovos (C. J. 37, 500), Staedel (Fr. 19, 452), Behmitt (J. pr. [3] 24, 444), Gladding (Am. 4, 42), Hempel (Fv. 17, 409), and Ilinski (B. 17, 1847). Frankland a Armstrong (C. J. 21, 77) connect the tube with a Sprengel's pump, which delivers the gas into a eudiometer at the end of the dropping tube (see also Gibbs, Fr. 11, 206; Hempel, Bn. 1, 9; Phüger, Fr. 18, 296; Johnson a. Jenkins, Am. 2, 27). Explosive substances may be analysed under diminished pressure.

The weight of nitrogen is calculated from its volume with the aid of the annexed table. From the barometric height, corrected for expansion of mercury and of the scale, the vapour pressure of water at the temperature of the nitrogen is deducted; from the corrected pressure and the temperature the weight of nitrogen is at once given by the table.

In the combustion of substances containing chlorine white fumes of ouprous chloride might pass into the chloride of calcium tube, and even chlorine might be given off by the action of the exygen (Städeler, A. 69, 335; Kraut, Fr. 2, 242); these sources of error may be prevented by placing a roll of silver foil between the copper exide and the cork into which the chloride of

(Arnold, Ar. Ph. [2] 24, 785). Kreusler recommends a mixture of conc. H₂SO₄ (9 pts.) and P₂O₅ (1 pt.) as a substitute for fuming H₂SO₄ (v. also Warington, C. N. 52, 162).

Substances containing Halogens.

Determination of Carbon and

Hydrogen.

Pressure of Aqueous Vapour, in mm.

Temp. | 10° | 11° | 12° | 13° | 14° | 15° | 16° | 17° | 18° | 19° | 20° | 21° | 22° | 23° | 24° | 25° | 27° | 28° | 10°5 | 11°2 | 11°9 | 12°7 | 13°5 | 14°4 | 16°4 | 16°3 | 17°4 | 18°5 | 19°7 | 20°9 | 22°2 | 23°6

Reduction of Barometric Height.

If the barometer has a glass scale, the necessary reduction will be found by multiplying the following numbers by the temperature, viz.:—

mm.	Reduction	mm.	Reduction
720	123	[] 750	128
725	·124	7 55	129
730	·125	760	·130
735	.126	765	·131
740	·127	770	·132
745	127	775	·133

Vapour-Pressure of Aqueous KOH.

When nitrogen is measured over aqueous potash, the correction for vapour-pressure is less than that given above, as is seen from the following table, which relates to a solution of 1 pt. of potash in 2½ pts. water (S.G. 1.258).

10°	6.19	li 18°	10.47
110	6.58	19°	11.20
12°	7.02	20°	11.97
13°	7.48	21°	12:80
140	7.99	22°	13.70
15°	8.53	23°	14.62
16°	9.13	24°	15.60
17°	l 9·77	25°	16.65

(Krensler, Fr. 24, 445).

In the course of an elaborate discussion of the satious methods of estimating nitrogen, Kreusler (Landwirthschaftliche Versuchstationen, 31,207; cf. Fr. 19, 92; 24, 438) recommends that the copper oxide be mixed with asbestos. Cuprio sulphate (150 g.), water (400 g.), gr.d light asbestos (50 g.), are evaporated until almost dry; the mass is then thrown in small quantities into boiling water (2500 g.) containing KOH (160 g.), and finally washed, dried, and heated until red hot. Kreusler also uses copper-asbestos prepared by reducing this copper-oxide-asbestos in place of a copper spiral.

Nitrogen may often be converted into NH, by KMnO, and boiling NaOHAq (Wanklyn, Chapman, a. Smith, C. J. 20, 445), or by KMnO, and fuming H₂SO, (Kjeldahl, Fr. 22, 370). In the latter case it is better to add CuSO, (Hilfahrt, C. C 16, 17), benzoio acid, sugar, and mercury (Schulze, B. 17, 1675).

calcium tube is inserted. This part of the tube is kept at a dull red heat throughout the combustion; cuprous chloride and silver form silver obloride and copper.

If the substance contains nitrogen as well as halogens, a copper spiral need not precede the silver spiral.

Determination of Halogens.

This is usually effected by placing 4 o.c. fuming nitrio acid and about a gram of silver nitrate in a strong glass tube, then sliding down a little tube containing the weighed substance in such a manner that it may stick to the wet glass and not at once fall into the acid; the open end of the strong glass tubo is then fused, drawn off to a stout point and scaled. A little tapping will now cause the tube containing the substance to fall into the acid, after which the whole is heated at 180° for seven hours in a gunbarrel. Aromatic substances require a higher temperature, 250°-300°. Silver chloride (bromide or iodide) is formed, and, after opening tho tube, diluting and boiling, it is collected, dried, and weighed (Carius, A. 116, 1; 136, 129). A still easier method is that lately proposed by Plimpton and Graves (C. J. 43, 119), in which the organic substance is burnt in the flame of a small Bunsen burner; the halogen is left chiefly combined with hydrogen but partly in the free state. The products are sucked through aqueous NaOH, which is then boiled with SO₂ and subsequently mixed with HNO₃. The halogen is then estimated volumetrically (best by sulphooyanide method) or gravimetrically.

Another method is to heat the substance in a combustion tube through which exygen charged with nitrous fumes is passing (Klason, B. 19, 1910).

Halogens may in many cases be determined by strongly heating with lime; with a mixture of Na₂CO₃ and KNO₂ (Volhard, A. 190, 40); with Fe₂O₃ (E. Kopp, B. 8, 769; Klobulowski, B. 10, 290); or with alcoholio KOH; or by reducing with sodium amalgam (Kokulé, A. Suppl. 1, 340).

Halogens in the side-chains of aromatic compounds may be estimated by boiling with a saturated alooholic solution of AgNO. (Schulze, B. 17, 1676).

1.05499

1-00001

1.06616

1.07166

1.07708

1.08246

1.08774

1.09304

1.09828

1.10346

1.10859

1.11369

1-11875

1.12376

1.12881

1.13380

750 744 753 748 1-13073 770 1.12770 768 1.07619 1-07316 1.07013 1.08225 1.07922 1.115581.11255 1.10649 1-10346 1.10043 1.09740 1.09437 1.08831 1.08528 1.06710 1.06407 1.06104 1.12467 1-12164 1.109521.09134 1.05801 1.11861 1.10924 1.13659 1.13355 1.10620 1.10316 1.10012 1.09708 1-094041.00100 1.08796 1.08493 1.08189 1.07885 1.07581 1.06973 1.06669 1.06365 1.12443 1.11532 1-12747 1.11835 1.11228 1.13051 1.12133 1.07277 Table showing the weight in milligrammes of 1 c.c. nitrogen at 10° to 25° C. and 720 to 770 mm. pressure (Dietrich, Fr. 5, 38) 1.13936 1.10886 1.10581 1.10276 1-09666 1-03056 1.03751 1.084.16 1.08141 1.07836 1.07531 1.07226 1.06921 1.142411.13021 1.12716 1.12106 1.11496 1.09971 1.09361 1.13631 1.13326 1.11191 1.12411 1.11801 230 1.07778 1.14818 1.14512 1-11145 1.10839 1.09308 1.08330 1.07472 1.14206 1.13900 1.13594 1.13288 1.12982 1.12675 1-12369 1.12063 1.11757 1:11:151 1.10533 1.10227 1.09921 1.09614 1-090021-08696 1.08084 ŝ 1.08015 1.09243 1.08936 1.08629 1.08322 1.13850 1.13236 1.10779 1.10472 1.10165 1.09857 1.09550 1.15385 1.15078 1.14771 1.14464 1.14157 1.13543 1.129281.12621 1.12007 1.11700 1.11393 1.11036 1.12314 210 1.08862 1.11943 1.09478 1.10139 1.09614 1.09083 1.08554 1.15640 1.14716 1.12559 1-12251 1.11018 1.10710 1.10402 1.10004 1.09786 1.09170 1.15948 1.153321.15024 1.14408 1.14099 1.13791 1.13483 1.13175 1.12867 1.116351.11327 8 1.09392 1.10320 1.1001.1 1.16504 1.16195 1-15267 1.14958 1.14649 1.14340 1.13412 1.13103 1.12794 1.12484 1.12175 1.11866 1-11557 1.11248 1.10938 1.10629 1.09702 1.15886 1.14030 1.15576 1.13731 61 1.11073 1.10544 1-09924 1.12095 1.11475 1.10854 1.10234 1.16746 1.16436 1.15816 1.14266 1.13955 1.13335 1.13025 1.12715 1.12405 1.11785 1.111651.170561.16126 1.15506 1.14576 1.13645 1.15196 1.14886 8 1.12006 1.11384 1.10450 1.17297 1.10985 1.13562 1.12629 1.11695 1.10761 1.17609 1.16674 1.16363 1.16052 1.15429 1.15118 1.14496 1.14185 1.13873 1.13251 1.12940 1.12317 1.15741 1.11307 170 1.17844 1.12845 1.12533 1.12220 1.10971 1.10658 1.18156 1-17531 1.17219 1.16906 1.16594 1.16282 1.15969 1-15344 1.15032 1.14720 1.14407 1.14095 1.13782 1.13470 1.13158 1.11908 1.11596 1.11283 1-15657 180 1.11486 1.18694 1.13366 1.12426 1.11799 1.11172 1.18381 1-18067 1.17754 1.17440 1-17127 1.16814 1.16500 1.16187 1.15873 1.15560 1.15247 1.14933 1.14620 1.14306 1.13993 1.13680 1.13053 1.12739 1.12113 120 1.12942 1.18920 1.18605 1.14515 1.13886 1.13572 1.19234 1.18291 1-17976 1.17661 1.17032 1.16718 1.10088 1.15459 1.15145 1.14830 1.14201 1.13257 1.12628 1.12313 1.11999 1.11684 1-17347 1.16403 1.15774 3 1.19452 1.18505 1-15663 1.13454 1-13138 1.125.36 1.12191 1.19136 1.18820 1.18189 1.17873 1.17558 1-17242 1.16926 1.16611 1.16295 1.15979 1.15348 1.15032 1.14716 1.14401 1.14085 1.13769 1.12822 1.19768 130 1-19344 1.16177 1.15543 1.14593 1.13010 1.12693 1.19977 1-19660 1.19027 1.18710 1.18394 1.17444 1.17127 1.16810 1.16493 1.15860 1-15227 1.149:0 1-14277 1.13960 1.13643 1.13326 1.20294 1.18077 1.17760 ន្ទ 1.13517 1.17650 1.17014 1.16696 1.16378 1.16060 1.15742 1.15494 1.15107 1.14471 1.14153 1.13835 1.13199 1.20511 1.20193 1-19875 1-19557 1.19239 1.18921 1.18603 1.17968 1.17332 1.14789 $1 \cdot 20829$ 1.18286 å 1.15613 1.14975 1.14018 1.21036 1.20717 1.20398 1.200791.19760 1.19441 1.18484 1.181651.17846 1.17527 1.17208 1.16889 1.16570 1.16251 1.15932 1.15294 1.14656 1.14337 1.13699 1.191221.213551-18803 န္ 736 742 740 738 734 754 750 746

Compounds containing Sulphur. Determination of Carbon and Hydrogen.

In the analysis of compounds containing sulphur there is danger that SO₂ may be absorbed in the weighed tubes. This is obviated by using lead chromate, in the form of small fused lumps, instead of cupric oxide. The PbCrO₄ (10 pts.) may be mixed with K₂Cr₂O₇ (1 pt.). Sulphur remains in the tube as PbSO₄. Volatile substances containing N as well as S must be burnt slowly (V. Meyer a. Städeler, B. 17, 1577). According to Ritthausen (Fr. 22, 108), reduced lead chromate can be re-oxidised by heating in a current of oxygen. It is of course not necessary to fill the whole tube with lead chromate, the posterior half may contain CuO. A mixture of CuO and PbCrO₄ is sometimes used.

Determination of Sulphur.

Sulphur may be estimated by the method of Carius, by heating with fuming IINO₂ in a tube as described for halogens; the sulphuric acid is ppd. by BaCl₂. This method cannot usually be employed in the case of organic sulphides, since these are converted into sulphonic acids.

The most rapid method is that of Plimpton, which consists in burning the substance in the flame of a small Bunsen, sucking the products of combustion through dilute NaOIIAQ, exidising with Cl, and ppg. as BaSO, (Morley a. Saint, C. J. 43, 401). Or the substance may be burnt in a stream of exygen and the SO, collected in HClAq containing Br (Sauer, Fr. 12, 32, 178; Mixter, Fr. 22, 581). In this experiment it is better to pass the exygen through fuming nitric acid so that it may be charged with nitrous funes (Klason, B. 19, 1910). In many cases sulphur may be determined by fusion with a mixture of Na₂CO₄ and KNO₄ or KClO₅. In the case of albuminoids it is convenient to evaporate with HNO₄ (10 pts. of S. G. 1·4) before fusing (Kochs, C. C. 1886, 894). Oxidation may also be effected by Na₂CO₃ and HgO (Russell, C. J. 7, 212).

Phosphorus.

The estimation of phosphorus resembles that of S. It is weighed as $Mg_2P_2O_7$.

Boron.

If compounds containing boron are analysed by combustion with copper oxide the hydrogen will come cut too high, owing to the volatilisation of boric acid. This is prevented by using lead chromate instead of cupric oxide (Councler, J. pr. [2] 18, 375).

Silicon.

SiO₂ is left behind when non-volatile compounds are heated. Velatile compounds are heated with cene. or fuming H₂SO₄ and KMnO; the product is poured into water. The ppd. II₂SiO₂ is freed from traces of manganese by fusion with Na₂CO₂ and KNO₂ (Polis, B. 19, 1024).

Alkalis or Alkaline Earths.

In an ordinary combustion these would be left as carbonates, the estimation of carbon thus being too low; but if the substance is thoroughly mixed with PbCrO₄ (10 pts.), and K₂Cr₂O₂ (1 pt.), the CO₂ will be turned out by CrO₄, the bases being left as chromates. Schaller (Bl. (2) 2, 93) mixes the alkaline salts of organic acids with an equal weight of SiO₂ and thon with CuO.

The alkaline metals are determined by strongly heating and analysing the inorganic residue.

Silver, Platinum, and Gold

Are determined by strongly heating the substance and weighing the metallic residue.

Explosive Substances

Must be well mixed with CuO before being put into a combustion tube; very explosive substances are analysed in vacuo.

Combustion with platinum black.

Kopfer (C. J. 29, 660) introduced the use of finely divided platinum as a carrier of oxygen. The anterior half of a combustion-tube is filled with platinum black intinately mixed with asbestos, the posterior half of the tube contains the weighed substance in a boat; air or oxygen is passed through the tube, the combustion being conducted in the usual way.

Additional References.

Gay-Lussac a. Thenard, A. Ch. 74, 47 (KClO₃); Saussure, A. Ch. 78, 57; Berzelius, Thomson's Annals of Philosophy, 4, 401, P. 44, 391; Liebig, P. 21, 1; Liebig a. Wöhler, A. 26, 270; Hofmann, C. J. 11, 30; Cloez, A. Ch. [3] 68, 394 (irou tube); Bl. [2] 1, 250; Fr. Schulze, Fr. 5, 289 (KClO₃); Wheeler, Am. S. [2] 41, 33; Marchand, J. pr. 41, 177; Gottlieb, A. 78, 241; Melsens, A. 60, 115; Frankland, T. 117, 63; Thorp, C. J. 19, 359; Maxwell Simpson, C. J. 6, 289; A. 95, 63; Pflüger, Arch. ges. Phys. 1878, 117; H. Schiff, A. 195, 293; Warren, Am. S. [2] 42, 156.

Determination of oxygen: Wanklyn a. Frank, P. M. [4] 26, 554; Baumhauer, A. 90, 228; Ar. N. 1, 179; Ladenburg, A. 135, 1 (AgIO₃); A. Mitscherlich, P. 130, 536; B. 1, 45; 6, 1000; Cretier, Fr. 13, 1; Stromeyer, A. 117, 247.

Simultaneous determination of C, II, and N: (Schulze, Fr. 5, 269; Frerichs, B. 10, 26; Hempel, Fr. 17, 409; Jannasch a. V. Meyer, B. 19, 949).

PROXIMATE ANALYSIS.

Separation of Mixtures.

A complete account of proximate organic analysis would include descriptions of the properties of every known organic substance, since the methods to be adopted vary in almost every case.

The simplest method is separation by solvents which dissolve some but uct all of the components of a mixture; when solid substances have been dissolved, they may usually be further purified by crystallisation.

An unknown mixture is treated with water; the insoluble portion is shaken with dilute H₂SO, which dissolves bases, then with dilute Na₂CO₅ which dissolves acids, then with dilute NaOH which dissolves phenols; the bases ars ppd. by adding KOH to the H₂SO₄Aq; the acids and phenols by acidifying the alkaline solutions containing them.

The neutral residue is fractionally distilled and the various portions are boiled with alcoholic KOH which saponifies: compound ethers, and gives a pp. of KCl in the case of fatty chloro-derivatives. The alcohol is distilled off together with volatile alcohols, &c., and the residue treated with water, which dissolves salts of acids that may have been formed. The insoluble residue is a hydrocarbon, high-boiling alcohol, alkyl oxide, ketone, haloid aromatic compound, &c.; the hydrocarbon will usually be left on treating the residue with cold H₂SO₄ if aromatic chloro-derivatives are absent.

The aqueous extract of the original substance is neutralised (if necessary) and distilled. The distillate is treated with K2CO3 which separates alcohols, lactones, methyl acetate &c. The residue is made alkalino by KOII and distilled: bases pass over; it is then acidified with H.SO, and distilled: volatile acids pass over. The residue is exactly neutralised and neutral substances are extracted by ether, olderoform, and benzone, the aqueous residuo is acidified and non-volatile acids are extracted by these solvents; the residuo is now made alkaline and again extracted, finally it is evaporated to dryness, and, if any organio matter is still present, extracted with solvents. (V. also Alkaloids, p. 120; Acros, p. 56.)

Many substances, even of very high boiling point, distil in a ourrent of steam.

Detection of radicles.

Hydroxyl.—Compounds containing OH evolve IICl when treated with PCl₂ (p. 54); but if PCl₃ is used chlor ration may occur e.g. C₆H₂OMe + PCl₅ = C₆H₄Cl.OMe + PCl₃ + IICl. AcCl acts upon hydroxylic compounds with evolution of IICl; when water is added to the product the acetyl derivatives of alcoholic hydroxyls are not affected, but CO.OAc is converted into CO.OH. AcCl acts also upon amines in the following way:

C₆II₈NH₂ + AcCl = C₆II₈NAcH + HCl.
BzCl and Ac₂O also displace hydroxylic hydrogen by acid radicles. In compounds whose molecules do not contain NH₂ or Nl1 the number of hydroxyls (other than those in carboxyls) present oan be determined by estimating the acctic acid obtained by sapouifying the product after treatment with water. The groups C.CO.NH and C.CO.ClI₂ in many cases act as if they contained hydroxyl and were C.C(OH):N and C.CO(OH):CH.

Very dilute, colourless solutions of Fe_zCl_d give a distinct colouration with compounds containing alcoholic hydroxyl (Landwehr, B. 19, 2726).

Zino ethide evolves ethane gas when mixed with compounds containing hydroxyl or amidogen (Japp, C. J. 37, 665); compounds containing imidogen evolve gas when heated with ZnEt, at 100° (Japp, C. J. 39, 224). (V. also Alcohols and Acids.)

Amidogen is indicated by the reactions just mentioned, and also by the readiness with which methylecan be introduced by heating with Mol. Methyl iodide does not act upon hydroxyl except in presence of an alkali. Compounds containing NH₂ evolve nitrogen when heated with nitrous acid.

The number of amidogens in the molecule of a compound may be found by treating the substance with KNO₂ and H₂SO₄; the escaping nitrogen is freed from NO by FeSO₄ and them measured (Sachsse a. Kormann, Fr. 14, 380). In easily diazotised aromatic amido compounds, the number of amidogens may be determined by dissolving in cone. HClAq and titrating with a normal solution of KNO₂, until a drop of the solution gives a blue colour with KI and starch (Green a. Evershed, S. C. I. 5, 633).

Imidogen is indicated by the production of a nitrosamine when treated with nitrous acid. (See also Amines, Amido-Acids, Amides.)

Carbon yl is indicated by the reaction with phenyl-hydrazino and hydroxylamine (p. 107).

Carboxyl is indicated by the easy production of metallic sults, and by the splitting off CO, when the substance is heated alone or with lime.

Want of saturation is indicated by instant bleaching of bromine.

Acetylenie hydrogen is indicated by the formation of explosive pps. with ammoniacal cuprous chloride or silver nitrate.

Nitroxyl.—Nitro compounds may be reduced to compounds containing anidogen. The number of nitroxyls is found by reducing with standard SnCl., and titrating with iodine (Limpricht, B. 11, 35).

pricht, B. 11, 35).

Methoxyl. The number of MeO groups can be found by boiling with aqueous H1 (S.G. 1 68) the distillate (Mel) being received in alcoholic AgNO₄ and the resulting AgI weighed (Zoisel, M. 6, 989).

Halogens in the side chains of aromatio compounds can be estimated by boiling with a saturated alcoholic solution of AgNO₃ (Schulze, B. 17, 1675).

ANAMIRTIN C₁₉H₂₁O₁₀(?). Occurs in grains of cocculus indicus (Anamirta cocculus) along with picrotoxin (n. v.), picrotiu, and glyceryl stearate (Barth a. Kretschy, Sitz. B. [iii] 81, 7; Francis, A. 42, 254). Short needles (from water), insol. benzene; becomes brown at 260°. Neither bitter nor poisonous.

ANCHOIC ACID = AZELAÏC ACID.

ANCHUSIN v. ALKANET.

ANDROMEDOTOXIN $C_{si}H_{si}O_{in}$ [228°]. **A** poisonous substance in *Rhododendron ponticum* (Zaaijer, R. 5, 313).

ANEMONE.—The plants Anemone nemorosa, A. pratensis, and A. pulsatilla when distilled with stam yield anemonin, anemone-camphor, anemonic acid, and a yei.owish oif, which are extracted by shaking the distillate with chloroform (Löwig a. Weidmann, P. 46, 45; Schwarz, Mag. Pharm. 10, 193; 19, 168; Febling, A. 38, 278; Dobraschinsky, J. Ph. [4] 1, 319; 11. Beckurts, C. C. 1885, 776).

Anemonin C_{1,1}H_{1,2}O₆ [150°]. Trimetric crystals; sol. CHCl₃, sl. sol. alcohol and water, insol. cther. Reduces AgNO₃. - PbOC₁₂H₁₂O₆.

Anemonic acid. Amorphous powder; insol. water, alcohol, and ether.

Anemone-compher. Trimetrio prisms; unstable.

ANETHOL C, H, O i.e. CH, O C, H, CH: CH, O C, H, CH: CH. CH, (1:4). Methyl p-properlyl-phenol. [21 3°]. (232°). S.G. 21 987. µn 1-6167 (Gladstone, C. J. 49, 623). R_∞ 77 97 (Nasini, G. 15, 93). S.V.S. 149·34.

Occurrence.- In oil of anise (from Pimpinella anisum) together with a terpene, in China or star anise (from Illicium anisatum), in fennel (Anethum fæniculum), and tarragon (Artemisia Dracunculus) (Laurent, Revue Scient. 10, 6; Gerhardt, A. 44, 318; 52, 401; Cahours, A. Ch. [3] 2, 274).

Formation. - Together with CO. CH,O.C.H. CH:C(CH,).CO.H is heated (Perkin, C. J. 32, 669).

Properties. — Plates; v. sl. sol. water, v. e. sol. alcobol and ether. Easily polymerised.

Reactions. - 1. Oxidation produces anisic aldehyde, anisic acid, and acetic acid (Hempel, A. 59, 104). — 2. Alcoholic potash produces $C_{1e}H_{1e}O_{2}$ [87°] and $C_{1e}H_{1e}O_{2}$ [65°]; the latter forms an acctyl derivative [40°] (Landolph, B. 13, 147; C. R. 81, 97; 82, 226).—3. Nitrosyl chloride produces C₆H₄(OMc).C₃H₃NOCl which reduces to C₆H₄(OMe).C₃H₆NII₂ (Tönnies, B. 12, 169).—4. Nitrous acid produces two bodies: C.H. (OMc).C.H. N.O. and C.H. (OMe).C.H. N.O. C.H. (OMc). C.J.H., N.O. and C.H. (OMc). C.J.H., N.O. 197°.

J. The former may be reduced to C.H. (OMc). C.J.H., (OII). M.J.III. the latter to (CH., O.C., H., C.J.H., 2N.O., (?) (T., B. 13, 1845).—5. Boron fluoride produces C.H., OMc, and a liquid C., H., (OC. 227°) (L., C. R. 86, 601).—6. HI at 260° forms C.J.H., (150°) and C., H., (210°) (L., C. R. 82, 849).—7. PCI, forms C., H., (10°) (C. 22°), S.G. 22° 1.191 (Landolph): Ladenburg gives different proper. (Landolph); Ladenburg gives different properties for chloro-anethol, viz.: [6°], (258°), S.G. 2 1.125.—8. Br forms C₁₀H₁₂Br₁O, [65°] (Ladenburg, A. Suppl. 8, 87; Z. [2] 5, 576).

Anethel - tetrahydride C₁₀H₁₀O. Anethol-

camphor. [190°-193°]. Has a camphor-like smell. Formed together with anisic aldehyde by the oxidation of anethol with HNO, (L.). Smells like camplior. On oxidation with K.Cr.O. it gives an acid which forms long needles of melt-

ing point [175°] (anisic acid?).

Anethol - hexahydride $C_{10}H_{18}O$. Anethol. borncol [19°]. (198°). Formed together with an acid by heating anethol-tetrahydrido with alcoholic KOH (L.). Long slender needles.

Insol. KOHAq.

Anethol dihydride = Methyl-Propyr-Phenol. Anethein (C₁₀II₁₂O)_a. Anisoin. [140°-145°]. From anethol by shaking with a little II SO, or P₂O₃ (C.), or by the action of SnCl, (Gerhardt, J. pr. 36, 267), I in KIAq (Will a. Rhodius, A. 65, 230) or BzCl (Kraut a. Uelsmann, J. pr. 77, 490). Prepared by distilling oil of anise with cono. H.SO. Needles (from ether); insol. water and alcohol. Decomposed on distillation into liquid metanethol and solid isanethol.

Metanethol $(C_{10}H_{12}O)_a$. $[132^{\circ 1}]$. (above 300°) Prepared by heating anothor (210 g.) with ZnCl₂ (750 g.) in a copper retort; in a few minutes white fumes appear in the receiver, superheated steam is then passed into the retort; the metanethol then passes slowly over. The yield is 5 to 10 p.c. (Perrenoud, A. 187, 63). Silky needles (from ether). Not volatile with

steam at 100°.

Metanethol sulphonic acid (C₁₀H₁₁(SO₂H)O). Formed by cono. H₂SO₄ in the cold. Salts.—CaA', aq: lamine.—BaA'₂.

Chloride C₁₀H₁₁(SO,Cl)O. [183°].

Liquid metanethol (C₁₀H₁₂O)_n. (233°).

S.G. H 971. Formed by distilling anethon

(Kraut a. Schinn, #, 1868, 859); and by distilling anethol with ZnCl₂. Converted by cono. H₂SO₄ into anethom. At 320° it partially changes to isanethol.

Sulphonic acid.—(C10H11(SO2H)O), (Gerhardt, J. pr. 36, 275). Salt.—CaA'2 aq: gummy. Isanethol (C10H12O)n. A viscid mass left in

the retort when anethoin is distilled; cone. H.SO, converts it into anethoin.

ANGELICA, OIL OF. The seeds of Angelica archangelica yield an essential oil (S.G. 2 872; [α]_D 13° 8') containing a terpene, $C_{10}H_{16}$, (175°), S.G. α 833, [α]_D = 12° 38'. The rotatory power of this terpeno is reduced to 4° 52' by heating for 432 hours at 100°; it is readily polymerised and easily oxidised (Naudin, C. R. 93, 1146). The essential oil also contains methyl-ethylacetic and oxymyristic acids (R. Müller, B. 14, 2476). The roots of Angelica archangelica yield an essential oil (S.G. 2 .875) containing a terpene (166°), S.G. $^{\circ}$.870, $[\alpha] = 2^{\circ}$ 50′. terpeno polymerises readily under the influence of heat or sodium (Naudin, C. R. 96, 1152; Bl. [2] 39, 406)

ANGELIC ACID C,H,O, i.e.

CH2:CH.CH(CH3).CO2H. Pentenoic acid. Mol. w. 100. [45°]. (185° i.V.).

Occurrence. — In the root of Angelica archangelica (Buchner, A. 42, 226). Oil of chamomile (Anthemis nobilis) may be separated by fractionating into isobutyl isobutyrate, isobutyl angelate, amyl angelate, and amyl tiglate; the residue contains hexyl tiglate and anthemol, C₁₀H₁₀O (Köbig, A. 195, 95). Angelic acid is formed, together with laserol by heating laserpitin with alcoholic KOH (Foldmann, A. 135, 236).

Preparation.-1. Angelica root (50 lbs.) is boiled with lime (4 lhs.) and water, and the filtrate acidified with H₂SO₄ and distilled (Meyer a. Zenner, A. 55, 317).—2. Oil of chamonile is saponified by alcoholic KOH (Kopp, A. 195, 81; Pagenstecher, A. 195, 108; Beilstein a. Wiegand, B. 17, 2261).-3. Sumbul or moschus root contains a resin which when boiled with alcoholic KOII yields angelic and methyl-crotonic acids (Reinsch, Jahrb. pr. Pharm. 7, 79; E. Schmidt,

Ar. Ph. [3] 24, 528).

Properties.—Monoclinic prisms or needles: sl. sol. cold water, v. sol. hot water, alcohol, and ether.

Reactions.-1. Hydriodic acid at 190° reduces it to valeric acid (Ascher, Z. [2] 6, 217) .--2. Potash-fusion produces acetate and propionato (Demarcay, C. R. 80. 1400) .- 3. Bromine forms a dibromido [86°].-4. Conc. H2SO, converts it into tiglic acid. -5. KMnO, gives CO, and aldehyde (B. a. W.). Salts. -- BaA', 4 aq: crystalline mass. --

CaA', 2aq: long needles, much more soluble in cold, than in hot, water. — AgA': fcathery crystals, sl. sol. water.—PbA'₂: crystals, sl. sol. water.

Ethyl-ether.—Eth. (1415°). S.G.º 935. Isobutyl ether C.H.A. (177°). Isoamyl ether C.H.A. (201°).

Oil (Chiozza, Anhydride (C,H,O)2O. A. Ch. [8] 39, 210).

Hydriodide CH,.CHI.CH(CH,).CO,H. [46°]. Iodo-valeric acid. Prisms. Formed by very conc. HI.

Constitution. - Angelic acid is isomeric with

allyl-acetic acid, CH,:OH,OH, CH, CO, H, methyl-erotonic or tiglic sold CH, OH:C(OH,).CO, H, 8-8-di-methyl-acrylic acid (CH₂)₂C:CH.CO₂H (Ustinoff, J. pr. [2] 34, 484), propylideneacetic acid, CH₂CH₂CH₂CH.CO₂H, and tetra-

methylene carboxylie acid.

The same valerie acid, (173'-175' uncor.), 8.G. 15 941, is formed by reducing the hydriodides of angelie and of tiglic acids by Zn and H.SO. It is probably CH3.CH2.CH(CH3).CO2H (Schmidt, B. 12, 252). Angelie acid changes when long kept, or when treated with AgNO, into tiglic acid, and hence the two acids have probably the same carbon skeleton. The hydriodide of angelic acid is, however, different from that of tiglic acid, CH3.CH2.CI(CH3).CO2H. This would be explained by assigning to angelic acid the formula CH., CH.CH(CH.), CO.H, its compound with HI being CH., CHI.CH(CH.), CO.H. The latter indevalence acid is not CH. L.CH., CII(CH.), CO.H for it gives no lactone on neutralisation with Na2CO3, but butylene CH_s.CH:CH.CH₃ (Fittig, A. 216, 161). An alternative formula, CH,:CEt.CO,H, would form with III either CH, I.CHEt.CO, II or, more probably, CH3.CIEt.CO2H; sodium carbonate would convert the latter into an oxy-acid, or back into angelie acid, while the former would give the butylene CII :: CII.CH 2. CH3.

ANGELICO BENZOIC OXIDE C₁₂H₁₂O₃ i.e. C₂H₁O.O.Bz. From potassium angelate and BzCl

(Chiozza, A. 86, 260). Oil.

ANGELICO LACTONES. C_sH_sO_x. β-acetylpropionic (levulic) acid splits up on distillation into H.O and a mixture of these lactones, transition compounds being doubtless the two isomeric oxy-acids CII₂.C(OH):CH.CH₂.CO₂H and CH₂.C(OH).CII₂.CO₄H. They are dried over K2CO3 and separated by fractional distillation (Wolff, A. 229, 249; B. 20, 425). The (a) lactono is converted into the (B) modification by combining it with HCl and distilling the product. Both lactones combine with bromine, and both are changed to β -neetyl-propionic acid by boiling with water or by treatment with cold aqueous baryta. This reaction is easily explained, for the oxy-acids CH2.C(OH):CH.CH2.CO2H and CH2:C(OH).CH2.CII ... CO2H into which the lactones should be converted, would both change (by Erlenmeyer's rule) into CH3.CO.CH2.CH2.CO21I.

CII, C; CII.CII, CO.O (a)-Angelico-lactone

[18°]. (168°). V.D. 3.6. S. 5 at 15°. Is also formed by the action of water on the bromido of B-bromo-B-acetyl-propionie acid, Colourless neutral liquid, gradually turns yellow. It has a pleasant odour and bitter taste. At 0° it solidifies to white needles which are not hygroscopic and are volatile. The lactone dissolves in most colvents. It is separated by K2CO3 from its aqueous solution. If left a few hours with celd water the liquid becomes acid.

Reactions.-1. NH₃ converts it into β-acetylpropion-amide.-2. Combines with bromine in CS forming the lactone of di-bromo-oxy-valcric soid (q. v.).—3. Combines with IICl forming the lactone of chloro-oxy-valerie acid (q. v.)

(β)-Angelico-lactone CH₂:C.CH₂·CH₂·CO.O. (209°) at 750 mm. (84°) at 25 mm. S.G. 21 1084. Colourless neutral liquid. It does not solidify at -15°. It is partly converted into its (a) isomer-

ide every time it is distilled under atmospherie pressure. Miscible with water, has a pleasant odour. May be left for 4 hours with cold water without production of an aoid.

Reactions .-- 1. Boiled with water it is very slowly converted into levulic acid. - 2. Combines with bromine in CS2-3. Does not combine with

ANGELYL. The radicle C.H.O. Also applied by Hofmann to monovalent Pententi

(q. v.)ANGUSTURA OIL C, H2,O. (266°). S.G. 93. Obtained by distilling true Angustura bark (Cusparia febrifuga) with steam (Herzog, J. 1858, 444). The bark contains also cusparine and gasipeine (q. v.).

ANHYDRIDES.—Oxides which react with water to form acids (q.v.), or are obtained from acids by withdrawing water, or which react

with basic oxides to produce salts: e.g.

 $SO_3 + H_2O = H_2SO_4$; $2HNO_3 - H_2O = N_2O_5$; $SO_3 + BaO = BaSO_4$; $CrO_3 + BaO = BaCrO_4$. Solutions of anhydrides in ether or other liquid quite free from water do not exhibit an acid reaction towards litmus. The greater number of the oxides of non-metals are anlivdrides; the metallic oxides which belong to this class are usually those containing the greatest quantity of oxygen relatively to the metal. The moro negative the character of an element the more do the lower oxides of that element exhibit the properties of anhydrides; the most positive clements do not form anhydrides. Metallie anhydrides do not, as a rule, produce acids by reacting with water, but most of them may be obtained from the corresponding hydrated oxides, having feebly-marked acid characters, by the action of heat; e.g. Nb2O5, PtO, PtO2, TiO2, SnO, &c. The formation of salts from these anhydrides is usually accomplished by fusing them with more basic oxides or hydrates; e.g. $Ta_2O_5 + K_2O$ (fused) = $K_2Ta_2O_6$; in some cases the metallie anhydride dissolves in strong aqueous potash or soda to form a salt; c.g

 $Au_2O_3 + 2KOHAq = K_2Au_2O_3Aq + H_2O_3$ nite connexion can be traced between the position of an element in the classificatory scheme founded on the periodic law and the existence or non-existence of anhydrides containing that element (v. Oxides; also Classification; and Periodic Law).

As a broad rule the anhydrides of the polybasic acids may be obtained from these acids by the action of heat (the anhydrides are usually the final products, before they are reached new acids are produced), but this rule has exceptions; e.g., $P_{2O_{5}}$ cannot be obtained by heating $H_{2}PO_{4}$. The anily drides of monobasic acids are usually obtained indirectly, often by the action of an acid chlorido on a salt, e.g. ClCl + HgClO = IlCl + HgClO = IlcHgCl + Cl.O (this method is largely used in preparing organic anhydrides, v. next article); these anhydrides are sometimes obtained from their acids by withdrawing water by the action of deliydrating agents; e.g. $2IINO_3 + P_2O_5 = N_2O_5 + P_2O_5H_2O$. A few anhydrides are produced by heating salts of the corresponding acids; e.g. FeSO, when heated in air forms Fe,S,O, and this on further heating gives Fe₂O₃ and 2SO₃.

Besides their characteristic reactions with water and basic exides, many anhydrides combine

with normal salts to produce 'acid' ealts (c. Salts); e.g. K₂OrO₄ + CrO₅ = K₂Or₂O₇.

K₂WO₄ + WO₅ = K₂W₂O₇. Many of the non-unstallic anhydrides combine with their own or other acids to form new acids; e.g. the following compounds are thus produced, (HNO₃)₂N₂O₄, H2SO4SO3, HCISO3, HSO2NO3, &c. A fow anhydrides react with basic oxides to form salts not of their own but of other acids; e.g., N₂O reacts with Na₂O₂Aq to produce NaNO₂Aq (not NaNO); such anhydrides seem to bo obtainable, indirectly, from mors than one acid, thus N2O is get by heating HNOAq or by heating solid NII, NO,

Most anhydrides may be regarded as constituted of two or mors acid radicles united by oxygen atoms; on this viow such formulæ as these would be applicable: -NO2.O.O2N; Cl.O.Cl, &o. The mutual relations between acids and anhydrides are repeated to a great extent in tho relations of basic hydrates, or hydroxides, to thoir oxides; e.g. Fe₂O₆H₆ when heated yiolds 3H₂O and Fe₂O₂; CaO when added to H₂O

produces CaO.H₂, &c.
At one time the name anhydrids included both base-producing and acid-producing oxides; then the formor class was distinguished as basic-anhydrides; but now the name is almost universally employed with the meanings given to it in this article. The following are the best marked anhydrides containing metals :- Sb2O3, Sb₂O₃, As₂O₃, As₂O₅; (? Bi₂O₃); CrO₃; (? Di₂O₃); Au₂O₃; Ir₂O₃; PbO₂; MnO₂; MoO₃; Nb₂O₅; OsO₄; PtO, PtO₂; Ta₂O₅; SnO, SnO₂; TiO₂; WO₄; UO₅; V₂O₄, V₂O₅; ZrO₂. The sulphides, are hydroughlyides of containing or hydrosulphides, of certain elements react with the auhydrides of thio-acids; e.g. As, S, dissolvss in KHSAq to form KASS, Aq, but the acid corresponding to this thio-arsenite is unknown; again, WS, dissolves in KHSAq to form K, WS, Aq, which reacts as the potassium salt of thiotungstic acid (H₂WS₁), which acid has not itself been prepared. The thio-acid corresponding to the thio anhydride SnS2, viz. H.SnS3, has been prepared; CS, again is the thio-anhylride of thio-carbonic acid H2CS3. It is not, however, customary to apply the term anhydride to any sulphides even when an acid, or a series of salts, can be obtained from them. M. M. P. M.

ANHYDRIDES, ORGANIC. - The anhydride of an organic substance is a body derived from it by elimination of water. The water may bs derived from one molecule, or several molecules may become united in the process; in the latter case 'condensation' is said to take place a term which is also used when closed chains are produced. A molecule of water may be derived:

A. From two carboxyls.

B. From one carboxyl and one hydroxyl.

C. From two hydroxyls.

D. From one hydrogen and one hydroxyl.

E. From carboxyl and amidogen. F. From carbonyl and amidogen.

G. From hydroxyl and amidogen.

A. FROM TWO CARBOXYLS. Acid Anhydridos.

Formation.-1. On application of heat most monocsrboxylic acids distil undecomposed, while di-carboxylic acids in which the carboxyls are attached to adjacent atoms of carbon give anhydrides, e.g.

 $C_2H_4 < \frac{CO.OH}{CO.OH} = C_2H_4 < \frac{CO}{CO} > 0 + H_2O$. When the oarboxyle are attached to the same atom of carbon, CO, ie split off :

 $CH_3 \cdot CH(CO_2H)_2 = CH_3 \cdot CH_2 \cdot CO_2H + CO_2$ 2. Anhydrides of monobasio acide are got by the action of acid chlorides on alkaline salts (Gerhardt, A. Ch. [3] 37, 285). Mixed anhydrides of monobasio acids may be got in the same way. Instead of the alkalins salt the frse acid may be used (Linnemann, A. 161, 169). An acid heated with its chloride gives its anhy. drids in the following cases amongst othere: acetic, trichloro-acetic, butyric, benzoic, and succinic acids. Acetyl chloride heated with dibasic acids gives anhydrides of the dibasic acid in the following cases amongst others (a mixed anhydrids is psrhaps first formed): succinic, oblorosuccinic, bromo succinio, maleio, acetyl-malio, diacstyl tartaric, diacetyl racemic, citraconic, itaconic, camphoric, phthalic and diphenio acids. These anhydrides of dibasic acids readily absorb water from the air forming the corresponding hydrates, from which, howsver, they may bs ssparated by chloroform which dissolves the anhydrides only. Benzoyl chlorids acts like acotyl chlorids. In neither case are mixed anhydrides formed.

Acctic anhydride at 120°-150° also converts dibasic acids into their anhydrides, e.g.: succinic, camphoric, phthalic, and diphenic acide (Anschütz, A. 226, 12).

3. Anhydrides ars also formed by the action of lead nitrate on acid chlorides (Lachowicz, B.

17, 1281), e.g.:

2Accl+Pb(NO₃)₂=Ac₂O+PbCl₂+N₂O₅.
4. By passing phosgene over heated salte (Hentsehel, B. 17, 1285):

 $2NaOAc + COCl_2 = Ac_2O + 2NaCl + CO_2$. 5. By warming the chlorides of the aoids with dry oxalic acid, e.g. 2Ph.COCl + H2C2O4 = $(Ph.CO)_2O + 2HCl + CO_2 + CO$ (Anschütz, A. 226,

Reactions.—1. Simple anhydrides usually be distilled or sublimed, but mixed anhydrides such as BzOAc are split up by heat into two simple ones: $2BzOAc = Bz_zO + Ac_zO$. 2. They are insoluble in water, but slowly converted by it into the corresponding acid; a conversion that is more rapidly effected by alkalis .-3. Alcohol forms ethyl ethers of the corresponding acids.-4. Ammonia forms an amide and an ammorrium salt: $Ac_2O + 2NH_3 = AoNH_2 + AcONH_4$ or, in the case of anhydrides of dicarboxylio acids, an amioacid .- 5. Sodium-amalgam raduces them to aldeliydes and alcohols (Linnemann, A. 148, 249). -6. PCl, forms POCl, and acid chlorides.-7. Heated with NaOAo in sealed tubes at 200° some anhydrides yield ketones (Perkin, C. J. 49, 325):

 CII_s . CO_s Na + (CH_s CO),O = CH_3 , CO, $CH_3 + CO_2 + CH_3$, CO_2Na , CH_3 , $CO_2Na + (C_3H_3CO)_2O =$ C_3H_1 , $CO.CH_3 + CO_2 + C_2H_1$, CO_2Na . Intermediats addition-products, such as CH₂.C(O.CO.C₂H₁)2.ONa, are perhaps the cause of this reaction.

B. FROM HYDROXYL AND CARBOXYL. This is the ordinary process of etherification: EtOH + $CH_1 \cdot CO_2 H = H_1O + CH_1 \cdot CO \cdot OEt$.

When hydroxyl and carboxyl are both present in the same molecule, spontaneous etherification may take place, the reaction taking place between two molecules, as in the formation of lactide, O < CHMo.CO > O, from lactic acid HO.CHMe.CO2H, or by'eplitting off water from one molecule, e.g.

 CH_3 .CH(OH). CH_2 . CH_3 . $CO_2H =$ CH₃.CH.CH₂.CH₂.CO.O.

In the latter case the alcoholic part of the molecule etherifies the acid part, and the product is called a lactone (v. Lactones). Hydroxyl in tho y and o positions gives riso to lactones.

C. FROM TWO HYDROXYLS.

The elimination of water between two hydroxyls in different molecules produces an oxide or simple ether. The result is brought about by first forming an intermediate compound, e.g.

 $EtOH + H_2SO_4 = EtO.SO_3H + H_2O$ EtO.SO,H + EtOH = EtOEt + H.SO, (v. ETDER). When two hydroxyls are attached to one atom

of carbon they usually split off water spoutaneously, producing carbonyl.

Two hydroxyls attached to contiguous atoms of carbon may give riso to an oxido or internal ethor, an intermediate body being first prepared,

 $HO.CH_{o}.CH_{o}OH + HCl = Cl.CH_{o}.CH_{o}OH + H_{o}O$ $Cl.CH_2.CH_2OH + KIIO = CH_2.CH_2 + KCl + H_2O.$

D. FROM ONE HYDROOEN AND ONE HYDROXYL.

Water can be climinated by the union of hydroxyl with hydrogen when they are attached This readily to adjacent atoms of carbon. ocours with \$\beta\$-oxy acids, e.g.:

 $CH_2(OH).CH_2.CO_2H = H_2O + CH_2.CH.CO_2H$ $\dot{C}_{a}II$, $\dot{C}H(\dot{O}H)$. $\dot{C}H$... $\dot{C}O_{a}H=$ $\mathbf{H}_{2}\mathbf{O} + \mathbf{C}_{6}\mathbf{H}_{4}$.CH:CII.CO.H.

E. From Carboxyl and Amidooen.

Water can be formed either from the hydroxyl and hydrogen, e.g.

Gody, and syntogen, e.g., $C_0H_4 < CO.CO.OH = H_2O + C_0H_4 < CO.CO.OH$ or from the oxygen of the carbonyl and hydrogen: $C_0H_4 < CO.CO.OH = H_2O + C_0H_4 < CO.CO.OH$ Compounds resulting from the first mode of dehydration are called 1 act a ms, those resulting from the second mode of dehydration below.

from the second mode of dehydration being termed lactims.

It is very difficult to say which formula best represents a given compound; Baeyer considers that the arrangement represented by the lactim is the more stable, but that before undergoing chemical reactions it usually changes to the transition or labilo condition represented by the lactam. The prefix pseudo is frequently applied to distinguish a lactam from a lactim. In the aromatic series the elimination of water takes place spontaneously when the two side-chains are in the ortho position, and when the condensation can produce a ring, containing 5 or 6 atoms.

Thus C.H. CH: CH: CH: CO.H

condenses to C₄H₄ CH: OH OH while C₂H₄ CH₂·CH₂·CH₂·CO₂H does not produce an appendix anhydrido.

F. FROM CARDONYL AND AMIDOOEN.

The case in which carbonyl forms part of carboxyl has already been montioned.

Mono-alkoyl-o-diamines and o-amido-alkoylphenols exhibit a tendency to split off water and form Anhydro compounds, e.g.

 $\begin{array}{ll} \text{Torm Analysis of compositions, e.g.} \\ C_{c}\Pi_{c}^{NHLCO.CH_{3}} = H_{2}O + C_{c}\Pi_{c}^{NH} \leqslant N \\ NH_{2} \\ C_{b}\Pi_{c}^{NLCO.CC_{c}\Pi_{3}} = H_{2}O + C_{c}\Pi_{c}^{N} \geqslant C.C_{c}\Pi_{5} \\ \text{(Hübner, A, 208, 278; 203, 339; 210, 328).} \end{array}$

first class of compounds may be viewed as amidines. These anhydro-compounds are formed: (1) From aromatic alkoylamides or alkyl phenols by nitration and roduction with tin and glacial acetic acid. (2) From [1:2] amido- (or oxy) nitro-compounds by heating with acid chlorides or anhydrides, and reducing the product. (3) From o-diamines or o-amido-phenols by heating with acid chlorides or anhydrides.

G. FROM HYDROXYL AND AMIDOOFN.

Alkyls can be introduced into amidogen by heating an amino with an alcohol or phenol, especially in presence of dehydrating agents: thus anilino boiled with (B)-naphthol gives phenyl-(\$)-naphthylamine.

ANHYDRO-ACET-DI-AMIDO-BENZENE v.

Ethenyl-phenylene-diamine

ANHYDRO - ACET - DI - AMIDO - BENZOIC ACID v. ETHENYL-DI-AMIDO-BENZOIC ACID. ANHYDRO-ACETYL. v. ETHENYL-

ANHYDRO-DI-ACETYL-ACETAMIDIL ACETAMIDINE.

ANHYDRO-DI-ACETYL-ACETAMIDINE

Acetambine ANHYDRO - o - AMIDO - PHENOL - ACETO -ACETIC ETHER v. Phopenyl. o-amido-phenol &-CARBOXYLIC ETHER.

ANHYDRO - AMIDO - PHENOXY - ACETIC ACID v. GLYCOLLIC ACID.

ANHYDRO-AMIDO-TOLYL-OXAMIC ACID

v. DI-OXY-METHYLQUINOXALINE. ANHYDRO-ATROPINE v. ATROPYL-TROPEÏN.

ANHYDRO-BENZ. v. BENZENYL-

ANHYDRO-BENZ-DIAMIDO-BENZENE . BENZENYL-PHENYLENE-DIAMINE.

ANHYDRO-BENZ - DIAMIDO - TOLUENE v. BENZENYI,-TOLYLENE-DIAMINE.

ANHYDRO-BENZ-DI-AMIDO-TOLVIC ACID v. Benzenyl-prenylene-diamine carboxylio aoid.

ANHYDRO-BENZOYL- v. BENZENYL-; or

named as derivatives of benzamidine. ANHYDRO BENZOYL - AMIDO-DI - TOLYL-

AMINE v. Benzenyl-tolyl-tolylene-diamine, ANHYDRO - CHLORO - FORMYL - AMIDO .

PHENYL MERCAPTAN v. CHLORO-METHENYL-AMIDO-PHENYL MERCAPTAN.

ANHYDRO-CINNAMOYL- v. CINNAMENYL-ANHYDRO-TRI-ETHYL-SULPHAMIC ACID v. Tri-Ethyl-amine

ANHYDRO-FORMYL- v. METHENYLnamed as derivatives of formamidine.

ANHYDRO-GLYCOLYL- v. OXY-ETHENYL-. ANHYDRO-LUPININE v. LUPININE. ANHYDRO-NAPHTHOL SULPHONIC ACID

v. Napothol-sulphonio acid.

ANHYDEO - OXALYL - AMIDO - PHENYL soluble spangles - B'HClAuCl : yellow needles MERCAPTAN C, H, N, S, i.e.

 $C_{\epsilon}H_{\epsilon} < N > C - C < N > C_{\epsilon}H_{\epsilon}$. [about 300°].

Formation .- 1. By heating anido-phenyl mercaptan with oxalic acid and PCl, -2. By the action of the chloro-methenyl-amide-phenyl mercaptan on methenyl-amido-phenyl mer-captan.—3. By heating chloro methenyl-amido-phenyl mercaptan with zinc.—4. By tho action of acetyl chloride or benzoyl chloride at 150° on methenyl-amido-phenyl mercaptan.

Preparation.—I. By heating acctantilide (5 pts.) with sulphur (3 pts.) to boiling for 30 bours; yield 25 to 30 p.c.—2. By leading (CN)₂ gas into an alcoholio solutiou of amido-phenyl mercaptan.

Properties .- Sublimable. Colourless glistening plates. Nearly insoluble in all solvents:

dissolves best in tolnene. Bitter taste.

Reactions.—By fusing with KOH at 200° it ie readily split up into amido-phenyl mercaptan and oxalic acid. On reduction with HI and Pat 150° it gives aniline and othenyl-amido-phenyl msrcaptan (Hofmann, B. 13, 1226).

ANHYDRO - OXALYL - D1 - PHENYLENE -

TETRA-AMINE $C_1H_{10}N_4$ i.e. $C_9H_4 < N_H > C.C < N_{NH} > C_9H_4$. [above 300°]. From o-di-nitro-oxanilide, Sn, and glacial HOAc (Hübncr, A. 209, 370). Yellow needles; insol. water, CS, and light p-trolenn, m. sol. glacial HOAc, sl. eol. alcobol, ether and benzene.

Salts.—B"2HCl 2aq.—B"H,SO, 2aq. ANHYDRO-OXALYL-D1-TOLYLENE-TETRA-

 $C_aH_a(CH_a) < N \\ NH > C - C < N \\ NH > C_cH_a(CH_a).$ [193°?]. Formed by heating oxalyl-di-tolylene-

diamine to above 200° (Hinsberg, B. 15, 2691). Or from di-nitro-di-tolyl-oxamide, Sn, and HCl (Hübner, A. 209, 373).

Salts. -B"H,Cl2 -B"(AcOH)2: glistening plates. - B", H,SO, 4aq : needles.

ANHYDRO-PHENYL-ACETYL- v. PHENYL-ETHENYL-

ANHYDRO - PHTHALYL - AM1DO-PHENYL MERCAPTAN C₂₀H₁₂N₂S₂ i.e.

$$C_{\bullet}H_{\bullet} <_{S}^{N} > C - C_{\bullet}H_{\bullet} - C <_{S}^{N} > C_{\bullet}H_{\bullet}.$$
 [112°].

Prisme or needles. Insol. water, sol. alcohol. Weak bass. Prepared by heating amidophenyl mercaptan hydrochloride with phthalyl chlorids. Salts. -- B'HCl: decomposed by water (B'HCl) PtCl4: slender needles (Hofmann, B. 13, 1233)

ANHYDRO-PROPIONYL. v. FROPENYL. ANHYDRO-PYROGALLO-KETONE v. Hexa-

OXY-BENZOPHENONE. ANHYDRO-SALICYL. v. OXY-nENZENYL-ANHYDRO - SUCCINYL - AMIDO - PHENYL

MERCAPTAN
$$C_{i_8}H_{12}N_{2}S_{2}$$
 i.e.

$$C_{i_8}H_{i_8} \searrow C_{i_8}C_{i_8}C_{i_8}C_{i_8} \searrow C_{i_8}C_{i_$$

Prepared by the action of succinamide on amidephenyl mercaptan. Colourless needles. Dis-solves in acids forming unstable salts. Vory stable towards reducing agents. By fusing with KOH amido-phenyl mcreaptan is reproduced. Salts.—BHCl: yellow nesdles, decomposed by water.—(B'HCl), PtCl,: eparingly

(Hofmann, B. 18, 1231).

ANHYDRO-SULPHAMIDO- v. SULPRO-ANHYDRO-TOLUYL-DI-AMIDO-BENZ**ENE** v. Toluenyl-phenylene-plamine.

ANHYDRO. TOLUYL-DIAMIDO. TOLUENE v. TOLUENYL-TOLYLENE-DIAMINE.

ANHYDRO-VALERYL- v. PENTENYL-.

ANIL- v. PHENYL-IMIDO -. ANIL - ACETOACETIC ACID- v. p. 19.

Reaction 18.

ANIL-BENZYL-MALONIC ETHER C₂₀H₂₁NO, i.e. C₆H₃.N;C(C₆H₃).CH(CO₂Et)₂, [75°]. Formula by the action of exo-chlore-benzylidene-aniline C.H. CCl:NPh upon sodio-malonie cther (Just B. 18, 2624). Large crystals; v. sol. alcohol and other, insol. water. It contains a hydrogen atom readily displaceable by sodium. Heated with dilute HCl at 120° it is split up into acctophenone, aniline, ethyl chloride, and CO. By heating alone to about 156° it eliminates alcohol and is converted into (Py. 1:3:2) oxy-phenyl-quinoline-oarboxylic ether

C₆H, C(OH):C.CO₂Et.

Di-anil-bsnzyl-malonio ethor C33H30N2O4 i.e. (PhN:CPh).C(CO.Et)... Formed by the action of exo-chloro-bouzylidene-aniline upon the sodium compound of mono-anil-benzyl-malonic ether (J.). Plates. By heating with dilute HCl or H,SO, at 120' it is split up into benzoic acid, anilins, acetic acid, ethyl chloride, and CO.

ANILIDES. - Substances derived from acids by displacement of the hydroxyl by phenyl-ami-dogen (NHPh). They are usually described under the acids to which they belong. The term amilide may also be applied more generally to

phenyl-amides and phenyl-imides. Anilides of acids.

Formation. -1. From aniline and acid chlor $ides: C_aH_aCOCl + NPhH_2 = HCl + C_aH_aCO.NPhH.$ 2. By boiling amides with the equivalent quantity of aniline until no more NH, comes off, and purifying by washing with other (Kelbe, B. 16, 1199). X.CO.NH. + NPhH. = X.CO.NPhH + NH. 3. In some cases, e.g. formic and acctio acids, anilides are formed by simply heating aniline with the dry acid (cf. Tobias, B. 15, 2866).— 4. By action of aniline upon compound cthors .-5. By the action of aniline on acid anhydrides.

· Properties .- Solid crystallins substances, v. sl. sol. water.

Reactions. - Split up into acid and aniline hy boiling aqueous or alcoholic KOH, by heating with HCl in a sealed tube, or by heating with oone. H.SO, at 100°.

Anilidee of phoephorous acid.

Tri-anilide P(NHPh), v. Aniline, react. 29. Di-anilide P(NHPh), (OH). Prepared by heating a mixture of aniline (3 pts.) and PCI. (1 pt.), extracting with etbor and ppg. with water (Jackson a. Menks, Am. 6, 89). White amorphons mass; eol. alcohol and ethor.

Anilides of phosphorio acid.

[208]. Tri-onilide PO(NHPh), [208]. From aniline and POCl₃ (Schiff, A. 101, 302; Michaelis a. Soden, A. 229, 335). Thin neodlss or eix-eided trimetric plates (from alcohol). Insol. water, aqueons acids, or alkalis. Forms a hexa-bromo derivative, [258°].

Di-antitids PO(OH)(NHPh), [1979]. From aniline (2 pts.) and POCl₂ (1 pt.), the product being treated with water (M. a. S.). Insol. water. Saponified by water or acids, not by

Anlide of thic-phosphoric acid PS(NHPh), [78°]. From PSCI, and aniline (Chevrier, Z. 1868, 539). Insol. water.

Anlide of arsenio acid AsO(OH)₂(NHPh).

Formed by heating aniline arsenate (Bechamp, C. R. 56, 1172).

Anilide of boric acid B₂O₃NPhH₂(?). From ethyl borate and anilino. Decomposed by water (Schiff, A. Suppl. 5, 209).

ANILIDO- v. PRENYL-AMIDO-

ANILINE C.H., N i.e. C.H., NH₂. Phenylamine. Mol. w. 93. [-8°] (Lucius, B. 5, 154). (185° cor.). (Private communication from R. J. Friswell); (183·7°) (Thorpe, C. J. 37, 221). S. G. \$\frac{1}{2}\text{ 1·0379 (T.).} \frac{2}{2}\text{ 1·0216 (Brühl); \$\frac{1}{2}\text{ 1·0212 (Friswell). C.E. (0°-10°) ·000866; (0°-100°) ·000925 (T.). (14°-25°) ·000818 (F.). H.F.p. -17450 (Thorpeon). 2747 (Reugen). -17450 (Thomsen); 2747 (Ramsay). H.F.y. -19190 (Th.). \(\mu_{\textit{B}}\) 10643 (B.). R \(\infty\) 49-83 (B.). S.V. 106-37 (T.); 106-08 (R. Schiff, B. 19, 566); 100 1 (Ramsay). Vapour pressure: Ramsay a. Young (C. J. 47, 647, 655). S. 5 at about 15°; the S.G. of the saturated aqueous solution is 15 1.0023; 21 1.001. 100 pts. of a solution of (W. Alexcieff, B. 10, 709). Aniline saturated with water has S.G. 4: I-025 (Friswell).

Formation.—1. Discovered by Unverdorben and the saturated with water has S.G. 4: I-025 (Friswell).

(P. 8, 397) among the products of distillation of indigo, and ealled by him crystalline .- 2. Rediscovered in coal tar by Runge (P. 31, 65, 513; 82, 331) and called by him cyanol. -3. Obtained by distilling indigo (from Indigofera 'Anil') with potash (Fritzsehe, J. pr. 20, 453; 27, 153; 28, 202) and then first called aniline.—4. Obtained from nitro-benzene by reducing with 250.—5. In supports animal oil (Anderson, A. 70, 32).—6. Be dary distillation of amido-benzoic acids (Hofinain a. Muspratt, A. 53, 221).—7. By distillang isatin with potash (Hofinain, A. 58, 11).—8. From di-phenyl-urea or di-phenyl-thio-urea by action of P₂O₃, zino chloride, or HCl (Hofinain, Pr. 9, 274): CO(NPhII).—9. NPhH₂+CO:NPh.—9. Among products of distillation of Peak (Yoll, J. Ph. [3] 36, 319).—10. By heatin ssium benzeno sulphonate with sodamide so son a. Wing, B. 19, 902).—11. By the act as Br in alkaline solution upon benzamido (Hoffinain Br. B. 18, 2737).—12. From phenol, and N. B. 19, 902).—1 immine by he amine by he amine by he amine by he addition of N. B. 18, 2737).—12. From phenol, and N. B. 18, 2737).—12. From phenol with 4 ph. 18, 18, 2737).—19. The addition of N. B. 18, 2737).—19. From phenol with 4 ph. 18, 18, 2737).—19. The addition of N. B. 18, 2737).—19. From phenol with 2 phenol with 2 phenol with 3 phenol with 3 phenol with 4 ph. 18, 18, 2737).—19. The addition of N. B. 18, 2737).—19. From phenol, and N. B. 18,

5,15

with a mixture of NH Cl and ZnO; also in this case an excess of NH Cl diminishes the quantity of secondary amine formed. The best yields (c. 55 p.c. aniline and 20 p.c. diphenylamine) are obtained by heating 2 pts. phenol with 2 pts. ZnO and 3 pts. NH,Cl at c. 330° for 20 hours under pressure.—(c) About the same results are obtained by substituting Zn(NHs)2Br2 NII Br for the chlorides in (a) and (b). -(d) Together with diphenylamine by heating plienol with NH,Cl and MgO. A yield of 45 p.c. aniline and 20 p.c. diphenylamine was obtained by heating 20 pts. phenol with 8.8 pts. MgO and 24 pts. NH₁Cl for 40 hrs. at 340°-350°.-(e) Small quantities of aniline and diphenylamine (e. 4 p.e. aniline and 15 p.e. diphenylamine) are formed by heating phenol (1 pt.) with NH₁Cl (2 pts.) alone, at 370°-400° (Morz a. Müller, B. 19, 2901).—13. Diphenylamine heated with cone. HCl at 320° yields small quantities of aniline and phono!.

Preparation. - By reducing nitro-benzene with iron filings in presence of a small quantity

of hydrochlorie or aectic acid:

 $4\text{PhNO}_2 + 4\text{H}_2\text{O} + 9\text{Fe} = 4\text{PhNH}_2 + 3\text{Fe}_3\text{O}_4$ Nitrobenzene (100 pts.), water (40 pts.), iron borings (25 pts.), and HClAq (9 pts.) are mixed in a east iron vessel and the reaction started by admission of steam; more iron borings (90 pts.) are then slowly added. When the reduction is complete, limo is added, and the aniline (67 pts.) distilled over with steam. Pure aniline is best prepared from pure benzeue. Aniline may be purified by conversion into its acetyl derivative, recrystallising this from water, and saponifying it with alkalis or acids. Aniline phosphate is less readily soluble in water than o-toluidine phosphato (Lewy, C. J. 46, 46)

Properties.-Colourless oil; sl. sol. water. miseiblo with most other menstrua. Turns red in air. Soluble in aqueous solutions of aniline hydrochloride. Its aqueous solution does not change the colour of red litinus or yellow turmerio but it changes the violet colour of dahlia to green. Cougo red may also be used as an indicator (Julius, S. C. I. 9, 109). Aniline pps. ferrous, ferrie, aluminium, and zine, hydrates from their salts; it forms double salts with PtCl4, AuCl3, HgCl2, SbCl3, and SnCl4. It gives a brownish pp. with tannin. It congulates albumen. Potash, soda, and lime expel aniline from its salts. NH, is expelled when aqueous ammonium salts are boiled with aniline, but aniline is liberated when NH, is added to cold aqueous solutions of its salts.

Detection.—1. Very dilute aqueous solutions give a violet colour with bleaching powder; the colour is destroy. I by shaking with ether (Runge). 2. Extremely dilute aqueous solutions treated successively with bleaching powder and a drop of ammonium sulphide givo a roso colour (Jacquemin, Bl. [2] 20, 68). -3. A solution of aniline in cone. H₂SO₄ mixed with a little solid K₂Cr₂O₇ gives after some time a splendid blue colour; the reaction is hastened by gently warming (Beissenhirz, A. 87, 376).-4. If a drop of CuSO Aq is added to an aqueous solution of aniline an applo-green crystalline pp. is formed even if the solution is very dilute; in extremely dilute solutions a green coloration is produced

(Friswell).

Reactions.—1. Aniline vapour passed through a red-hot tube forms C, NH, HON, benzene, benzonitrile (Hofmann, Pr. 12, 383), oarbazol (Graebe, A. 167, 125), iso-benzidine (v. di-AMIDO-DIPHENYL), and quinoline (Bernthssn, B. 19, 420)

2. Electric sparks passed through liquid anilino form earbon, and a gas containing hydrogen (65 p.c.), acetylene (21 p.c.), prussio aoid (9 p.o.), and nitrogen (5 p.o.) (Destrem, C. R. 99, 138).

3. Dilute H₂SO₄ and MnO₂ form NII₃ (Matthiessen, Pr. 9, 637), and a little quinone

(Hofmann, Pr. 13, 4).

4. Chromic acid, CrO3, sots fire to aniline. 5. Chromic acid mixture produces quinone.

6. Potassium chlorate and hydrochloric acid give tetrachloroquinone (chloranil) and trichlorophenol (Hofmann, A. 47, 67; 53, 28).

7. Potassium permanganate gives some azobenzeno (Giaser, A. 142, 364), NII, and oxalic aoid (Hoogewerff a. Dorp, B. 10, 1936; 11, 1202).

8. Hydrogen peroxide also produces azobenzene (Lecds, B. I4, 1384); which is also formed by passing aniline vapour over heated PbO (Behra. Dorp, B. 6, 755).

9. Potassium permanganate in acid solution

forms aniline black.

10. Strong nitric acid violently attacks aniline; picrio acid is among the products. Aniline nitrato dissolved in a large quantity of H2SO, produces m-nitro-aniline and a small quantity of p-nitro-aniline (Lavinstein, B. 18, Ref. 203).

11. A mixture of aniline, o-toluidine, and ptoluidine, is converted by oxidising agents such as nitric acid, mercuric chloride, lead nitrate, silver nitrate, arsenic acid, and stannic chloride

into aniline red (v. Rosaniline).

12. Nitrous acid converts cold aqueous salts of aniline into salts of diazo-benzene (v. Di-Azo-COMPOUNDS); on boiling the solution nitrogen is ovolved and phenol formed. Nitrous acid passed into a cold alcoholic solution of aniline produces liazo-benzene anilide.

13. When aniline is boiled with sulphur H.S. is evolved and di-amido-di-phenyl sulphide (thioaniline) is formed together with other products of substitution of hydrogen by sulphur (Merz a.

Weith, B. 3, 978).

14. Potassium produces NH, and azobenzene

(Girard a. Caventou, Bl. [2] 28, 530).

15. Chlorine acts upon dry aniline with great violence, producing a black mass containing trichloro-aniline.

16. Bromine behaves like chlorine. Brominewater added to solutions of salts of audine gives a pp. of tri-bromo aniline C, H. Br, NH2 [2:4:6:1]. Bromine has no action on a solution of anilino in conc. H.SO. (Morley, C. J. 51, 582).

17. Iodine dissolves in aniline forming hydriodide of p-iodo-anilinc.

18. Hot conc. sulphuric acid forms p-amidobenzene sulphonic acid; hot fuming H.SO, forms amido-benzeno disulphonio acid (Buckton a. Hofmann, C. J. 9, 260).

19. Sulphide of carbon forms di-phenyl thiopres

20. Sulphide of carbon and ammonia produce crystals of C₁₄H₁₈N₄S₃ or (PhNH₈.NH.CS)₂S₄ decomposed by boiling water into CS₂, NH₃, and di-phenyl-thio-ures (Hlasiwetz a. Kachler, A.

21. Carbon tetrabromide forms diphenyl-p-

amido-benzamidine hydrobromide

PhNH.C(NPh).C₆H₄.NH₂HBr (Bolas a. Groves, A. 160, 174). CCl₄ acts similarly (Hofmann, Pr.

22. Cyanogen forms a 'cyan-aniline' (C₆H₈NH₂),C₂N₂, [210°-220°] (Hofmann, A. 66, 129; 73, 180; B. 3, 763). Insol. water, sl. sol. alcohol. Boiling acids decompose it into phenyl-manida. oxamido, di-plenyl-oxamide, oxamido, aniline, and NH₃. Nitrous aoid produces a base C₁₄H₁₅N₅O₅ (Senf. J. pr. [2] 31, 543). Salts of Cyananiline: B"2HCl. — B"H₂PtCl₆. — B"2HAuCl .- B"2HBr. - B'2HNO.

23. Chloride of sulphur diluted with CS. forms tri-phenyl-guanidino (di-phenyl thiourea being first formed, Claus a. Krall, B. 3. 527; 4, 99).

24. Iloated with persulphocyanic heid it -

gives phenyl-thio-biuret. 25. Carbonyl chloride, COCl2, gives di-phenyl-

urea.

26. Cascous cyanogen chloride forms iii-phenylguanidine (melanilins); in presence of water
phenyl-urea is formed (Hofmann, A'' 70, 130).

CyCl passed into an ethereal soluti on forms
phenyl cyanamide (cyanamilide, Grahours a.

Cloëz, C. R. 38, 354). Solid chloride: if cyanogen
forms 'chlorocyanamilide' C₁₃H₁₂ClN₁₁ (Laurent,
4 Gn 973) A. 60, 273).

27. Cyanic acid forms phenyl-urecia.

28. When treated in othereal so ation with perchloromethylmercaptan CCl₃ SCl id t yields the compound CCl₃.S.NHC₆II₃. If the control the latter body is mixed will the alcoholio KOH or NH₃ it splits off HCl and 196 crystalline compound separates which has t 1 to probable constitution CCl₂S.NC₆H₃, (Rathke, I₃ 19, 395).

29. Phosphorus trichloride acts of vigorously, producing 'phosphaniline hyd Hrochlorida' PCl₃2C₆H₁N (Tait, Z. [2] i., 649). 13 it is perhapa the anilide of phosphorous acid P(N mchill),3HCl. PtCl₂PCl₃ forms (PhNH)₃PtCl₄(E_NNI,PhHCl) whence water produces (PhNH)₄N. PPtCl₄OH) (Quesneville, Monit. scient. [3, 6, 659). PCl₃PtCl₄ alcohol, and anility cats produce P(OEt)₃PCl₄(C₄H₁N)₄ and P₂(OEt)₄PCl₄(C₄H₁N)₄. (Cochin, C. R. 86, 1402).

30. Phosphorus orychloride pild goduces very unstable anilide of phosphorio acide it PO(NPhH). (Schiff, A. 101, 302; Michaelis a. S. st, PO(NPhH);

335).

31. Aniline, isobutyric acid, algorid ZnCl, give iso-butyric anilide (Bardwell, Am. Abrila 1116). 32. Aniline-zine-chloride and by 116). soamyl alcohol give amido-plienyl-isopeutane id.

hol give annoto-phenyl-isoperature id.

33. Chloroform at 130° for Antims di-phenylformamidine, CH(NPh)(NPhH).

34. Chlorides, bromides, and (3 todides of alcoholic or acid radicles act upon and mailline as they do npon other primary amines 6, (v. AMINES).

Alvyls may also be introduced by ol and heating aniline Altyis may also be introduced upol and heating annu-hydrochlorido or, better, hydrobr bid, omido with alco-hols (Stabdel a. Reinhardt, B. 12h). Methyl-anilino hydrochloride is oor 101 verted by heat into toluidine hydrochloride (1 n new Johnson, B. 5, 720); similarly aniline hydrochloride hated with MeOH at 200° forms Ford C.H.MeNMes. C.H. Me, NMe, C.H. Me, NMe, C.H.Me, NMe, and C.Me, (Holmann a. Martius, B. 4, 742).

85. Aldehydes act npon aniline with elimina-

tion of water: e.g. OH_2 . OH_2 . OH_3 . OH_4 . OH₄·CHO + H₂NPh = H₂O + CH₂·CH:NPh (Schiff, A. Suppl. 3, 344). The last formula onght perhaps to be written (CH3.CH)2(NPh)2 (v. METHYLENE-ANILINE; ETHYLIDENE-ANILINE; CHLORO-ETHYLIDENE-ANILINE, &c.). Aniline acts similarly upon glucose, levulose, and galactose, forming C₆H₁(OH)₅(NPh) (Schiff, A. 154, 30; Sorokin, B. 19, 513).

36. Aniline sulphite gives with an ethereal solution of allehyde prisms of PhNH₂C₂H₄OSO₂ or PhNH₂SO₂.CHMe.OH (Schiff, A. 140, 127; 210,

129).

37. Chloral and aqueous SO2 produce unstable crystale of PhNII, SO, CH(OII).CCl, (Schiff, A. 210, 129).

38. Acetone and aqueous SO2 form an unstable compound, PhNII₂C₃II₆O SO₂ which is perhaps Me₂C(OII). SO₂. NHPh (S.).

39. Aectone (1 mol.) and P₂O₅ two days at 180° form Mc.C:NPh, (200°-220°) (Engler a. Heine, B. 6, 642; cf. Pauly, A. 187, 222).

40. Aniline liydrochloride heated acetone or mesityl oxide at 190° forms some (Py. 1, 3)-di-metliyl-quinoline (Engler a. Riehm, B. 18, 2245, 3296).

41. A mixture of aldchyde and acetone at 100° give (Py. 1, 3)-di-methyl-quinoline (Beyer,

J. pr. [2] 33, 393).

42. Paraldchyde and cone. HCl at 100° form (Py. 3)-methyl-quinoline (quinaldine, Doebner a.

von Miller, B. 16, 2464).

43. Aniline (1 pt.) distilled with glycerin (1 pt.) and H₂SO₄ (2 pts.) forms quinoline (Königs, B. 13, 911). Quinoline is also formed by distilling aerolein-aniline, or by heating aniline with glycerin, nitrobenzene, and H.SO (Skraup, M. 2, 141). Aniline-zinc-chloride heated with glycerin forms skatole (Fischer a. German, B. 16, 710).

44. Aceto-acetie other at 120°-150° gives aceto-acetic anilide CH3.CO.CH2.CO.NPhH (cf. ACETO-ACETIC ACID, reaction 18), [85°]. Sl. sol. water and NII, Aq, v. sol. NaOHAq and acids. Distilled with aniline it gives s-di-phenyl urea. Boiling potash produces aniline, acetone, and acetic acid. Fe, Cl, colours its aqueous solution Br produces CH3.CO.CHBr.CONPhH, violet. Cone. H₂SO, forme (Py. 3, 1)-oxy-quinoline. Nitroue acid forms methyl - quinoline. CH, CO.C(NOII).CO.NPhH, [100°] (Knorr, A. 236, 75).

45. Acctophenone eyanhydrin gives rise to CaH3.CMe(NIIPh)CN (Jacoby, B. 19, 1515).

46. An alcoholic solution of quinone produces C₆H₂(NHPh)₂O₂ and hydroquinone. The former dissolves in conc. H2SO4 forming a crimson solution.

47. Tetrachloro-quinone (chloranil) produces, similarly, C₀Cl₂(NIIPh)₂O₂ (Hofmann, Pr. 13, 4; Hesse, A. 114, 292; Knapp a. Schultz, A. 210, 164).

48. Aniline mixed with an anilide and PCl,

produces a phenyl-amidine: e.g. $3PhNH_2 + 3CH_3 \cdot CO.NHPh + 2PCl_3 =$ 8CH, C(NPh).NHPh + P.O. + 6HCl (Hofmann, Z. 1866, 161).

Vor. 1.

49. Aniline hydrochloride and acetonitrile at 170° produce phenyl-acetamidine: CH,CN+H,NPh=CH,C(NPh).NH,

50. With benzo-trichloride, PhCCl, (1 mol.), aniline (2 mols.) on warming acts violently forming the hydrochloride of di-phenyl-benz. amidine, C,H,C(NPh)(NPhH)2HCl: aniline in glacial acetic acid, ZnCl, and PhCCl, give, chiefly, the same body.

51. But aniline hydrochloride (40 pts.), nitrobenzene (45 pts.), benzo-trichloride (40 pts.), and iron filings at 180° form the chloride of di-

amido-tri-phenyl-earbinol (q. v.).

52. Aniline (2 mol.) boiled with chloro-acctic acid (1 mol.) and water produces plienyl-amidoacetic acid and phenyl-imido-di-acetic acid; the aniline salt of the latter, PhN(CH., CO., NH, Ph) erystallises in needles, [99°] (P. Meyer, B. 14, 1325).

53. An alcoholic solution of aniline, chloroacetie acid, and ammonium sulphocyanide at 100° deposits crystals of phenyl-thio-hydantoic acid NIL, C(NPh). S. CH₂, CO₂H, [148 -152°]. This acid is decomposed by boiling with dilute (20 p.c.) H2SO4 into plienyl-urea and thio-glycollic acid (Jaeger, J. pr. [2] 16, 17; Claesson, B. 14, 732; Liebermann, A. 207, 129).

54. Acetamidoxim hydrochloride is converted by heating with aniline into acctanilidoxim, CH₃.C(NOH).NHPh, [121°] (Nordmann, B. 17,

2753).

55. Zincethide forms zine anilide Zn(NHPh), which is decomposed by water into Zn(OII)2 and

aniline (Frankland, Pr. 8, 504).

Salts.—(Beamer a. Clarke, Am. 1, 151; B. 12, 1066; Hjortdahl, Z. K. 6, 471).— B'HCl. [192°] (Pinner, B. 14, 1083). Needles or plates, v. sol. water and alcohol: may be sublimed.— B'₂H₂PtCl₃: yellow needles. — B'₂H₃NnCl₃: monoclinic. — B'₄H₂CuCl₁. — B'HB': trimetric, c:b:c = 723:1: 818. — B'HCdBr₃: trimetric. — B'HI.-B'HBiI, (Kraut, A. 210, 323).-B'HF: pearly scales, sol. water and boiling alcohol.-B'HClO,: long white prisms, sol. alcohol and other, m. sol. water; explodes at 75° .- B'HClO, -B'HIO₃. S.G. ¹³ 1.48. Explodes at 125°-130°. -B'HCNHg(CN)... [88°]. White needles tables (Claus a. Merek, B. 16, 2737). White needles or B',H,FeCy,: small micaceous erystals, v. sol. water, insol. alcohol and other (Eisenberg, A. 205, 267). — B'_H,FeCy, — B'_9H,CoCy, — B'_2H,PtCy,: triclinie (Scholtz, M. 1, 904). — B'_4H_3PO_4: laminæ, v. sol. water, ether, and hot alcohol (Nicholson, A. 59, 213; Lewy, B. 19, 1717). — B'₃H₃PO₄. — B'HPO₃. — B'₄H₄P₂O₅. — B'₄H₄SO₄: ni. sol. water, sl. sol. alcohol, insol. ether. Does not form an alum with aluminium sniphate (Wee. C. N. 38, 1).—BIL,SO₄: large plates; converted by water into the neutral sulphate (Wollington a. Tollens, B. 18, 3313). sulphate (Wolfington a. Toliens, B. 18, 3343),

— B'₂H₂S₂O₆ (Malczewsky, J. R. 11, 364). —

B'₁(H₂SO₄)₃H₁ (Jörgensen, J. pr. [2] 14, 384). —

B'INO₅ — Chloro-acctate. [88°]. — Dicehloro-acctate. [122°]. — Tri-chloroacetate. [145°]. -Oxalate B'2H.C2O4: trielinic columns, v. sol. water, el. sol. alcohol, insol. ether. — Phenate, B'HOPh. [30°]. (181°) (D.). (195°) (D. a. S.). Crystals resembling naphthalene (from alcohol or light petroleum) (Dyson, C. J. 43, 466). Formed by boiling equivalent quantities of phenol and aniline

together Dale a. Schorlemmer, O. J. 48, 186).

— Phihalatie, [1467]: needles.—(3)-Naphtholatie, [82-47]: orystalline powder (from light
petroleum) (Dyson, C. J. 43, 469). Other salts
of aniline are described under the various acids.

Combinations. (Schiff, C. R. 56, 268, 1095; Yohl, Ar. Ph. [2] 148, 201; Leeds, J. 1882, 500). —B's(SiF₁): minute needles, insol. benzene or petroleum-spirit; may be sublimed; converted by water or alcohol into anilino silicofluorido (Jackson a. Comey, B. 18, 3195). — B',Ag.SO, 2aq: hair-like crystals (Mixter, Am. 1, 239). —B',ZnSO, —B',ZnCl... —B',ZnBr... —B',Znl...—C,H,NH.HgCl: amorphous pp. got by mixing hot alcoholio solutions of aniline and HgCl., (Forster, A. 175, 30).—B',HlgCl.; needles, got by mixing cold alcoholio solutions of aniline and HgCl.,—B',HgBr., [112°] (Klein, B. 13, 835). —B',Hg(NO₃); pp., converted by hot water into O₂H,NH.HgNO₃, 3aq and (PhN)₂Hg.(NO₃), 2aq.—B',CaCl.,—B',CaCl

Acetyl derivative C.H.NO i.e. C.H.NH.C.H.O. Acetanilide. Mol. w. 135. [1142]. (295). S. 34 at 14°; 6·59 at 102°; Grievell evigate con. V.D. 48 (calc. 4.7).

Friswell, private com.). V.D. 4'8 (calo. 4'7).

Formation. — 1. From aniline and AcCl
(Gerhardt, A. 87, 164).—2. By beiling aniline
with glacial acctic acid (Greville Williams, C. J.
17, 106; v. also Chemical Chanop).—3. By heating
aniline with acctamide (Kelbe, B. 16, 1199).—4.
From acctophenone-oxim and H₂SO₄ at 100°
(Beckmann, B. 20, 1507).

Properties.—Laminæ (from water); v. sol. alcohol, ether, and benzenc. A saturated aqueous solution boils at 102.5° (Friswell).

Reactions.— 1. Passed through a red-hot tube it forms di-phenyl-urea, aniline, benzene, and CNH-(Nietzki, B. 10, 476).—2. ZnCl., at 260° gives flavanilino C₁₆H₁₄N₂.—3. PCl₅ forms CH₂,CCl.NHPlı which readily splits up into HCl and CH₂,CCl.NPl. The latter is converted by water into HCl and acetaniliae, and by anilino into di-phenyl-acetamidine, CH₂-C(NHPh):NPl. CH₂-CCl:NPh changes a little above its melting point [50°] into the hydrochloride of CH₂-C(NPh).CH₂-CCl:NPh, [117°] which at 160° changes to the hydrochloride of amorphous C₁₆H₄,N₂ (Wallach, A. 184, 86; cf. Miehael, J. pr. [2] 35, 207).—4. P₂S₃ forms thio-acetanilide (Hofmann a. Simpson, B. 11, 33°), Jacobson, B. 19, 1071; v. Thioacetta Acid.—5. Heating with sulphur produces oxalyl-amido-phenyl mercaptan, C₆H₄<N₈>C.C<N₈-C₆H₄, and some ethenyl-amido-phonyl mercaptan (Hofmann, B. 13,

1926). — 6. Dry NaOEt at 170° gives ethylaniline AcNHPh+NaOEt = EthHPh+NaOAe.
7. Nitrous acid passed into a solution of acetanilide in glacial HOAo forms an unstable nitrosamina, PhNAo.NO, [41°] (O. Fischer, B. 9, 403). — 8. Nitric acid converts acetanilide dissolved in 4 pts. of H₂SO, chiefly into p-nitroacetanilide, some o-being formed. If the acetanilide is dissolved in 20 pts. H₂SO, a small quantity of the m-compound appears (Nölting a. Collin, B. 17, 261).

Sodium acetanilido C.H.,NAcNa (Runge, Z. [2] 6, 119). Formed by distilling off tha alcohol from an alcoholic solution of equivalent quantities of acotanilide and sodium ethylate (Seifert, B. 18, 1358). Crystalline powder Absorbs CO₂ in the cold, becoming sodium acetyl-phenyl-carbamate, C. H. NAc. CO. Na.

Mercury acetanilide (C.H.NAc). Hg. [215°]. Formed by melting acetanilide with

HgO (Oppenheim a. Pfaff, B. 7, 624).

Hydrochlorido (C₆H₈NIIAc)₂IICl crystallises in needles, decomposed by water. By
heating for half-an-hour at 250° it splits off
acetic acid and yields the hydrochloride of diphous' heating at 280° it gives flavaniline. At
a still higher temporature quinoline bases ara
formed in small quantity (Nölting a. Weingürtner, B. 18, 1340).

p. Chloro-acetanilide C₉H₃NClAc. Acetyl-phenyl-chloro-amide. [172"]. Prepared by adding a cone solution of bleaching powder to a cone aqueous solution of acetanilide containing excess of acetic acid, as long as a pp. is formed. Colourless needles. Scarcely soluble in water. Crystallises well from very dilute acetic acid. Heated to 172° it suddenly changes, with explosive violence, to the isomerle p-chloro-acetanilide. This change is also produced by cold cone. HCl with a violent reaction. It dissolves in warm absolute alcohol at first unaltered, but after a few moments a violent reaction sets in and tha above change takes place. Alkalis and amine bases replace the Cl by H. It converts acetic ether into chloro-aceto-acetic ether. Not attacked by boiling water (Bender, B. 19, 2272).

Diacetyl derivative C₁₀H₁₁NO₂ i.s. C_LH_{_N}NAe₂. [111°]. Diacetanilide. From phenyl thiocarbinide and HOAo at 140° (Hofmann, B. 3, 770): PhNCS+2HOAo=PhNAc₂+CO₂+H₂S. Plates. On distillation it gives acetanilide (Gumpert, J. pr. [2] 32, 293).

Benzoyl derivative C₄H₃NHBz. [159°], Formation.—1. From aniline and BzCl (Gerhardt, A. Ch. [3] 37, 327).—2. By the action of phenyl oyanate upon benzene in presence of AlCl₄; the reaction probably being: (a) PhN.CO+1ICl=PhNH.COCl.

(b) PhNH.COCI+C₄H₆=PhNH.COC₄H₄+HOl (Leuckart, B. 18, 873). 8. From benzophenone oxim and H.SO, at 100° (Beckmann, B. 20, 1507)

Properties.—Volatile plates; insol. water. Reactions.—1. PCl. forms PhNH.CCl.Ph and then PhN:CClPh, [40°] (Wallach, A. 184, 79).—2. Boiling with sulphur produces benzenylamido-phenyl mercaptan.

Oxim C₁₃H₁₂N₂O i.e. C₆H₅.C(NOH)NHC₆H₅ Bens-anilidoxim. [136°] Obtained by heating taiobenzanilide with hydroxylamine hydro-chloride and Na₂CO₂ in alcoholic solution for about an hour (Müller, B. 19, 1669). Slender needles. Sol. hot water, alcohol, ether, chloroform, and benzene, sl. sol. ligroin. Dissolves both in acids and alkalis. Salts. - B'HCl: sol. alcohol. — B'2H2Cl2PtCl4x. Ethyl ether C.H., C(NHPh): NOEt. [56°]; white pp. Bensoyl derivative C.H., C(NHPh) NOB2. [116°]; white glistening needles; sol. alcohol, ether, and benzene, insol. water and ligroin.

Di-benzoyl derivative NPhBz,. [136°]. From benzanilido and BzCl (Gerhardt a. Chiozza, C. R. 37, 90).

Preparation .-- Benzanilide (18 g.) ie heated

with B2Cl (14 g.) for 3 hours.

Properties.—Needles (from alcohol). With dilute HCl at 120° it gives aniline and benzoic

acid (Higgin, C. J. 41, 132).

C.H.NBz. Di-benzoyl derivative [160°] (isomeric with preceding?). From benzoic acid (13 g. and phenyl thiocarbimido (6 g.) at 220° for 6 hours (Losanitch, B. 6, 176; Higgin, C. J. 41, 133): PhN:CS + 2HOBz = PhNBz2+CO2+H.S.

Properties.-Plates (from alcohol). With dilute HCl at 120° it gives aniline and benzoic acid.

ANILINE BLACK C₃₀II₂₂N₅. Nigraniline. Formed by mixing aniline, a chlorate (of K, Na, Nigraniline. or Ba) and a metallio salt (of Cu, Fe, V, Mn, or Cc) (Lightfoot, C. N. 11, 65; Lauth, Bl. [2] 2, 416; (Higgin, Bl. [2] 7, 93; Guyard, Bl. [2] 25, 58; Rosonstiohl, Bl. [2] 25, 356; C. R. 81, 1257; Kruis, D. P. J. 222, 347; Goppelsröder, C. R. 82, 331, 1392; Coquillion, C. R. 81, 408; Glenk, B. B. P. J. 200 D. P. J. 218, 234).. The quantity of metallie salt may be small; ammonium vanadate added to a solution of aniline hydrochloride (6 pts.) and NaClOs or KClOs (4 pts.) in water (100 pts.) eonverts 100,000 times its weight of aniline into black. An aniline black is formed at the positive pole when a solution of an aniline salt is electrolysed. Aniline black is purified by washing with water, alcohol, ether, and benzene. It is the hydroohloride of a base, C30 H25 No 2 HC1; and can be reduced by tin and HCl, or by III and P, to p-phenyleno-diamino and p-diamido-diphenylamine NH₂.C₂H₄.NH.C₅H₄.NH₂ (Nietzki, B. 11, 1093). Electrolytic aniline black appears to be C₂₄H₂₉N₄2HCl (Goppelsröder).

ANILINE CARBOXYLIC ACID v. PHENYL-

CARBAMIO ACID

ANILINE DYES v. ROSANILINE, MAUVEÏNE, CHRYGANILINE &C. ANILOTIC ACID is Nitro-salioylio soid v.

NITRO-OXY-BENZOIO ACID. ANILOXALBENZAMIC ACID v. PHENYL-

OXAMIDO-BENZOIO AOID.

ANIL - PYRUVIC C, H, NO, i.e. ACID CH2.C(NPh).CO2H. [122°]. Formed by the action of aniline on pyruvio acid (Böttinger, B. 16, 1924; A. 188, 336). Cryetals, v. sol. water. In contact with water it condenses to aniluvitonic or (Py. 3, 1)-methyl-quinoline carboxylic acid (q. v.). On bromination it yields tribromanil-dibromopyruvic acid. Salt.—BaA'₂: v. o. sol.

ANIL-UVITONIC ACID v. (Py. 3)-METHYL QUINCLINE (Py. 1)-carboxylic soid.

ANISAMIDE. Amide of methyl-p-oxy-

ANISANILIDE. Anilide of methyl-p-oxy-BENZOIO ACID

ANIS-BENZ-ANIS-HYDROXYLAMINE HYDROXYLAMINE.

ANIS - BENZ - HYDROXAMIC ACID HYDROXYLAMINE.

DI - ANIS - BENZ - HYDROXYLAMINE v.

HYDROXYLAMINE. ANISE, OIL OF. The essential oil obtained by distilling the seeds of Pimpinella anisum or Illicium anisatum with water. It contains anethol (q.v.). If the oil is heated with dilute IINO, (S.G. 1.2) and the resulting heavy oil shaken with warm NaIISO_aAq, so-called anisoïc or thianisoïc acid C₁₈H₁₈SO₄ [below 100°] is formed (Limpricht a. Ritter, A. 97, 364; Staedeler a. Wächter, A. 116, 169). It is a very soluble crystalline mass. NH, A'aq.—NaA'aq.— MgA'₂5aq.—CaA'₂2aq.—BaA'₂3aq.—AgA'. ANISE CAMPHOR. A name for anothol (q. v.).

ANISHYDRAMIDE C24H21N2O3 i.e. Tri-melhyl-tri-oxy-hydro-(MeO.C, H, CH), N₂. benzamide. [c. 120°]. Formed by action of cono. NH, Aq upon p-methoxy-benzoic (anisic) aldehyde. Prisms, insol. water, eol. boiling alcehol, ether, or cone. HClAq (Cahours, A. Ch. [3] 14, 487; Bertagnini, A. 88, 128). Changes at 170° into Anisin, a basic isomeride which forms ealts: B'HCl aq.—B',H,PtCl.

ANIS-HYDROXAMIC ACID v. HYDROXYL-

AMINE.

ANISIC ACID v. p-Methyl-Oxy-Benzolo Aoid. ANISIC ALDEHYDE v. p-Methyl-Oxy-Benzoid ALDERYDE

ANISIC ALCOHOL v. p-Methyl-Oxy-BENZYL ALCOHOL.

ANISIDINE v. Methyl derivative of AMIDO.

ANISIL $C_{16}H_{14}O_{4}$ i.e. $C_{6}H_{4}(OMe) - CO - CO - C_{6}H_{4}(OMe)$. [133°]. Prepared by oxidation of anison with Fehling's solution (Bösler, B. 14, 327). Yellow needles. Sol. hot, sl. sol. cold, Distils undecomposed. alcohol, insol. water. On boiling with alcoholic KOH it gives anisilic acid.

ANISILIC ACID C₁₅H₁₆O₅ i.e. (MeO.C₆H₄)₂C(OH).CO₂H. [164°]. Prepared by boiling anisil with alcoholic KOH (Bösler, B. 14, 328). Slender white needles. Sol. alcohol, sl. eol. water. Dissolves in strong IL SO, with a violet colour. On oxidation with CrO, it gives dimethoxy-benzophenone.

ANISIN v. ANISHYDRAMIDE,
ANISON J., H., O. i.e.
MOO.C., H., CII (CII). CO.C., H., O.M. Di-methyl-trioxy-phenyl-benzyl-ketone. [113°]. Propared by boiling p-methoxy-benzoic (anisic) aldehyde with alcoholic KOH (yield 60 p.o.; Bösler, B. 14, 826), or KCy (Rossel, Z. [2] 5, 562). Needles; v. sl. sol. water, sl. sol. cold alcohol and other. Strong H2SO, dissolves it with red colour, rapidly changing to pale green, and on heating, to yellow and then purple. See also Hydro-Anisoïn.
ANISOIC ACID v. Anethol.

ANISOL v. Methyl-Phenol.

ANISOL-ISATIN v. Di-methyl-di-Oxy-Di-PHENYL-OXINDOLE.

ANISOL-PHTHALIC ACID v. Methyl-Oxx. BENZOPHENONE CARBOXYLIC ACID.

ANISO-DIUREIDE v. Methyl-Oxy-BENZENYL-

ANISURIC ACID C10H11NO4 i.e. MeO.C0H4.CO.NH.CH2.CO2H. Methyl-oxy-bensoyl-glycocoll.-From silver amido-acetate and methyl-p-oxy-benzoyl chlorido. Also secreted when methyl-p-oxy-benzoic (anisio) acid is taken into the stomach. Lamino. Split up by acids into glycocoll and anisic acid.

Salts: CaA', 3aq.-AgA' (Cahours, A. 103, 90; 109, 32; Graebe a. Schultzen, A. 142, 348).

ANISYL. This name is usually given to the radicle methyl-p-oxy-phenyl, CH,O.C,H, but sometimes also to methyl-p-oxy-benzoyl, CH₃O.C, H, CO, which could more appropriately be called anisoyl.

ANISYL BROMIDE v. p-BROMO-PHENOL, Methyl-ether and Methyl-oxy-BENZOYL BROMIDE. ANISYL CARBAMIDE v. Methyl-p-Oxx-

PHENYL-UREA.

ANISYL CHLORIDE v. p-Chloro-Phenol, Methyl-ether and Methyl-oxy-nenzoul culoride. ANISYL METHYL KETONE v. Methyl-Oxy-

ACETOPHENONE ANISYL-THIO-UREA v. Methyl-Oxy-PHENYL-

THIO-UREA.

ANISYL-UREA v. Methyl-Oxy-PHENYL-UREA. ANOL C, II, O i.c. IIO.C, II, CH:CII.CII, [92·5°]. (c. 250°). Formed by heating anethol with KOII. Shining lamine (from alcohol, ether, and chloroform) sol. KOHAq and reppd.

by acids (Ladenburg, B. 2, 371).

ANTHEMENE $C_{1_0}II_{1_0}$. [64°]. (440°).

S.G. 15 942. V.D. 127 (calc. 131). S. (alcohol) $^{10}3$ at 25°. Extracted from flowers of chamomile (Anthemis nobilis) by light petroleum (Naudin, Bl. [2] 4I, 483). Minute needles, insol. water, sol. ether, petroleum, CS2, chloroform, and hot alcohol.

ANTHEMOL C. III, O. (214°). Occursinoilof chamomile (Anthemis nobilis). Thick liquid with camphor-like smell (Koebig, A. 195, 104). Boiling dilute HNO, produces terephthalic and p-toluio scids. Acetul derivative C10H15OAc. (235°).

ANTHOCYANIN. Blue colouring-matter of flowers v. Pigments

ANTHOXANTHIN. Yellow colouring matter of flowers, v. Pigments.

ANTHRACENE C, H10 i.e.

Mol. w. 178. [213°]. (o. 360°). H.F.p. – 115,000 (v. Rechenberg). S.V. 195 8 (Ramsay). Alcol.olic solutions containing less than '03 p.c. exhibit absorption bands in the ultra-violet part of the association that they, C. J. 39, 162). S. (ether) 1-17 at 15°; S. (HOAc) '44 at 15°. S. (benzene) 1-7 at 15°. S. (alcohol) '08 at 16°; (dilute alcohol, S.G. '84) '46 at 15° (Versmann, J. 1874, 182). Peaki, J. 1876. 423; Becchi, B. 12, 1978).

Occurrence .- In coal tar (Dumas a. Laurent, A. Ch. [2] 50, 187). From crude anthracene

ANISONITRILE v. Nitrils of Methyl-p-Oxx- the following bodies may be extracted by acetic ether:—(a) Soluble in cold alcohol: carbazol, phenanthrene, fluorene, and a hydrocarbon [130°]. Carbazol is insoluble in CS2, the others dissolve, but oan be separated by their picrates. (b) Soluble in cold benzene: synanthrene and two hydrocarbons [104°] and [97°].—(c) Soluble in hot benzene: anthracene (insoluble in dilute alcohol) and pseudophonanthrene (soluble in dilute alcohol).-(d) Insoluble in benzene : carbazol (Zeidler, A. 191, 302).

Formation.—1. By passing through red-hot tubes a mixture of ethylene with bonzene, diphenyl, or chrysene; or a mixture of benzene and styrene (Berthelot, Bl. [2] 7, 223; 8, 231; 9, 295) or benzeno and petroloum (Letny, B. 10, 412; 11, 1210), or benzene and oil of turpentine (Schultz, B. 7, 113).—2. By passing benzyltolnene, Ph.CH₂C₄H₄CH₅, through a red-hot tube, or over heated litharge (Behr a. Dorp, B. 6, 754; A. 169, 216).—3. By action of AlCl, on a mixture of benzene and s-tetra-bromo-ethane (Anschütz a. Eltzbacher, B. 16, 623).-4. By action of sodium on o-bromo-benzyl bromide (Jackson a. White, Am. 2, 392; B. 12, 1965).— 5. By heating benzyl cthyl oxide, PhCH, O.Et, with P.O. (J. Schulze, J. pr. [2] 27, 518).—6. By action of CHCl, or CH₂Cl₂ on benzeno in presence of AlCl, (Friedol, Crafts, a. Vincent, Bl. [2] 40, 97; 41, 325) .- 7. By heating benzyl chloride with water at 200° and distilling the product (Limpricht, A. 139, 303; Zincke, B. 7, 278).-8. By heating a mixture of alizarin with zine dust at low red heat (Graebe a. Liebermann, A. Suppl. 7, 297).-9. By distilling o-tolyl phenyl ketone with zinc dust (Behr a. Dorp, B. 7, 17) .- 10. Together with toluene by the action of AlCl, on benzyl chloride (Perkin a. Hodgkinson, C. J. 37, 726).-11. By distilling benzyl-phenol with P2Os (Paterno a. Fileti, B. 6, 1202). - 12. By heating o-benzoyl-benzoic acid with zinc dust (Gresly, A. 231, 238).

Properties.-Four- or six-sided monoclinic white tables with blue fluorescence. Insol. water, sl. sol. alcohol, ether, benzene, glacial IIOAc, chloroform, CS,, and light petroleum. Changed in sunlight into para-anthracene (paraphotene) [244°], which is insoluble in most menstrua, but is reconverted into anthracene by fusion.

Estimation.—Anthracenc (1 g.) is dissolved in boiling glacial HOAc (45 c.c.), filtered if necessary, and a solution of CrO₂ (10 g.) in glacial HOAc (5 c.c.) diluted with water (5 o.o.) is slowly added; a slight excess of CrO, should bo left after long boiling. The liquid is diluted with water (150 c.c.) and the ppd. anthraquinone washed, dried at 100°, and weighed (Luck, B. 6, 1347; v. also Meister, Lucius, a. Brüning, Fr. 16, 61; Paul a. Cownley, C. N. 28, 175; Lucas, C. N. 30, 190; Nicol, C. J. 30, 553; J.T. Brown,

C. J. 31, 232; Versmann, C. J. 30, 347).

Reactions.—1. Cone. HI at 280° forms anthraceno hydrides, toluene, and various paraffins. 2. Chronic acid produces anthraquinone .-Nitric acid also produces anthraquinono, and not nitro-anthracene .- 4. Sulphuric acid forms sulphonates .- 5. COCl, forms the chloride of anthracene (A)-carboxylio acid. - 6. H.O. produces anthraquinone (Leeds, B. 14, 1382).

Combinations.—With pierio acid it forms

C, H, O, H, (NO.) OH [188°]; red needles.—With dinitrothic phene: O₁₄H₁₀C₄H₂(NO₂)₂S [162°] (Resenberg, B. 18, 1778).—With nitric soid: C₁₄H₁₀NO₃H [125°]. Formed by passing nitrons fumes into anthracene suspended in glacial HOAc at 20°. White needles or prisms, sol. alcohol or benzene, nnstable when damp; con-warted by alkalis into nitrose-anthrone C₁₄H₂NO₂, [146°] and nitroso-hydroanthrone O14H11NO2 (Liebermann a. Lindermann, B. 13, 1535).— With nitric peroxide: C_{1.}H₁₀N₂O₄ [194°]. Formed by passing nitrous fumes into anthracene dissolved in glacial acetic acid cooled below 15°. Needles or plates, sl. sol. alcohol and benzenc; unstable when damp; converted by alkalis into nitroso-anthrone.—Hydro-anthraceno-nitrito, C_eH₄:C₂H₂(ONO)₂:C₂H₄ (?) [125°] is isomerio with the last body. It is formed by the action of HNO, on an acetio acid solution of anthracene-di-hydride. Small crystals. Easily soluble in benzeno. On boiling with alkalis nitroso-oxanthranol dissolves whilst nitronitrosoanthrone remains insoluble. Oxidation with CrO, converts it into anthraquinone (Liebermann a. Landshoff, B. 14, 467).

Constitution.—The constitutional formula given above is founded chiefly upon Formation 3, 4, and 2, and upon the formula assigned to anthraquinone (q. v.). Substitution in one of the benzene nuclei may be indicated by B prefixed to a number exactly as in the naphthalenederivatives; substitution in the $C_{\nu}H_{\nu}$ group is indicated by the prefix A. But in this dictionary the notation employed for anthraceno derivatives is usually exactly the same as that used for other benzene compounds (v. Benzene).

Additional References.—Graebe a. Liebermann, Z. [2] 4, 279, 503, 724; 5, 502; 6, 257; Fritzsche, N. Petersb. Acad. Bull. 9, 385; 13, 531. V. also Methiyl-anthracene, Ethyl-anthracene, Butyl-anthracene, Amyl-anthracene, and their hydrides; also Chloro-, Bromo-, Nitho-, anthracene, Anthramine, Anthrol, Anthranol and Sulpho-anthracene carboxylic actio.

Iseanthracens $C_{14}H_{10}$. [134°]. Obtained by passing di-benzyl-toluene, $C_{24}H_{20}$ (from toluene, benzyl eliloride, and zinc dust) through a redhot tube (Weber a. Zincke, B. 7, 1156). Januine; more soluble than anthracene. Converted by CrO_3 into a quinone $C_{14}H_8O_2$.

Para-anthracene C₁₄H₁₀. [244°]. Deposited when solutions of anthracene are exposed to sunlight. V. sl. sol. alcohol, ether, and benzene. Changed into anthracene by fusion. Does not combine with picrio acid (Schmidt, J. 2r. [2] 9, 248).

ANTHRACENE BROMIDE C₁₄H₁₆Br₂. Crystals formed by adding bromine to a solution of anthracene in CS₂ at 0° (Perkin, J. N. 39, 145). Split up by heat into HBr and bromoanthracene.

ANTHRACENE CARBOXYLIC ACID

C₁₅H₁₆O₂ i.e. C₁₄H₉.CO₂H. [280° corr.]. Two anthracene carboxylio acids can be obtained by distilling dryerade potassium anthracene sulphonate with K₄FeCy₆, and saponifying the mixture of nitriles that results (Liebermann, B. 8, 246; 13, 48). They may be separated by barytawater which forms a soluble salt with the (β)-

acid, and an insoluble one with the (γ) -acid. Nevertheless the two soids may be identical.

(8)-Arithracene carboxylic acid. [o. 260°]. Yellow needles (from alcohol). Insol. water, sl. sol. benzene and ether. Gives anthracene when heated with soda-lime, and anthraquinone carboxylic acid when exidised by CrO. Salts with heavy metals are v. sol. water. The acid and its salts show blue fluorescence.

(y). Anthracens carboxylic acid. [o. 280°]. Formed also by reducing anthraquinone carboxylic acid with zinc dust and ammonia (Börnstein, B. 16, 2609). Small plates or needles. May be sublimed. Sol. acetic acid and hot alcohol, sl. sol. cold alcohol and chloroform. Its solutions have a blue fluorescence. Cro., gives anthraquinone carboxylic acid [285°].

Salts.—NaA': small spangles, sl. sol. water; fluorescent. All the salts of the heavy metals are tolerably insoluble.

Ethyl ether A'Et: [135°], white plates, with blue fluorescence.

Chloride C₁₄H₂.COCl: yellow crystals, soluble in alcohol, ether, benzene, and CS₂ with an intense green fluorescence.

Amide C₁₄H₂.CONH₂: [295°], slender yellow needles or plates, sl. sol. alcohol with a blue fluorescenco, insol. wator, benzene, CS₂ and

Anthracene (A.)-carbexylic acid

C₆H₄:C₂H(CO₂H):C₆H₄. [206°]. From anthracene and COCl₂ at 200°, and saponifying the chloride by Na₂CO₂Aq (Graebe a. Liebermann, B. 2, 678). Formed also by heating (A. 1)-chloro-anthracene (A. 2)-carboxylic acid with alcoholic potash (Behla, B. 20, 701). Silky needles (from alcohol); decomposed by heat into CO₂ and anthracene. V. sl. sol. water. CrO₂ gives anthraquinone. Cl or Br (1 mol.) gives (A. 1)-chloro- (or bromo-) anthracene (A. 2)- carboxylic acid. A larger quantity (2 mols.) of Cl or Br give (A.)- di-chloro- (or bromo-) anthracene. Conc. H₂SO₄ gives (B.)-sulpho-anthracene (A.)-carboxylic acid. Salt—AgA.

carboxylio acid. Salt.—AgA'.

Methyl ether MeA'. [111°]. Yellowish
prisms or tables.

Anthracene-di-m-carboxylic acid

$$C_0H_4$$
 C_0H_2
 C

by reduction of an ammoniacal solution of the corresponding anthraquinone-m-di carboxylic acid with zinc - dust. Crystallino powder. Scarcely sol. water, m. sol. other solvents (Elbs a. Günther, D. 20, 1365).

Anthracene-di-carbexylic acid

reduction of an ammoniacal solution of anthraquinone-di-carboxylic acid [340°] with zinc-dust. Yellow crystalline powder. Scarcely sol. water, sl. sol. other solvents (Elbs a. Eurich, B. 20, 1363).

ANTHRACENE CHLORIDE C₁,H₁₀Cl₂. From anthracene dissolved in CS₂ at 0° by passing in Cl (Perkin, C. J. 31, 209). Needles (from

benzene). Splits up into HOl and chloranthracene even in the cold.

ANTHRACENE HYDRIDES.

Anthracene di-hydride C₁₄H₁₂. [108°]. (318°). Formed by heating anthracene at 160° with HI and red P, or by treating a solution in alcohol (95 p.c.) with sodium-amalgam. Prepared by heating anthraquinone (30 pts.), HI (140 pts. of S.G. I.8), and red P (10 pts.) with inverted condenser for one hour on a sand bath (Liebermann, A. Suppl. 7, 265; 212, 5). Large monoclinic plates (from alcohol) or needles (by sublimation). Insol. water, v. sol. alcohol, ether, and benzene. Volatile with steam. Its solutions fluoresce blue.

Reactions .- I. Warm conc. II, SO, forms anthracene and SO2.-2. Br added to its solution in CS2 forms di-bromo-anthracene. - 3. Conc. nitric acid forms hydro-anthracene-nitrito (v. sup.) and dinitroanthrone.-4. CrO, gives

anthraquinone.

Anthracene hexa-hydride $C_{14}H_{18}$. [63°]. (290°). From oxy-anthraquinone (or anthracene diliydride), fuming HI and red P by boiling for 20 hours (Liebermann, A. 212, 25; Suppl. 7, 273). Plates (from alcohol). Volatile with steam; v. sol. alcohol, ether, and benzene. At a red heat it is split up into hydrogen and anthracene.

ANTHRACENE . HYDRIDE CARBOXYLIC

ACIDS.

Anthracene-di-hydride carboxylie acid C15H12O2 i.e. C14H11(CO2H). [203°]. Formed, together with the following acid, by reduction of anthracene carboxylic acid, [280° cor.], with sodium-amalgam (Börnstein, B. 16, 2612). Colourless plates. V. sol. ordinary solvents.

Anthracene-tetra-hydride carboxylic acid C,H, (CO,H). [165°]. Colourless trimetric tables. Anthracene-bexa-hydride carboxylic acid

C14II15(CO2H). [232°]. Formed by reduction of anthracene-carboxylic acid by heating it with HI (1.7) and P at 220° (B.). Slender needles.

ANTHRACENE-DI-HYDRIDE SULPHONIC ACID C, H1. HSO. Prepared by reduction of sodium anthraquinono sulplionato with HI (S.G. 1.8) and red P (Liebermann, B. 12, 189, A. 212, 44). Decomposed by fusion with KOH with formation of anthracene and anthracene hydrido. NaA'aq: long soluble needles.—BaA'... CaA'..

ANTHRACENE SULPHONIC ACID

O,H,SO,H.

Formation.-From anthraquinone sulphonic soid, HI (S.G. 1.7), and red P, by boiling for half

an hour (Liebermann, A. 212, 48).

Preparation.-From sodium anthraquinone sulphonate (500 g.), zino dust (750 g.) and ammonia (3 litres of S.G. 88), at 100° (A. 212, 57; B. 15, 852). On oxidation by HNO, it gives anthraquinone sulphonic acid.

Salts.-NaA'4aq, v. sl. sol. water.-BaA'2.-

The existence of (a)- and (β) -anthraceno sulphonic acids amongst the disulphonic acids obtained by sulphonation of anthracene (Linke, J. pr. [2] 11, 222) has been denied by Liebermann (B. 12, 592).

(a)-Anthracene-disulphonic acid [1 2] C,H3(SO,H):C,H2:C,H3SO,II [3 4].

Preparation .- 1 pt. of anthracene is gently heated on a water bath with 3 pts. of H.SO, for

an hour. After dilution with water, the filtered solution is neutralised with PbCO, and the lead salts converted into the sodium salts.

Since the sodium salt of the (a)-acid is much less soluble in water containing Na₂CO₃ than the sodium salt of the (β) -acid it can be readily separated from the latter (which is formed simultaneously) (Liebermann a. Bocck, B. II, 1613; I2, I82, I287).

Properties .- Minute needles. By fusion with KOH it gives (α) -dioxyanthracene (chrysazol), which is converted on oxidation into chrysazin v. Di-oxy-anthraquinone.

Salts. -Na A" 4aq. -K A" aq. - CaA" 5aq. -BaA" 4aq.

(B)-Anthracene-disnlphonic acid

 $\begin{bmatrix} 1 & 2 \\ 3 \end{bmatrix}$ $C_6H_3(SO_3H):C_2H_2:C_6H_3.SO_3H$ $\begin{bmatrix} 2 \\ 3 \end{bmatrix}$ 1

Preparation .- I pt. of anthracene is heated to 100°C, with 3 pts. of H.SO, till half has dissolved. It is separated from the (a)-acid, simultaneously formed, by conversion into the sodium salt. By fusion with KOH it gives a dioxyanthracene, which on oxidation is converted into anthrarufin v. DI-OXY-ANTHRAQUINONE.

Salts.—Na, A" 3aq; white leaflets, easily soluble with a blue fluorescence.—BaA" 4aq; white leaflets.—A"Pb: crystalline pp.—CaA" 3aq (Liebermann a. Bocck, B. 11, 1613; 12, 182,

1287).

Anthracene - di - sulphonic acid (Flav-). C14H8(SO3H)2. Prepared by reduction of sodium (a)-anthraquinone di-sulphonate with zinc-dust and NH₃ (Schüler, B. 15, 1807).

Salts. - A"Na : soluble crystals, its solutions have a blue-violet fluorescence.—A"Ba: white

crystalline powder.

ANTHRACHRYSONE v. tetra-Oxy-ANTHRA-

ANTHRACYL-AMINE v. ANTHRAMINE.

ANTHRAFLAVIC ACID v. DI-OXY-ANTHRAQUI-NONE.

ANTHRAGALLOL v. (1, 2, 3)-tri-Oxy-ANTHRA.

ANTHRAHYDROQUINONE v. OXANTHRANOL ANTHRAMINE C, II, N i.e.

C.H.: (C2H2): C.H.NH2. Anthracylamine. Amido.

anthracene. Anthrylamine. [237°].

Formation. - I. By heating amide-anthraquinone with HI and P .- 2. By heating anthrol with acctamide at 280° and saponifying ths acctyl derivative so produced .-- 3. By heating anthrol with 10 p.o. aqueous NII, at 250°; the yiold is nearly theoretical .- 4. By heating anthrol with alcohol and ammonia at 170°.

Properties.-Yellow plates (from alcohol). May bo sublimed; sl. sol. alcohol, the solution having a splendid green fluoroscenco. Is a weak base, dissolving with difficulty in boiling HOL. Forms a blue mass when fused with arsenic

acid. Is readily methylated.

Reactions.—I. Does not give the carbamine or mustard oil tests.—2. Boiling HOAc gives di-anthramine. -3. Chloroform and alcoholio potash give rise to di-anthryl-formamidine C₁,H₂,NH.CH:N.C₁,H₂,—4. Nitrous acid gives C₂₈H₂₁N₃O, [250°], a body which forms a red solu-

tion in CS2, and a blue solution in H2SO4.
Salts.—BHCl: white iridescent plates, al. sol, water; formed by adding HCl to an alcoholic solution of the base. Its solution does not fluoresce. -B'2H2SO4: v. al. sol. water.

Acetyl derivative O, H, NHAo. [240°]. Plates. Its alcoholic solution fluoresces blue. CrO, gives acetyl-amido-anthraquinone.

C14H9NH.CHO. Formyl derivative [242°]. Small yellowish-green crystals, sl. sol. alcohol, with green fluorescence.

References. - Roemer, B. 15, 223; Liebermann

a. Bollert, B. 15, 226, 852; 16, 1635; A. 212, 57.

Di-anthramine C₂₄H₁₈N i.e. (C₁₄H₁)₂NH. Di-anthracylamine. Prepared by boiling anthramine with acctic acid (Bollert, B. 16, 1636). Does not melt at 320°. Small glistening plates. Very sparingly soluble in all solvents.

ANTHRAMINE-DI-HYDRIDE C, H, , NH, Slender colourless needles. Very soluble in alcohol. Formed by reduction of anthramine with sodium-amalgam. - B'HCl: sparingly soluble white needles (Liebermann a. Bollert, B. 15, 853).

ANTHRANIL C, H, NO i.e.
$$C_0H_4 < {}^{CO}_{NH} > (?)$$
.

o-Amido-benzoic lactam. (o. 213°). V.D. 4:14 (obs.). Formed by reduction of o-nitrobenzaldehyde with tin and acetic acid (Fried-Iander a. Henriques, B. 15, 2105). Colourless oil; volatile with steam. Soluble in ordinary solvents, but sparingly in water. Very weak basic properties. Reduces salts of gold and silver to the metal. By alkalis it is converted into anthranilic acid of which it is the anhydride. Double compound C,H,NOHgCl.: [174]; slender needles, sol. alcohol and hot water, sl. sol. cold water.

(above 360° with decomposition). Formed by heating isatoïc acid with BzCl (E. v. Meyer, J. pr. [2] 33, 19). Long white needles; readily takes up H₂O forming benzoyl-authranilic acid (Friedlander a. Wleugel, B. 16, 2229).

ANTHRANIL v-CARBOXYLIC ACID v. ISATOIC ACID.

ANTHRANILIC ACID v. o-Amido-Benzoic

ANTHRANOL O, HIO i.e.

thraquinone (30 g.), HI (140 g. of S.G. 1.75), and red P (8 g.), by 15 minutes' digestion (Liebermann, A. 212, 6). Needles (from benzene). Its alcoholic solution shows blue fluorescence. Decomposed by heat, becoming greenish. Dissolves in aqueous KOH, forming a yellow liquid, whonce CO, pps. the anthranol. The alkaline solution is oxidised by air, some anthraquinono being formed. CrO, in glacial HOAc completely exidises anthranol to anthraquinono.

Acetyl derivative. [126°-131°]. White needles (from dilute alcohol).

ANTHRANOL DIHYDRÍDE

O₆H₄<\frac{CH_2}{CH(OH)}>O₆H₄. [76°].

Preparation .- 50 grms. of anthraquinone are mixed with 100 grms. of zino dust and heated over a water bath with 300 c.c. ammonia and 200 o.c. of water. The liquid turns at first bloodred from oxanthranol, but after three hours this is reduced, the liquid becoming yellow. The chloride by heating with benzene and zinc-dust liquid is filtered, the pp. dried at 15°, and in sealed tubes at 220° (Piccard, B. 7, 1785) or

extracted with boiling benzoline (40°-60°), from which the anthranol dihydride crystallises on cooling (H. R. v. Perger, J. pr. [2] 23, 189).

Properties .- Slender satiny needles, which dissolve in benzoline, forming a solution with bluish fluorescence. May be crystallised from boiling water, but by long-continued boiling with water or with alcohol it is converted into anthracene:

with algorithms to converted that an interact
$$C_sH_s < \frac{C_{H_s}}{C_{H_s}} > C_sH_s = C_sH_s < \frac{C_{H_s}}{C_{H_s}} C_sH_s + H_sO_s$$

DIANTHRANYL $C_{2s}H_{1s}$ i.e.

C.H. C.H.

Formed by heating anthrapinacone C₂₈H₂₂O₃ with acetyl chlorido (K. Schulze, B. 18, 3035).

AN1HRAPINACONE C,H, $C_{28}H_{22}O_2$ i.e. CH_2 $C(OH) \cdot C(OH)$ CH_2 CH,

[c. 182°]. Formed as a by-product of the reduction of authraquinone to dihydroanthranol by means of zine-dust and NH, Long slender white needles. Sol. hot benzene, toluene, or xyleno, sl. sol. alcohol, insol. petroleum-ether. On heating with acetyl chloride, 2H₂O is removed giving dianthranyl C., H₁₀ (Schulze, B. 18, 3034).
ANTHRAPURPURIN v. Tri-OXY-ANTHRAQUI-

ANTHRAQUINOL v. OXANTHRANOL.

Formation. -1. By heating alizarin-blue with zinc-dust .- 2. By heating a mixture of anthramine, nitrobenzenc, glycerin, and H₂SO₄ (Graebe, B. 17, 170; A. 201, 344).

Properties.-Tables, insol. water, sol. alcohol and ether; its solutions show intense blue fluorescence. Its salts are yellow and possess in solution an intense green fluorescence. B'HCl.-

B'_H_P(Cl_e.—B'H_.—B'H_.SO_. Combinations.—With pieric acid it forms C_{1.H1}N C_eH_.(NO_.)₃O11: slender yellow needles. With ethyl iodide: B'EtI; golden needles, v.

sol. hot, sl. sol. cold, water.

Quinone C₁-H₂NO₂. [185°]. Formed from the preceding by CrO₂. Yellow prisms or needles, insol. water, sol. in alcohol and other. Salts .-B'HCl: sulphur yellow needles, sl. sol. water, but slowly decomposed by it. - B'2H2PtCl8. Picric acid compound B'C.H.N.O.; golden needles.

C14H8O2 i.e. C6H4 CO C6H4. Mol. w. 208. [273°]. S. (alcohol) '05 at 18°; 2.25 at 78°. V.D. 7.33 (calc. 7.20) (Graebe, B. 5, 15).

Formation.—1. By exidation of anthracene (Laurent, A. Ch. [2] 60, 220; 72, 415; A. 84, 287; Anderson, C. J. 15, 44).—2. From phthalyl chloride by heating with benzene and zinc-dust

by treatment with AlCl, (Friedel a. Crafts, Bl. [2] 29, 49).—3. By dry distillation of calcium phthalate (Panactovits, B. 17, 312).—4. Together with benzophenone by distillation of calcium benzoate (Kekulé a. Franchimont, B. 5, 908).—5. By heating o-(but not p-) benzoylbenzoic acid with P.O. at 200° and extracting with benzene (yield 20 p.c.; Behr a. Van Dorp, B. 7, 578).—6. In small quantity, by distilling benzoic acid with P₂O₃ (K. a. F.).—7. From phenyl o-tolyl ketone, MnO₂, and H₂SO₂Aq (Behr a. Dorp, B. 6, 753; 7, 16).—8. By acting on obromo-benzyl bromide dissolved in ether with Na and oxidising the product (anthracene) with Cro. (Jackson a. White, Am. 2, 390) .- 9. By action of water on 'anthraquinone chloride O14H2Cl2O, obtained by passing ohlorine into phenyl o-tolyl ketone at 110° (Thörner a. Zincke, B. 10, 1479).

Preparation.-Anthracone is dissolved in glacial acctio acid; K2Cr2O, or CrO3 is added; the liquid is then heated to 100°, the acetic acid is distilled off and the anthraquinone ppd. by water. Large quantities are prepared by oxidising anthracene (100 kilos.) with K₂Cr₂O, (150 kilos.) sulphuric acid (200 kilos.) and water

(2.000 kilos.).

Properties.—Yellow needles (by sublimation). Insol. water, v. sl. sol. alcohol, sl. sol. benzene. Not attacked by alcoholio KOH at 200°; or by

cold Br. Reactions .- 1. Bromine at 100° forms dibrome-anthraquinene (q. v.) .- 2. HI and P form anthranol and anthracene dihydride. -3. Heated with zinc dust to dull redness it is reduced to anthracene .- 4. Zinc dust and aqueous NaOH giveoxanthranol, C,H;(O,H,O);:C,H,(q,v); when alkyl iodides are added alkyl oxanthranols are got. When stronger soda is used and the alkyl iodide is not added until the reduction is complete, alkyl-hydro-anthranols (v. Hydro-ANTHRANOL) are got (Liebermann, A. 212, 100).— 5. Zinc dust and aqueons NH, give dihydro-oxanthranol, C₆H₄:(C₂H₄O₂):C₆H₄.—6. PCl, di-luted with PCl₃ at 200° forms chlorinated anthraquinones (Graebe a. Liebermann, A. 160, 121) .- 7. Potash-fusion at 250° forms potassium benzoate.

Constitution.—The formation of anthraquinone from phthalyl chloride (q. v.) and beazene might be thought to indicate the formula

This formula is open to O ZUH. several objections :- (a) the group CaH.: O is unknown; (b) it represents a lactone which should be converted by KOH into an oxy acid: (c) anthraquinone reacts with hydroxylamine while phthalido and its derivatives do not (E. v. Meyer, J. pr. [2] 29, 139, 496; V. Meyer, B. 17, 818). There remains the formula

C.H. CO C.H., which agrees with that of

C.H. and must thereanthracene C.H

fore be adopted. Bromo-phthalio acid, benzene, and AlCl, give bromo-benzoyl-benzoio acid, CO,H.O,H.Br.CO.O,H., in which it is evident that the carbonyls are o to one another. Cone. H.SO. condenses this acid to bromo-anthraquinone,

whence potash-fusion forms an oxy-anthraqu none from which phthalic (not oxy-phthalic) acid can be obtained by nitrio acid. Hence the two carbonyls are o to one another in both benzene nuclei (Pechmann, B. 12, 2125).

Derivatives of anthraquinone are described as CHLORO-ANTHRAQUINONE, BROMO-ANTHRAQUINONE, OXY - ANTHRAQUINONE, OXY - METHYL - ANTHRAQUI-

NONE, METHYL-ANTHRAQUINONE.

ANTHRAQUINONE CARBOXYLIC ACID C₁₅H₈O₄ i.e. C₅H₄:(C₂O₂):C₅H₄CO₂H. [282°-284°]. Obtained by boiling methyl-anthracene (Weiler, B. 7, 1186; O. Fischer, B. 7, 1196; Liebormann, A. 183, 166; Japp a. Schultz, B. 10, 1051), methyl-anthraquinone (Hammer-schlag, B. 11, 82), or anthracene earboxylio acid [280°] (Liebermann a. v. Rath, B. 8, 248), with CrO, and HOAc, or the compound C19H14O (obtained by action of cone. H.SO, on amyloxanthranol) with CrO, and H.SO, (Liebermann, A. 212, 97).

Properties.—Compact yellow prisms (from alcohol); yellow needles (by sublimation); v. sl. sol. HOAc, benzene, and alcohol, v. sol. acetone. Decomposed by heat into CO2 and The sodium salt is insol. anthraquinone. NaOHAq.

Salts.—BaA'2(?Aq) needles, v. sol. hot

water. - CaA'2(?Aq).

The following derivatives are got from the acid obtained by oxidising methyl-anthraquinone (Liebermann a. Glock, B. 17, 888).

Ethyl ether A'Et: [147°], needles, easily

soluble in alcohol.

Chloride C₁,H₇O₂.COCl: [147°], needles, very stable towards water. Amide C₁₄H₇O₂.CO.NH₂: [above

ncedles, very stable compound.

Anilide C, H,O2.CO.NHPh: [260°], needles, very sparingly soluble in most solvents.

(γ)-Anthraquinene carboxylic acid. [285°]. From the corresponding anthraceno carboxylic acid (Liebermann a. Bischof, B. 13, 49). Yellow ncedles (from glacial HOAc). Its alkaline solutions do not fluoresce. Its barium salt is v. sl. sol. water. This acid may be identical with the preceding.

Anthraquinene-di-m-carboxylic acid

$$O_8H_4$$
 $\stackrel{(1)}{\underset{CO}{CO}}$ $C_6H_2(\stackrel{(2:1)}{CO_2H})_2$. [above 330°]. Formed

by oxidation of the corresponding m-di-methylanthracene. Yellow needles. Nearly insol. water, sl. sol. other solvents. Dissolves in aqueous NH, with a red colour; the NII, salt crystallises in easily soluble small red warts; its solution gives with AgNO, a reddish pp. of A"Ag. By zinc-dust and aqueous NH, the acid is reduced to anthracenc-m-di-carboxylic acid (Elbs a. Günther, B. 20, 1364).

Anthraquinene-di-carboxylic acid

$$C_{e}H_{*} \stackrel{(1)}{\underset{(6)}{\text{CO}}} C_{e}H_{z} \stackrel{(3.4)}{\underset{(6)}{\text{CO}_{2}}} H_{z}.$$
 [340°]. Formed by

oxidation of di-methyl-anthraquinone [183°] by HNO, (1·2) at 220°. Yellow needles. Scarcely sol. water, sl. sol. most other solvents. Dissolves in aqueous NH, with a red colour. On heating it loses H.O. giving the anhydride. By sincdust and aqueous NH₂ it is reduced to anthrasene-di-carboxylic acid [345°]. The solution of the NH₄ salt gives pps. with CaCl₂, Pb(OAc)₂, and AgNO₂.

Anhydrids $C_{e}H_{1}C_{2}O_{2}.C_{e}H_{2} < \stackrel{CO}{CO} > 0$:

[290°]; sublimes in small yellow needles (Elbs & Eurich, B. 20, 1362).

ANTHRAQUINONE-OXIM

$$C_eH_e < CO > C_eH_e$$

Formed by heating anthraquinone with hydroxylamine hydrochloride and alcohol at 180° (Goldschmidt, B. 16, 2179). Red crystalline powder. Sublimes without melting above 200°. Dissolves in H₂SO₄ with an intense yellow colour.

ANTHRAQUINONE SULPHONIC ACID C_{1,}H₂SO₃ i.e. C₃H₁;(CO)₂:C₄H₂SO₃H. Formed together with the disulphouic acid by heating anthraquinone (1 pt.) with H₂SO₄ (2½ pts.) at 260°. Also from diamido-anthraquinone sulphonic acid by diazo reaction; and from obenzoyl-benzoic acid and furning H₂SO₄ (Liebormann, A. 160, 131; Suppl. 7, 805; v. Perger, J. pr. [2] 19, 218).

Properties.-Yellow scales, v. sol. water and

alcohol, v. sl. sol. H₂SO₄ and ether.

Reactions .- 1. Fused with potash it forms alizarin, oxy-anthraquinono, and benzoic, oxy-benzoic and protocatechuio acids. - 2. HI and P form authracene sulphonic acid and its dihydride. -3. Anthracene sulphonic acid is also produced by sodium-amalgam, and by zinc-dust and ammonia .- 4. Ammonia at 190° forms amido-anthraquinone. - 5. Distillation of the sodium salt produces, besides small quantities of anthraquinone and oxyanthraquinone, chiefly a compound C₂₈H_AO₈ which melts far above 300°. This compound forms minute reddishyellow needles (from glacial HOAc), sl. sol. IIOAc, toluene, phenol, and aniline, v. sl. sol. alcohol. At a high temperature it may be sublimed. It is insol. alkalis, but forms a crimson solution in cono. H2SO4. Distilled with zine dust it yields anthracene. Fused with potash it result yields animatent. Fuscult with potent is gives alizarin. Its constitution is perhaps C_sH_s:(CO)₂:C_sH_sO:(CO)₂:C_sH_sO:(CO)₂:C_sH_sO:(CO)₂:C_sH_sO: This forms transport of the constitution of the cons parent plates, insol. aqueous alkalis, v. sl. sol. boiling benzene, sl. sol. acetic acid, m. sol. aniline. It may be sublimed. Alcoholic KOII forms a violet solution decolorised by sliaking with air. Distillation with zinc dust produces anthracene (A. G. Perkin a. W. H. Perkin, jun., B. 18, 1723; C. J. 47, 682).

Salts.—BaA'2aq; small leaflets, sl. sol. water.—CaA'22aq; sl. sol. water.—NaA'aq:

white leaflets, sl. sol. water.

Chlorids C₁,II,O₂,SO₂Cl. [193°]. Light yellow plates; sol. benzene and acetic acid, v. sl. sol. alcohol and ether. Converted by dimethyl-aniline into the sulphone

C₁₄H₁O₂SO₂C₄H₁NMe₂. [171°].

A m i d e C₁₄H₁O₂SO₂NH₂. [261°]. Long yellow nsedles; almost insoluble in alcohol, chloroform

and CS_2 .

Anilide C, H,O, SO, NHPh. [198°]. Long

prisms; sol. alcohol and acetic acid.

Additional References.—Liebermann, A. 212, 42; B. 12, 189, 1293, 1597; MoHoul, B. 13, 692.

C₁₄H₄O₄(SO₂H)₂. When anthraquinone (1 pt.) is heated with fuming H₂SO₄ (2½ pts.) at 170°, or when dichloro-anthracene or di-bromo-anthracene is similarly treated, a mixture of (a) and (b) disulphonic acids is get. The salts of the (a)

(a)-Anthraquinons disulphonic acid

(a) dissiphonic actus is got. The saits of the (a) actid are less soluble and less crystalline than those of the (β) acid. The (α) acid is converted by potash-fusion into anthraflavin (di-oxy-anthraquinone), oxy-anthraquinone sulphonic acid, and flavonurourin (tri-oxy-anthraquinone)

and flavopurpurin (tri-oxy-anthraquinone).
Salts.—Na,A" 7aq.—BaA" aq.—PbA" aq.
(\$\beta\$).—Anthraquinone di-sulphenic acid. Prepared as above. Potash-fusion produces iso-anthraflavin (di-oxy-anthraquinone) and iso-purpurin (tri-oxy-anthraquinone). The sodium salt heated with NH,Aq at 180° produces (C. H. (MIN)MI 1/80 MIN) produces (C. H. (MIN)MIN) produces (C. H. (MIN)MIN)MIN) produces (C. H. (MIN)MIN)MIN produces (C. H. (MIN)MIN)MIN)MIN produces (C. H. (MIN)MIN)MIN produces (MIN)MIN produces (C. H. (MIN)MIN)MIN produces (C. H. (MIN)MIN)MIN produces (MIN)MIN p

i.e. C._iH₃(OH)(NH₂)(SO₃H)(Bourcart, Bl. [2] 33, 264).
Salts.—Na,A" 4aq.—Ba A" 2aq.—Pb A" aq.
Anthraquinone-disnlphonic acid C₁,H₀o₅y
i.e. C._oH₃(SO₃H):(CO)₂:C₃H₃.SO₃II. Prepared by oxidation of (β) anthracene-di-sulphonic acid. On fusion with KOH it first gives chrysazin (di-oxy-anthraquinone) and then oxychrysazin (trioxy-anthraquinone).

Salts.-Na A" 4aq : yellow prisms.

(ρ)-Anthraquinone-disulphenic acid

C₁₄H_aO_aS₂ i.e. [3 ½] C₆H₄(SO₃II)(CO)₂C₆H₃SO₂H

[½ s]. Prepared by oxidation of (a)-anthracone-disulphonic acid. On fusion with KOH it first gives anthrarufin (di-oxy-anthraqninone) and then oxy-chryszzin. Salts.—Na₂A" 5aq: yellow leaflets, sol. water.

References.—Graebe a. Liebermann, A. 160, 134; B. 3, 636; 7, 805; Liebermann a. Dehnst, B. 12, 1288; Perkin, C. N. 22, 37; A. 158, 323; Schunck a. Roemer, B. 9, 379; 10, 1821.

ANTHRAQUINONE CHLORIDE v. ANTHRA-

QUINONE, Formation 9.

ANTHRARUFIN v. Di-OXY-ANTHRAQUINONE. ANTHROL C₁₄H₁₀O i.e. C₈H₄(C.H₂)C₆H₃OH.

Formation.—1. From oxy-anthraquinone and III.—2. By fusing anthracene sulphonic acid with potash.

Preparation.—Crude sodium anthraquinone sulphonate (1 pt.) is heated on a water bath for a few hours with 1½ pts. of zinc-dust and 7 pts. of cone. ammonia; the anthraceno sulphonate thus obtained is fused with NaOH. Leaflets or needles (from dilute alcohol). Insol. water, v. sol. alcohol, acetone, or ether. Decomposes at 200°. Insol. in NH₂Aq, soluble in KOHAq or barytawater, forming a yellow solution with green flucrescence. Its alcoholic solution shows a violet fluorescence. Cone. IL-SO, gives a yellow solution, turned blue by heat. A drop of fuming HNO₃ added to its solution in glacial acetic acid gives a transient green colour. Its alcoholic solution reduces w.rm AgNO₃.

Acetyl derivative [1989]. Microscopic leatlets. Difficultly soluble in cold acetic acid, easily in C₂H₂. CrO₃ in HOAc converts it into the acetyl derivative of oxy-anthraquinone.

Ethyl derivative [146°]. Needles. Methyl derivative [c. 178°]. References.—Lichermann a. Hörmann, B. 12, 589; L. a. Hagen, B. 15, 1427; L. a. Bollert, B.

226; L., A. 212, 26, 49.
 Isomerides have been described by Linke, J. pr. [2] 11, 227.

ANTHROL-SULPHONIC ACID

C1.4H4(OH)(SO3H). Formed by careful fuelon of anthracene-di-sulphonic acid with KOH.

Salts .- A'Na: crystals sl. sol. cold water. A',Ba: plates or needlos (Schüler, B. 15, 1808). ANTHROPIC ACID. Shewn by Heintz (P.

84, 238; 87, 233) to be a mixture of palmitic and etearic acids.

ANTHROPOCHOLIC ACID C18H29O, 2aq. [145°]. [a]_D 50°. The cholic acid of human bile. The bile is extracted with alcohol; evaporated; extracted with dry alcohol, and the pp. (probably a mixture of sodium glyco-, and tauroanthropocholates) is decomposed by boiling with baryta-water (Bayer, H. 3, 293).

Properties.—Groups of prisms, insol. water, v. e. sol. alcohol, v. sol. ether, m. sol. chloreform. Lævorotatory. Fusion changes it into an amorphous dyslysin C181I26O2.

Salts.-KA': v. c. sol. water.-BaA'2 (? Aq):

silky plates, sl. sol. water. ANTHROXANIC ACID

$$C_sH_sNO_s$$
 i.e. C_oH_s $\nearrow O$ (?)

[191°]. Obtained by oxidation of anthroxanic aldehyde with dilute KMnO, (Schillinger a. Wleügel, B. 16, 2224). White felted needles. Sol. acetone and hot water, v. sl. sol. cold water, sl. sel. alcohol, ether, and benzene. Strong acid. On reduction with FeSO, and NH, it yields isatic acid.

ANTHROXANIC ALDEHYDE

[73°]. Obtained by heating a solution of o-nitro-(B)-oxy-cinnamic acid in an equal weight of acetic acid to 100° for a few hours, diluting the product with water, neutralising with CaCO,, distilling with steam, and extracting the distillate with ether. Long yellowish needles. Sublimable. Volatile with steam. Easily seluble in het water and in most other solvents except ligroine.

The addition of zinc-dust to the dilute ammoniacal solution produces a reddish-violet colonration. It combines with bisulphites and reddens fuelisin-sulphurous acid. With aniling it yields a crystalline anilide which melts at about 40° (Schillinger a. Wleiigel, B. 16, 2222).

DI-ANTHRYL-FORMAMIDINE v. ANTHRA-

MINE, reaction 3.

ANTIARIN C₁₄H₂₀O₅ 2aq (?). [221°]. Poisonous substance in the milky juice of Antiaris toxicaria or Upas Antjar, used to poison arrows (Mnlder, A. 28, 304; Ludwig a. de Vry, Z. 1869, 351; Pelletier a. Caventou, A. Ch. 26, 57).

ÁNTIMONIDES. — Binary compounds, or rather alloys, of Sb with more positive metals. Most of these bodies are of somewhat vague compositions; some occur as minerale (v. Anti-

MONY, Combinations, No. 10).

ANTIMONY. Sb. (Antimonium, or Stibium metallicum; Regulus antimonii). At. w. 120. Mol. w. probably 120, v. Biltz a. Meyer, B. 22, 725. [about 425°] (between 1090° and 1450') (Carnelley a. Williams, C. J. 35, 566). S.G. 6-71 to 6.86; c.15°, 6.697 (Schröder, J. 1859. 12) 8.G. (melted) 6.53 to 6.65 (Playfair a. Joule, C. S. Mem. 8, 57). S.H. (0°-100°) '0495 (Bunsen P. 141, 1); '0528 (Kopp, A. Suppl. 3, 66); (0°-33°) '049; (-21° to 0°) '048; (-75° to -21°) '047 (Pebal a. Jahn, W. 27, 584). C.E. (cub. 0°-100°) '003161 (Matthiessen, P. 180, 50); (lin. 40°) '00001152 (Fizeau, C. R. 68, 1125). T.C. (Ag=100) 4'03 (Lorenz, W. 13, 422). E.C. (Hg et 0°-1) 2'05 to 0° 140° th 10° (T. E.C. (Hg at 0°=1) 2.05 at 0°, 1.42 at 100° (Lorenz, Lc.). Chief lines in spectrum, 6128.5, 6078, 6003.5 (Thalen, A. Ch. [4] 18, 243). Crystallises in rhombohedra, approaching cubes (Marx, S. 59, 211); isomorphous with As, Bi, and Te.

Occurrence. Nativo; but chiefly as sulphide Sb.S. (Stibnite), and as double sulphide with PbS, CuS, Ag, S, NiS, &c.; as oxide in small quantities; in various iron ores; in ferruginous mineral waters; in some gas coals; in certain river sands (Campbell, P.M. [4] 20, 301; 21, 318).

Preparation. The sulphide is fused, to separate gangue, and roasted in air; the oxide thus produced is reduced by heating with charcoal or coal. Or the sulphide is reduced by charcoal or by iron. The crude metal (16 parts) is purified by fusion with dry Na CO₂ (2 parts) and Sb₂S₂ (1 part), for an hour in a Hessian crucible; the regulus is separated and again fused for an hour with 12 parts Na₂CO₃, and this is repeated with 1 part Na₂CO₃ a second time (Bensch, A. 5, 20). Or the crude metal is fused with NaNO, and Na₂CO₃ (details, v. Meyer, A. 66, 238). Pure Sb is prepared by Dexter (J. pr. 18, 449) by fusing dry H, SbO, with lampblack, and then with a little Sb₂O₃. Capitaine (P. 100, 563) prepares the pure metal by heating tartar-emetic in a closed crucible. Bongartz (B. 16, 1942) digests pure SbCl₂ with (NH₄)₂SAq in Pt vessels, electrolyses, fuses with pure Na₂CO₃, treats with dilute IIClAq, cleans with sea sand, and dries. Cooke (P. Am. A. [2] 5, 1) reduces NaSbO, by KCN, and fuses the Sb nuder Sb2O3 for several hours. Pure crystalline Sb may be obtained, according to Pfeifer (A. 209, 161), by electrolysing a solution of SbCl, in HClAq containing not more than 7 p.e. SbCl, Metallic antimony seems to have been known since the end of the 15th century. The sulphide was known to the ancients as Stibium.

Properties. - Brittle, hard, silver white, metailike; easily crystallised, isomorphous with As and Te, melts easily [425°]; volatilises at bright red heat in open vessel with simultaneous production of Sb₄O₆; scarcely volatilises in absence of air; but slightly volatile in vacuo at 292° (Demarcay, C. R. 95, 183); may be distilled in H at white heat. Unchanged in air at ordinary temperatures; melted on charcoal before blowpipe and then expessed to stream of air, pure Sb burns easily to Sb.O.; if traces of lead or iron are present a yellow or reddish sublimate is produced on burning before blowpipe. By electrolysis of SbCl, in HClAq, under special conditions, a lustrous silver-like deposit is obtained on the negative electrode; this deposit when dry explodee when rubbed with a hard substance, or when heated to 200°, with formation of olouds of SbCl3; a similar change occurs when the deposit is heated under water to 75°, but at ordinary temperatures it may be rubbed with a hard body under water without change. This so called explosive antimony contains SbOl. varying from 8 to 20 p.c. A similar explosive sabstance is obtained by electrolysing ShBr, in HBrAq, or ShI, in HIAq; the former contains 18 to 20 p.c. ShBr,, it explodes at 160°; the latter contains 22 p.c. HI and also SbI₃, it explodes at 160° (Gore, Pr. 9, 70 and 304; *ibid. C. J.* [2] 1, 365; Böttger, J. pr. 73, 484; 107, 43). According to Böttger (C. C. 1875. 674) explosive antimony also contains

occluded hydrogen.

The atomic weight of Sb has been determined (i.) by analysing and determining V.D. of oertain gascous compounds, particularly SbCl₂ and Sb(CH₃)₃; (ii.) by measuring the S.H. of Sb; (iii.) by comparing isomorphous compounds of Sb, As, and Bi; (iv.) by analyses of Sb, S, (Schneider, P. 98, 293), SbCl, (Weber, P. 98, 455; Dexter, P. 100, 563; Dumas, C. R. 46, 200, Dealer, F. 100, 505; Burnes, C. R. 40, 951; Kessler, P. 95, 204, 113, 134), SbBr₃ and SbI₃, and by synthesis of Sb.S. (Cooke, P. Am. A. [2] 5, 1; 7, 251; 9, 1; B. 13, 951) (comp. also Kessler, B. 12, 1044; Schneider, J. pr. [2] 22, 131; and Bongartz, B. 16, 1942). Some of the carlier determinations gave the number 122; but the researches of Cooke have established the value 120. The atom of Sb is trivalent in gaseous molecules, SbCl3, Sb(CH3)3. Antimony combines with oxygen and chlorine with production of heat: $[Sb^2, O^3, 311^{\circ}O] = 167,420$, [Sb², O', 3H²O] = 228,780, [Sb, Cl³] = 91,390, [Sb, Cl³] = 104,870 (Th. 2, 240). Antimony is oxidised by strong HNO, or by licating with various metallic oxides, e.g. MnO₂, PbO₂; hot conc. H SO, forms Sb₂.3SO₄; Sb combines with Cl or Br with production of light. Pure Sb is unacted on by HClAq out of contact with air; in presence of a little HNO3Aq solution proceeds rapidly (Cooke, P. Am. A. [2] 5, 1). Antimony forms three oxides Sb₄O₆, Sb₂O₄, and Sb₂O₅; various compounds corresponding to the first and third are known. Antimony is more metallic than arsenic, whether considered physically or chemically. Hydrated antimonious oxide, Sb₂O₃.3H₂O(=Sb(OH)₃), is known and reacts as a feeble base; if one third part of the H is replaced by K the remaining OII groups may be replaced by the residue of tartaric acid, with formation of Sb.OK.C₄H₁O₄ (v. Antimonious onide, Reactions, No. 4). Various compounds of Sb.O₅ with SO₅ (v. Antimonious onide, Reactions, No. 3), and at least one with N₂O₅ are known. A few unstable salts derived from the hypothetical hydrate Sbo.OII (=Sb₂O₃,H₂O) are known, so that Sb₂O₄ acts both towards strong acids and strong alkalis as a feeble saltforming oxide: thus [2HClAq, 2Sb-O³Aq] = 4,720; whereas [2HClAq, 2Na²OAq] = 25,500 (Th. 2, 241). The thio-antimonites are few in number and their stability is decidedly less than that of the thio-arsenites. The compounds of Sb which most decidedly exhibit negative or acidic functions are Sb₂O₄ and Sb₂S₅; the same holds good for As. The haloid compounds of Sb form many well-marked double salts. Many oxyhaloid compounds are also known. SbH, does not combine with acids, but compounds of the type SbR,X where $R = C_n H_{2n+1}$, and X is a halogon or even OH, have been prepared (v. ANTIMONY, COMPOUNDS WITH OROANIO RADICLES). For further discussion of the chemical relations of Sb, v. BISMUTH, CERNICAL RELATIONS OF; and NITROOEN OROUP OF REAL PRINTS.

Reactions .- 1. Water is not decomposed at ordinary temperatures by Sb; but at a red heat it reacts with steam to produce oxids of Sb and H .- 2. Dilute nitric acid digosted with finely powdered Sb forms a compound of Sb₂O₆ with N₂O₅; stronger acid forms chiefly Sb₂O₆.xH₂O and Sb.Q . - 3. Sulphuric acid ronets with Sbonly when conc. and hot; SO, is evolved, and a compound, or compounds, of Sb.O3 with SO3 produced .- 4. By the action of aqueous sulphurous acid at 200° in a closed tube Sb.S, is produced. 5. Hydrochloric acid forms SbCl, when heated with powdered Sh; in absence of air no action occurs (Cooke, P. Am. A. [2] 5, 1).-6. Aqua regia dissolves Sb forming SbCl₃.-7. Solid phosphoric acid and carbon heated with Sb form phosphide of Sb.-8. Alkali nitrates and chlorates heated with Sb form alkali antimonates and generally also Sb.O₃. -9. Alkali sulphates form Sb.S3, alkali sulphide, alkali antimonate,

and Sb.O.

Combinations. - 1. With nascent hydrogen SbII, is formed (q. v.). — 2. With chlorine, bromine, or iodine, the compounds SbCl, and SbCl., SbBr3, and Sb1, are produced (q. v.).-3. With fluorine (action of HFAq on Sb.O. and Sb.O. 3HO) SbF, and SbF, are formed (q. v.) .- 4. Sb combines with oxygen to form $\mathrm{Sb_{i}O_{6}}$ and $\mathrm{Sb_{i}O_{1}}$ $(q.\ v.)$; $\mathrm{Sb_{i}O_{5}}$ $(q.\ v.)$ is produced by the action of conc. $\mathrm{HNO_{5}}$ -5. The sulphide Sb.S. (q. v.) may be obtained by hoating Sh with sulphur; the pentasulphide Sb₂S₃ (q. v.) is best produced by decomposing Na₂Sb₃. by an acid.-6. Heated with selenion Sb. Se. is formed as a greyish metal-like solid (Berzelius; also Hofacker, A. 107, 6; v. also Uelsmann, A. 116, 124). Sb, Se, fused with Se and an alkali forms alkali seleno-antimonate (e.g. Na. SbSo.); this compound is decomposed by acids, in absence of air, with ppn. of brown Sb_Se_s. (Hofacker, l.c.). — 7. With tellurium (Oppenheim, J. pr. 71, 277) Sb forms either iron-grey SbTc, or tin-white Sb.Te, (S.G. of latter $\frac{130}{1} = 6.47 - 6.51$; Bödcker a. Giesecke). 8. Phosphorus is said to combine with Sb to form a tin-white brittle phosphide containing 15.5 p.o. P (Landgrebe, S. 53, 469). By the action of P (in CS₂) on SbBr₂ (in CS₂) a red powder, PSb, is obtained (Macivor, B. 6, 1362) .-9. Sb combines with arsenic, by fusion under boric acid, to form crystalline Sb. As (Descampes, C. R. 86, 1065). The compound Sb₂As, occurs native as allamontite. - 10. Antimony forms alloys with many metals; they are usually produced by melting together Sb and the specified metal. The alloys with K and Na are produced by fusing Sb with K₂CO₃ (or Na₂CO₃) and C, or by reducing Sb₄O₆ with KH.C₄H₄O₆ at high temperatures; they decompose water with evolution of H and separation of Sb; if containing much K or Na they take fire in the air. The alloys of Sb are usually lustrons, hard, and brittle. The alloys with Cu and Sn will be described under those metals. An alloy of iron is formed when Sb₂S₃ is reduced with excess of Fe; a mixture of 7 parts Sb and 3 parts Fe heated to whiteness in a charcoal-lined crucible produces a very hard white alloy. Gold loses its malleability by the presence of about 1 2006 of Sb. Lead alloys with Sb in all proportions; the lead is hardened; type metal is an alloy of about

17-20 parts 8b with lead and sometimes Bi or Sn (v. Lead, alloys or). With nickel two alloys are known; NiSb sublimes in prisms; Ni₂Sb occurs as breithauptite containing a little Fe and PbS. Two alloys with silver, Ag, Sb and Ag.Sb, occur native as antimonial silver. With sinc at least two crystalline alloys of definite composition are known, SbZn₂ and SbZn₂ (Cooke, Am. S. [2] 18, 229; 20, 222).

Many of these alloys are used in manufactures. Antimony compounds are also used in

medicine.

Detection .- Most Sb compounds are insoluble in water and in oxcess of conc. HNO, Aq, but many dissolve in HClAq, especially if tartario aoid is added; insoluble compounds may be dissolved by fusion with KNO, and K,CO, and treatment with IIClAq; when Sb compounds are fused with NaNO, and Na₂CO₂, NaSbO₃, insoluble in water, is formed.

In dry way. Heated on charcoal with Na₂CO₂ and KCN, all Sb compounds yield a brittle lustrous metallic bead. In the upper reduction-flame of the Bunsen lamp, Sb compounds give a green colour to the flame; in the oxidation flame, a white oxide film is obtained (on porcelain) which, moistened with nentral AgNO. Aq and then blown on with ammoniacal air, gives a black spot

(Ag.O).

In wet way. I. Antimonious compounds. (i.) Sulphuretted hydrogenpps, orangered Sb₂S₃ from acidulated solutions, soluble in KOHAq or NaOHAq, less soluble in NII,Aq, insoluble in NH₄HCO₆Aq, soluble in K₂SAq and (NH₄)₂SAq, insoluble in dilute acids, but dissolved by boiling with conc. HClAq. Dilute tartar emetic solution is not ppd. by II2S, the liquid turns red; conc. solutions are completely ppd. (Schulze, J. pr. [2] 27, 320). (ii.) Heated with gold chloride solution in presence of HClAq, Au is ppd. along with Sb₂O₃. (iii.) Caustic and carbonated alkalis pp. white Sb₄O₆ soluble in KOHAq and NaOHAq; the ppn. is slow and incomplete in presence of tartaric acid. (iv.) Zinc pps. Sb as a black powder; in presence of acids and Pt the Sb is deposited on the Pt, and a little SbH, is also formed; the deposited Sb is insoluble in cold HClAq, but easily dissolves in HNOsAq. (v.) Zinc and iron powder, added to a solution of an antimonious compound in conc. NH, ClAq cor.taining NH, Aq, ppt. Sb on the Zn without production of any SbHz; under similar conditions arssnious compounds yield AsII₃. (vi.) Zinc and dilute H_2SU_4Aq , in absence of HNO₃Aq, evolve H, mixed with SbH, which may be burnt in air with production of Sb₂O₃, or decomposed by heat into Sb and II, or led into AgNO, Aq whereby silver antimonide is ppd. mixed with Ag, or passed over S in snnshine whereby orange Sb₂S₃ is formed (v. Jones, C. J. [2] 14, 649; this is a very delicate test; v. Marsh's test for arsenio, under Arsenic, Detection of). (vii.) Dissolved in KOHAq, and treated with silver nitrate, a brown black pp. is obtained, from which NH, Aq removes Ag,O, leaving black Ag,O.

II. Antimonic compounds. - (i.) Sulphuretted hydrogen pps. orange-red SbyS, from acidulated solutions, soluble in KOIIAq, in K₂CO₃Aq, in (NH₄)₂SAq, and more slowly in (ii.) Heated with hydrochloric acid and potassium iodide, SbI, and I are formed $(Sb_2O_5 + 10KIAq = 2SbI_2Aq + 5K_2OAq + 4I)$. antimony trioxide has no action on KI, this reaction may be used to detect Sb2Oa in Sb2Os. (iii.) Gold salts, chromates, and permanganates, are not reduced by antimonic compounds; ner is AgNO, Aq acted on (compare tests (ii.) and (vii.) for antimonious compounds). (iv.) Towards zinc, or iron, and acids, antimonio compounds beliave similarly to antimonious (v. Antimonoso-ANTIMONIO OXIDE). Antimony tetroxide Sb2O4 gives the reactions both of Sb₂O₃ and Sb₂O₅; an alkaline solution reduces AgNO, Aq and AuCl, Aq

(For details of procedure in cases of suspected poisoning by Sb compounds a manual of toxicology must be consulted, e.g. Taylor On Poisons,

or Taylor's Medical Jurisprudence.)

Antimony may be separated (qualitatively) from tin and arsenio by treating the sulphides with conc. (NH4)HCO3Aq, which dissolves only As,S, then dissolving the SnS (or SnS,) and Sb,S, in conc. HClAq, boiling off II,S, ppg. Sb in one portion by Zn, reducing SnCl, to SnCl, in another portion by boiling with Cu turnings, and ppg. by HgCl2Aq; or the solution containing SbCl, and SnCl, may be diluted and boiled with a slight excess of iron wire whereby Sb is ppd. and SnCl, is reduced to SnCl, (Classen, J. pr. 93, 477). Sb.S. is completely converted into SbCl. by dry HCl at ordinary temperatures, whereas SnS is unacted on (Tookey, J. pr. 88, 435). A little As in Sb compounds may be detected by fusing with 2 pts. Na₂CO₃ and 4 pts. NaNO₃, and dissolving in water, Na AsO, goes into solution and NaShO, remains.

Estimation.-I. Gravimetrio methods. Antimonious compounds are ppd. by H2S in presence of HClAq and tartario acid, excess of H.S is removed by CO2, the pp. of Sb2S3 is collected (after boiling for 15-20 mins.; Sharples, Fr. 10, 343), on a weighed filter, dried at 100° and weighed; a portion is then dried by heating in a stream of dry CO, and again weighed; if a portion of the pp. dried at 100° yields S on treatment with hot cone. IIClAq, the pp. contains Sh₂S₅ or free S; in this case the other portion must be heated in dry CO, until S is no longer volatilised; the residue is now pure Sb2S2 (v. also Cooke, P. Am. A. 13, 1; 17, 13). Or the pp. of Sb₂S₃ (perhaps mixed with Sb₂S₅ and S) may be converted into Sb₂O₄ by treatment with cohc. HNO, Aq (for details v. Bunsen, A. 106, 3). Schneider (P.110, 634) decomposes the Sb₂S₂ by HClAq, leads the H2S into an alkaline solution and determines it by volumetric methods (iodine method, or ppn. by excess of titrated As,O,Aq

and determination of excess of As.O. by iodino).

H. Volumetrio methods. (i.) Antimonious compounds are oxidised to antimonic by iodine in alkaline solutions (Sb₂O₃ + 4I + 2II₂O = Sb₂O₃+4HI). NaHCO₂Aq is the best alkaline solution; titrated I solution is run in until a blue colour is produced with starch. (ii.) Antimonious compounds are oxidised to antimonic in presence of tartaric acid, by K2Mn2O, solution (attention must be paid to details, v. Guyard, Bl. 6, 89). For other methods of estimating Sb, especially in presonce of As, or of As and Sn v. Bunsen, A. 106, 3; 192, 317; Clarko, Am. S. [2] 49, 48.

References .- In addition to those in the text,

Bergmann, Opusc. 3, 164; Thénard, A. Ch. 32, 257; Proust, G. A. 25, 186; Berzelius, S. 6, 144; 22, 69; 34, 58; P. 20, 365; 37, 163; Berthier, A. Ch. [2] 22, 239; 25, 379; H. Rose, P. 8, 441; 42, 532; 24, 165; Vauquelin, S. 2I, 219. Antimony, acids of, and their salts (compare arts. Acins; Acids, Basicity of; Hynroxides). The oxides Sb₂O₄ and Sb₂O₅ are searcely soluble in water, but each reddens moist blue litmus paper; the oxido Sb4O6 is slightly soluble in water, but is without action on litmus. A few feebly marked salts are known which may be regarded as dcrived from the hypothetical hydrate Sb₂O₃.H₂O (=SbO.OII); two sodium salts, so-called antimonites, are obtained according to Terreil (A. Ch. [4] 7, 380) by dissolving Sb.O. in boiling NaOHAq and allowing to cool (v. infra). Two hydrates of Sb₄O₆, viz. Sb₄O₆, 4H₂O and Sb₄O₆, 6H₂O have been obtained (v. Antimonious oxide); but neither seems to possess acidic properties. The oxide Sb,Oc reacts with Na CO3 when the two are fused together, but on adding water Sb₄O₆ is ppd. and NaOH remains in solution. The acid-forming character of Sb.O. is therefore extremely feeble (v. further Antimonious oxide).

the following older memoirs are important:-

No hydrate of Sb₂O₄ is known; but by fusing this oxide with KOII or K₂CO₃, a compound, Sb₂O₄.K₂O, insoluble in cold water, is produced; by dissolving this in hot water and adding various metallic compounds, several compounds of Sb₂O₄ with metallic oxides, e.g. Sb₂O₄CaO and Sb₂O₄CuO (which both occur nativo as romette and ammiolite respectively [? merely mixtures]) are obtained. A solution of Sb₂O₄ in KOIIAq (obtained by fusion) is easily decomposed: e.g. on boiling and then diluting, Sb₂O₃ is ppd. and Sb₂O₃xH₂O is then thrown down on addition of acids; on standing in air, without boiling, KSbO₂ is produced. The oxide Sb₂O₄ cannot therefore be regarded as a definite anhydride, nor can any acid, or well-marked series of salts, be said to exist corresponding with this oxide (v. Antimonx, oxides or).

Of).

Three hydrates of antimonic oxide are known (v. infra): Sh.O.H.O (= HSbO.), Sb.O.2II.O (= H.Sb.O.), Bh.O..2II.O (= H.Sb.O.), Bh.O..3H.O (= II.SbO.). The first and third may be obtained from H.Sb.O.2II.O which is a product of the action of water on SbCl.; dried over H.SO., H.SbO. is obtained, and at higher temperatures HSbO. is produced (v. ANTIMONATES). Soveral fairly marked salts are known, antimonates, derived from HSbO.; two series of metantimonates (M.Sb.O. and M.H.Sb.O.) exist (v. infra); no salts of the hydrate Sb.O.3H.O (= H.SbO.) have been obtained. Antimonates are usually obtained by fusion; aqueous alkalis dissolve the hydrate Sb.O.H.O without change; some metantimonates are produced from the hydrate Sb.O.2.2H.O by the action of alkalis in the wet way. Antimonic oxide is evidently a feebly marked acid-forming oxide. The only definite compounds of Sb hitherto obtained which exhibit acidic.characters are then HSbO. and H.Sb.O.

The following thermal data are given by Thomsen [Sb², O³, 3H²O] = 167,420; [Sb, O², H, H²O] = 117,890; [Sb², O³, 3H²O] = 228,780; [Sb, O³, H, H²O] = 145,570; [SbO³H³, O] = 30,680.

I. Antimonites.—Two sodium salts are desoribed by Terreil (A. Oh. [4] 7, 380): NaSbO₂3H₂O, lustrous ootahedral orystals, obtained by dissolving Sb₂O₃ in boiling NaOHAq and allowing to cool; NaSbO₂Sb₂O₄H₂O. Harge crystals, insoluble in water, obtained from very concentrated alkaline solutions.

II. Antimonoso-antimonates.—This name has been given to the compounds of Sb₂O, with metallic oxides; it implies that these bodies are compounds of antimonites with antimonates; very little, however, is known of their properties. Two potassium salts K₂O.Sb₂O₄ (?KSbO₃,KSbO₂), and K₂O.2Sb₂O₄ are said to be produced, the former by fusing Sb₂O₄ with KOH or K.CO₃ and washing with cold water, the latter by the action of a little HClAq on the former. An aqueous solution (the salt dissolves in hot water) of K₂O.Sb₂O₄ is said to give pps. with various metallic salts. These salts might perhaps be regarded as derivatives of the hypothetical hydrates Sb₂O₄.H₂O (=II_Sb₂O₃) and 2Sb₂O₄.H₂O (=II_Sb₂O₃); but our knowledge of them is almost nil.

III. ANTIMONATES, AND ANTIMONIO ACIDS .--Three hydrates of Sb₂O₅ are known. By ppg. KSbO₃Aq by HNO₃Aq, washing the pp. and leaving it for a whole summer, Geuther obtained the hydrate Sh₂0,3H₂O (=H₃SbO₃) (J. pr. [2] 4, 438); at 175° this hydrate gives HSbO₃ (=Sh₂O₃H₂O). The hydrate Sh₂O₃2H₂O = II,Sb.O,) is obtained by adding hot water to SbCl, and drying the pp. of H,Sb,O,.2H,O at 100°; it is also produced by decomposing the salts M,SbO₂ by acids: this hydrate is easily decomposed to HSbO₃, even by standing in contact with water (Dubrawa, A. 186, I10; Conrad, C. N. 40, I97). HSbO₃ may also be obtained by decomposing MSbO₃ by acids, or by oxidising Sb by HNO₃. The hydrate HSbO₃ is dishting soluble in water installable in MIA. slightly soluble in water, insoluble in NH,Aq, and easily soluble in KOHAq; II,Sb,O, is more soluble in water, and dissolves in both NHAAq and KOHAq: little is known of the hydrate Sb₂O₅.3H₂O. The antimonates belong to the two types MSbO2, and M,Sb2O2; the former are usually called antimonates, the latter metantimonates.

Antimonates: investigated by Berzelius, then by Fremy (A. Ch. [3] 12, 499; 22, 404), and by Heffter (P. 86, 418; 98, 293). These salts are obtained by fusing Sb or Sb₂O, with nitrates, or HSbO, with carbonates, or by double decomposition from the K salts; ank-ous alkalisdissolve HSbO₂ without clange. Some of the K and NH, salts are soluble in water, the others are slightly soluble or insoluble. The normal antimonates are converted into acid salts by the action of weak acids (e.g. CO₂Aq), they are decomposed by stronger acids with separation of HSbO₂; the antimonates are decomposed by fusion with NH₂Cl, the whole of the Sb being volatilised as SbCl₂; those which are soluble in water or acids are decomposed by (NH₄)₂SAq, with production of this-antimonates.

Ammonium antimonate NH₄SbO₂2H₂O; white crystalline powder, insoluble in water, easily decomposed with loss of NH₃: obtained by dissolving HSbO₃ in warm NH₄A₄.

Barium antimonate Ba(SbO₃)₂; obtained by adding BaCl₂Aq to KSbO₃Aq; pp. at first in

flocculent but becomes crystalline. By adding BaCl,Aq to boiling NaSbO,Aq a floceulant pp. of Ba(SbO,), 5H,O (air-drisd) is obtained.

Potassium antimonates.—The normal salt KSbO₃ is obtained by fusing 1 part Sb with 4 parts KNO, and washing with hot water; it dissolves after long boiling with water, and is obtained as a mass of white crystals when the solution is evaporated until a crust forms. Another form of this salt is described by Fremy as a guminy mass, obtained by evaporating this foregoing solution nearly to dryness, or mors easily by long-continued fusion, either alone or with KOH or K.CO3, of the product obtained by melting together 1 part Sb and 4 parts KNO3. The gum-like salt dried in vacuo is 2KSbO.5H.O. it is easily soluble in hot water; dried at 160° it leaves 2KSbO, 3H,O which is changed to the gum-like salt by boiling with water; at a red heat all the water is removed, and the product is gradually changed to the gum-like salt by contact with hot water. When the normal salt is boiled with water, a residus of 2KSbO2.Sb2O2.10II2O is ohtained; and a similar salt with 6H2O is produced by the action of CO2 on the normal salt (Heffter, P. 86, 418; v. also v. Knorrs a. Olschewsky, B. 18, 2353).

Sodium antimonates. - The salt 2NaSbO, 7H2O is obtained similarly to ths

normal KSbO3; also in octahedra by the action of NaOll Art on Sh.S. NaShO, 3H.O is said to be formed by the action of NaOHAq on Sh.S., filtration, and addition of more NaOHAq (v. also v. Knorro a. Olschewsky, B. 18, 2353).

Many other antimonates are described by Fremy and Heffter; the chief are the salts Ca(SbO₃), Co(SbO₃), Cu(SbO₃), 5H,O, Pb(SbO₃), LiSbO₃, Ma,2SbO₄,12H,O, Hg(SbO₃), S(SbO₃), 26H,O, Sn(SbO₃), 22H,O (v. also Schiff, A. 12O, 47; Unger, Ar. Ph. [2] 147, 195).

Metantimonates. These salts fall into two classes-normal salts M.Sh.O., and acid salts M2H2Sb2O; they are formed from the antimonates by addition of metallic oxide or water $(2MSbO_3 + M_2O = M_4Sb_2O_3; \text{ and } 2MSbO_3 + H_2O =$ M.H.Sb.O.); conversely the metantimonates lose M2O (or H2O) and form MSbO2. The metantimonates as a class are insoluble in water, the alkali salts are crystalline; they are decomposed by acids; they have been chiefly investigated by Fremy (A. Ch. [3] 12, 499).

Ammonium metantimonates .- H.Sb.O. dissolves slowly in cold NH3Aq; a conc. solution on addition of alcohol gives the acid salt (NH₄)₂H₂Sb₂O₁.5H₂O; this salt is easily decomposed by heating, either in presence or absence of water, into (NH₄)SbO₂. The normal salt has

not yet been isolated.

Potassium metantimonates .- By fusing KSbO, (best the gum-like salt) with about 3 parts KOH, dissolving in water, and crystallising, the salt K,Sb,O, is obtained as deliquescent, easily soluble, crystals. The acid salt K, H, Sb, O, 6ILO is produced by decomposing the normal salt by a little water (KOHAq is also produced), or by dissolving SbCl, in excess of KOHAq, oxidising by K.Mn.O.Aq, and crystallising (Reynoso, A. Ch. [3] 23, 325); at 200° the dehydrated salt K₂H₂Sb₂O₇ is obtained, and at 300° KSbO₃ is formsd. The acid salt is slightly soluble in cold water, more easily in water at 40°-50°, with

gradual production of the gum-like KSbO.; at aqueous solution of this salt precipitates sodium salts. Other metantimonates are described by

Fremy (l.c.).

Seleno-antimonates .- A fsw salts are known, derived from the hypothetical sslenoantimonic acid H2SbSs4. Na2SbSe4.9H2O forms orange-red tetrahedral crystals, and is obtained by fusing Na₂CO₃, Sb₂Se₃, Ss, and C; the salt Na₂SbS₃Se₂9H₂O is obtained as yellow tetrahsdral crystals by boiling Na,SbS,Aq with Se (Hofacker, A. 107, 6).

Antimeny, alleys ef, v. Antimony, Combinations, No. 10.

Antimeny, arssnids of, v. Antimony, Combinations, No. 9.

Antimony, bromids of. SbBr., No other bromido is known. Mol. w. 359.28; [90°-94°] (Scrullas, P. 14, 112). (275°-280°) (Kopp, A. 94, 257; Cooks, P. Am. A. [2] 5, 72). V.D. 180 (Worcester, P. Am. A. [2] 10, 61). S.G. 120 4·148 (Cooks, l.c.); fused 80 3·641 (Kopp, l.c.). If the vol. of fused 80 Bbr₃=1 for 40 =0°, then the vol. at $t^{\circ} = 1 + 000576d + 0000013465d^{2}$, where $d = t^{\circ} - 90^{\circ}$ (Kopp, *l.c.*). H.F. solid Sb, gaseous Br,[Sb,Br3] = 76,900 (Guntz, C. R. 101, 161).

Formation.-1. By shaking powdered Sb into a retort containing Br and connected with a condenser.—2. By distilling a mixture of Sb sulphate and KBr (Macivor, C. N. 29, 179).

Preparation .- 1. By adding powdered Sb to a solution of Br in dry CS, at 0°, distilling off CS, adding powdered Sb, distilling off the SbBr. and recrystallising it from CS, (Cooke, P. Am. A.

[2] 5, 72; Nicklès, C. R. 48, 837).

Properties and Reactions .- Deliquescent trimetric crystals; a:b:c=1.224:1:1.064 (Cooks, l.c.); sublimes when heated. Decomposed by water; cold water produces Sb₄O₅Br₂, hot water 10Sb₄O₅Br₂,SbBr₃ (Macivor, C. N. 29, 179). The compound Sh.O. Br., is also produced by heating ShBr., with alcohol to 160° (Macivor, i.e.). The action of air and sunlight on SbBr, in CS, produces an oxybroinide, probably SbOBr (Cooke, P. Am. A. [2] 5, 72). Combines with KCl to form SbBrg. 3KCl, which according to Atkinson is identical with SbCl2.3KBr obtained by action of SbCl2 on KBr in presence of a little water (C. J. 43, 290).

Antimeny, chlerides ef. Sb and Cl combins directly to form two compounds SbCl, and SbCl,; the former may be gasified, the latter is decomposed by heat at ordinary pressures into

SbCl₃ + Cl₂ (v. infra).

I. Antimonious ohloride. SbCl₃. 226·11. [73°·2] (Thorpe, C. J. 37, 387). (223·5°) 216°, Cooke, P. Am. A. [2] 5, 72). S.G. 36 2-6753 (Thorpo, Lc.). S.G. 3-064 (Cooke, P. Am. A. [2] 5, 72). V.D. 115-6. V = $1 + 0008054d + 00001032d^2$, where d = degreesabove M.P. (73.2°) (Thorpe, l.c.). [Sb,Cl] =

301,390 (Thomsen).

Formation.—1. By dissolving Sb,Sb₂O₄, or Sb₂S₃, in HClAq with a littls HNO₂Aq, syaporating, and then distilling .- 2. By the action of of lon Sb₂S₃,—3. By distilling together 1 part powdered Sb with 2 parts HgCl₂; or 3 parts Sb₂S₃ with 7 parts HgCl₂; or 1 part Sb₂(SO₂), with 2 parts dry NaCl.—4. By distilling 2 parts of the control o Sb₂O₄ (impure), with 6 parts dry NaCl, 4 parts H₂SO₄, and 2 parts H₂O and changing the

receiver as soon as the distillate begins to solidify

Preparation.—1. By passing dry Cl into a retort containing powdered Sb, until most of the Sb is transformed into SbCl,; a little more Sb is then added, the stream of Cl is stopped, and the SbCl, is distilled off into a dry receiver. Cooke (P. Am. A. [2] 5, 72) saturates warm CS, with SbCl, and cools by freezing mixture. A solution of SbCl, in cone. HClAq (generally prepared by the action of the acid on Sb.S.) is used

in pharmacy.

Properties.—A colourless, translucent, crystalline, mass. Melted and allowed partially to solidify, or dissolved in hot CS, and cooled, trimetric crystals are obtained, a:b:c = 1 263:1:1 09 (Cooke, P. Am. A. [2] 5, 72). Very caustic. Solublo in alcohol without change; on heating this solution oxyohlorides of Sb (q.v.), HCl, and C2H3Cl, are formed. It absorbs moisture from the air and forms a clear liquid, from which crystals of SbCl, are obtained by

standing over H₂SO₄.

Reactions.—1. With water various oxychlorides are produced (SbCl, dissolves unchanged in a very little water at ordinary temperatures); if a little cold water is added (about 2 parts to 1 part SbCl3), and tho pp. is washed with ether, SbOCl (q. v.) is obtained (Peligot, A. 64, 280; Sabanajew, Bl. [2] 16, 79). When from 5 to 50 parts H2O are added to 1 part SbCl2, tho compound Sb₄O₅Cl₂ (q. v.) is obtained (Sabanajew, l.c.). Other observations point to a varying composition for the product of the mutual action of SbCl, and H2O; by continued washing the whole of the Cl may be removed (v. Duflos, S. 67, 268; Johnston, J. pr. 6, 55; Malaguti, J. pr. 6, 253; Peligot, A. 64, 280; Schneider, P. 108, 407; Schäffer, A. 152, 314). Thomsen (Th. 2, 240) gives these data; [SbCl*,Aq]=8,910 when Sb₄O₅Cl₂ is formed, and = 7,730 when Sb₂O₅Aq and 11ClAq are formed. According to Williams (C. N. 24, 225) boiling water produces 10Sb₄Cl₂O₅.SbCl₃. Formation of oxychlorides is prevented by tartario acid.—2. Antimonious oxide dissolves in boiling SbCl, to form oxychlorides; SbOCl.7SbCl, is described by Schneider (P. 108, 407).—3. Alcohol heated with SbCl, in proportion C.H.O.SbCl, in a closed tube to 160° forms SbOCl; heated to 140° in the proportion 30, H_eO:SbCl_s, Sb₄O₃Cl₂ is formed (Schäffer, A. 152, 314).—4. Aqueous solution of sodium thiosulphate reacts on solution of SbCl, to form a double compound of Sb.O. and Sb.S. probably Sb₂O₃.Sb₂S₃ (v. Antimony, oxysulphides of).—5. Boiling SbCl₃ dissolves powdered antimony trisulphide, on cooling a crystalline mass of sulphochloride, SbSCI.7SbCl,, is obtained; on washing with alcohol 2SbSCl.3Sb,S, remains (Schneider, P. 108, 407).

Combinations .- 1. With chlorine; SbCl, is formed.—2. Ammonia forms SbCl₃.NH₂ which on warming gives off all its NH₂.—3. By mixing conc. solutions of SbOl, and alkaline chlorides and evaporating, double salts are formed, c.g. 2NH, Cl.SbCl₃; 2(BaCl₂,SbCl₃).3H₂O; 8KCl.SbCl₃; 3MaCl.SbCl₃. With KBr the salt SbCl₂3KBr is formed identical with SbBr₂.3KCl obtained by the action of KClAq on SbBr. (Atkinson, C. J. 43, 290).

II. ANTIMONIO CHLORIDE SbCl., Mol. w. un-

known; vapour obtained by heating consists of SbCl, +Cl, [-6°] (Kämmerer, B. 8, 507). S.G. 25° 2°346 (Haagen, P. 181, 117). (79° at 22 mm.; 68° at 14 mm.) (Anschütz a. Evans, C. J. 49,

Preparation .- Powdered Sb is heated in a retort in a rapid stream of dry Cl; SbCl, (and Cl) distils over, and SbCl, remains. molted SbCl, is saturated with Cl, and distilled in a stream of Cl (or under greatly diminished pressure, Anschütz a. Evans, C. J. 49, 708).

Properties. - Colourless, or slightly yellow, liquid, with an offensive smell, furning in moist air; solidifies at a low temperature $(<-6^{\circ})$; absorbs moisture from air and changes to a crystallino mass. According to Anschütz a. Evans (C. J. 49, 708) SbCl, may be distilled nn.

changed at low pressures.

Reactions .- 1. Dissolves in a very little water; solution over H₂SO, deposits crystals of SbCl₂4H₂O. Kept cold by ico, and water added drop by drop in proportion SbCl.: H.O, SbOCl. (q.v.) is formed (Dubrawa, A. 184, 118). Addition of more water produces SbO2Cl, which is decomposed by hot water, giving H,Sb,O,2H,O, soluble in HClAq. Decomposition by H2O hindored by tartaric acid. Thomsen (Th. 2, 212) gives the number [SbCl⁵, Aq]=35,200, when Sb₂O₃Aq and HClAq are formed.—2. Dry sulphuretted hydrogen produces white crystals of SbSCl, which are decomposed by heat into SbU, and S₂O₄ (Clocz, J. pr. 51, 459).—3. Heated in closed tube to 140° with antimonic oxide, in proportion 3SbCl,:Sb₂O₂, Sb₃OCl₁, and Sb₄O₄Cl, (q. v.) are produced (Williams, C. J. [2] 10, 122).—4. With phosphorus trichloride (in CHCl3) reacts to form PCl, SbCl, and SbCl, (v. Combinations) .- 5. Chlorinates many carbon compounds, e.g. CHCl, to CCl4, C2H4 to C2H1Cl2, &c. (v. Chloro-compounds). 6. With nitrogen tetroxide forms SbCl. NOCI (Weber, P. 123, 347).

Combinations.—1. With ammonia forms brown SbCl, 6NH, which may be sublimed unchanged. - 2. With hydrocyanic acid forms white crystals of SbCl, 3HCN, which volatiliso with partial decomposition under 100°, and are decomposed by 11,0 (Klein, A. 74, 85). -3. With gascous cyanogen chloride forms ShCl. CNCl (Klein, l.c.). - 4. Combines with some non-metallic chlorides to form double compounds which usually deliquesco in air and are decomposed by usually deliquesco in air and aire decomposed by heat; the more important are Slocl, PCl. (Weber, P. 125, 78; Kühler, B. 13, 875); SbCl., POCl., SbCl., SeCl., and SbCl., SCl. (Woben, Le.); and SbCl., SeCl., (Weber, P. 125, 325).—5. Also combines with C_eH, PCl₄ to form SbCl₃, C_eH, PCl₄ (Köhler, B. 13, 1626).—6. Combines with various deloids and with the (Weilberg, C. 19). alcohols, and with ther (Williams, C. J. [2]

15, 463).

Antimony, fluorides of. Sb.O. dissolves in HFAq to form SbF.; Sb.O. xH.O dissolves in HFAq to form SbF. Neither has been gasified, so that mol. ws. are unknown.

I. Antimonious fluoride SbF, [abt. 2920]

(Carnelloy, C. J. 33, 275).

Preparation .- (Berzolius, P. 1, 34; Dumse A. Ch. [2] 31, 435; Flückiger, A. 84, 248).
 By dissolving Sb₂O₃ in HFAq, evaporating at 70°-90° and crystallising .- 2. By distilling Sb with HgF2.

Properties.-White, trimetrio, octahedral

deliquescent, soluble in H₂O without decompo- of SbCl, in conc. HClAq on to granulated Zn; sition.

Reactions .- - 1. Solution in water on evaporation yields an oxyfluoride (composition unknown).—2. Deliquesced SbF₃ pressed between paper gives 2SbF₃.Sb₂O₃ (=3SbOF.SbF₃), which is decomposed by heating into ShF, and Sh.O.s.

Combinations. — With alkali fluorides to

form double compounds, SbF_3 combining with MF, 2MF, or 3MF, where M = K, Na, &o. These compounds are bost obtained by dissolving Sb.O. and M₂CO₃ in the proper proportions in HFAq, and evaporating. The principal compounds are SbF_{3.2}NH₁F; SbF_{3.2}KF, SbF_{3.4}KF; SbF_{3.2}LiF; and SbF₃.2NaF (v. Flückiger, A. 84, 248).

II. Antimonie fluoride, SbF_s. Obtained by Berzelius, investigated more fully by Marignae

(A. 145, 239).

Preparation.—By dissolving hydrated Sb.O.

in HFAq, and evaporating.

Properties. - A gum like amorphous mass decomposed by heat; very slowly decomposed, in solution, by H.S.

Combinations .- With the alkali fluorides, to form double compounds, which are easily soluble in water, orystallise badly, and yield oxyfluorides when evaporated in aqueons solutions, e.g. SbOF, NaF from SbF, 2NaF (Marignao, A. 145, 239). These solutions are very slowly decomposed by H.S. KOHAQ, and K.CO,AQ. The more important compounds are SbF,NH,F, 2(SbF,2NH,F).H₂O; SbF,KF, SbF,2KF.2H₂O; SbF. NaF.

Antimony, haloid compounds of. SbF3, SbF5; SbCl₃, SbCl₅; SbBr₃; SbI₃, (?SbI₅). Only SbCl₃ has been gasified and V.D. determined: SbCl. is decomposed by heat. The formulæ of the tribaloid salts are probably molecular, v. Axti-MONY, FLUORIDES OF, CHLORIDES OF, BROMIDES OF, TODIDES OF; v. also art. Haloid compounds.

Antimony, hydride of (v. also art. Hydrides). (Antimoniuretted hydrogen; Stibine). Only one hydride of Sb, SbH, is certainly known; and this has not been obtained except mixed with much H. Marchand (J. pr. 31, 381) described a black powder obtained by electrolysing conc. NH,ClAq with a rod of Sb as neg. and a thick Pt wire as pos. electrode. When a powerful battery was used, gas came off which burnt in the air; the powder was supposed to be a solid hydride of Sb, and the gas a spontaneously inflammable hydride (v. also Ruhland, S. 15, 418). But Marchand's results were not confirmed by Böttger (A. pr. 68, 372), who obtained only the ordinary products of the electrolysis of NH, ClAq, viz. H, NH₂, and N chloride. Wiederhold (C. C. 1864. 995) described a graphite-like powder obtained by the action of dilute HClAq on an alloy of 1 part Sb with 5 parts Zn; after drying at 100° this powder gave off '001 p.c. II at 200° (Sb.H requiros '004). The gaseous hydride is almost certainly SbH's (v. especially Jones, C. J. [2] 14, 617), but it has not yet been obtained free from H.

Preparation.—By treating an alloy of 2 parts Zn and 1 part Sb (Capitaine, B. J. 20, 89), or 3 Zn and 2 Sb (Lassaigne, B. J. 22, 101), with dilute H.SO.Aq. Schiel (A. 104, 223) decomposes an alloy of Sb with K with dilute HClAq. Humpert (C. C. 1865, 863) treats conc. SbCl₃Aq with Na amalgam. Jones obtained a gas containing about 4 p.c. SbH, by dropping conc. solution

the gas was partially decomposed as it was formed (C, J, [2]) 14, 641).

Properties. - A colourless gas, with nauseating smell and intensely disagreeable taste, slightly soluble in H.O but decomposed by long contact into Sb and H; easily decomposed by heat: burns in air to Sb₂O₅ and H₂O, or in limited supply of air to Sb, Sb₂O₆, and H₂O; decomposed by electric sparks into Sb and H.

The gas obtained by reacting on an alloy of 2 parts Sb with 3 parts Zn with dilute H.SO.Aq, and collecting the first portions only, solidified at -91.5°, and decomposed, with separation of Sb, between -65° and -56° (Olszewski, M. 7.

Reactions.-1. With oxygen and heat, explosion occurs and formation of Sb₄O₆ and H₂O; the same products are obtained by burning in air. - 2. Decomposed by chlorine, bromine, or iodine, with formation of SbCl, SbBr, or SbI3; passed through a hot tube containing a little I, an orange-yellow or brown ring of SbI_s is formed (Husson, J. pr. 106, 314).—3. Passed over sulphur in snnshine, or at temperatures over 100°, orange-coloured Sb.S. is formed; very minute quantities of the gas may be thus deteeted $(2ShH_3 + 6S = Sh_2S_3 + 3H_2S)$ (Jones, C. J. [2] 14, 649).—4. Decomposes sulphuretted hydrogen in sunshine forming Sb_2S_3 ($2SbH_3 + 3H_2S = Sb_2S_3 + 6IL_1$) (Jones, l.c.).—5. With antimonious chloride, Sh and HCl are formed .- 6. Easily oxidised by nitric acid .- 7. Decomposed by aqueous potash or soda with separation of a black powder (? SbOH, or ? Sb.O, v. Jones, l.c.; also Dragendorff, Fr. 5, 200) which is at once dissolved on shaking in air. -8. With aqueous silver nitrate the whole of the Sb is ppd. (as Ag, Sb mixed with Ag, Lassaigne, B. J. 22, 104; v. also Jones, l.c.).

References. - Thompson, B. J. 18, 135; Pfaff, P. 40, 339; Simon, P. 42, 369; Vogel, J. pr. 13, 57; Meissner a. Hankel, J. pr. 25, 243.

Antimony, hydroxides of. Several compounds of Sb, II, and O are known; some of them are probably best regarded as hydrated oxides: others react as acids, especially HSbO, and H4Sb2O7; v. Antimony, Acids of (v. also arts. Acids and Hydaoxides).

Antimony, iodide of. ShI2. Only one iodide of Sb is known with certainty; van der Espt (Ar. Ph. [2] 117, 115) asserts that SbI, is produced by heating 1 part Sb with 5 parts I, or by leading SbH, into I in alcohol; but as SbI, is known to be produced by such processes the existence of the pentiodide is extremely doubtful (comp. Pendleton, C. N 48, 97). Mol. w. 499.62. 167°) (Cooke, P. Am. A. [2] 5, 72). (401° at 760 mm.) (Cooke, P. Am. A. [2] 7, 251). V.D. 252 (Worcester, P. Am. A. [2] 10, 61). S.G. hexagonal 21° 4.818, monoclinio 21° 4.768 (Cooke, l.c.). H:F. solid Sb, gaseous I, [Sb, I] = 45,400 (Guntz, C. R. 101, 161).

Formation.-1. By the action of powdered Sb on I in CS2 .- 2. By the action of SbH2 on I. -3. By subliming together Sb₂S₈ with 3I in 8 globe (Schneider, P. 109, 609).

Preparation.—Powdered So is added little by little to I, with gentlo heating, until no further action occurs; the SbI, is then separated by

sublimation in H or CO.

Properties. - Red orystals, which melt on

heating and volatilise in red vapours : soluble in boiling CS, and boiling benzene, but separates cut on cooling; almost incoluble in CHCl.; soluble in HIAq. Exists in three forms (a) hexagonal ruby-red crystals, by crystallisation .from CS_2 , M.P. = 167° , a:c = 1:1:37; (b) trimetric greenish-yellow crystals, by subliming the hexagonal form at temperatures not above 114°; at 114° the change is sudden, the external form of the hexagonal crystals is preserved but each crystal is found to consist of a mass of trimetric crystals: heated above 114° the hexagonal form is reproduced; (c) monoclinic orystals (a:b:c = 1.6408:1:6682) obtained by exposing a solution of SbI3 in CS2 to direct sunlight; at 125° they are changed into the hexagonal form (Cooke, P. Am. A. [2] 5, 72).

Reactions.—1. Water decomposes SbI, with production of IIIAq, which dissolves part of the SbI, and oxyiodito of Sb (q. v.).—2. Aqueous alkalis and alkali carbonates produce Sb2O, and alkali iodide (Scrullas, J. Ph. 14, 19).—3. Conc. sulphuric acid or nitric acid separates I.—4. Alcohol or ether partly dissolves SbI, and partly changes it to yellow oxyiodide (Macivor, C. J. [2] 14, 328).—5. Chlorine torms SbCI, and ICI (Macivor, I.c.).—6. Antimony trisulphide reacts with molten SbI, to form SbSI; this sulphiodide is obtained as a lustrous brown-red powder by treating the fused mass with dilute HClAq; it is decomposed by H,O and KOHAq; boiled with H,O and ZuO the oxysulphide Sb,OS; is formed (Schneider, P. 110, 147).

Con.binations.—Dissolves in aqueous solutions of the iolides of the athali metals, on evaporation double compounds are obtained. These compounds are soluble in HClAq, H.C.,H.O.,Aq, and H.C.,H.O.,Aq; they are decomposed by H.O., yielding Sb oxyiodide; CS_dissolves out SbI_3. The following salts are described by Schneffer (P. 109, 611): 28bI_38K13H_2O; 28bI_38K13H_2O; 48bI_38KH_19H_O;

25bL,3Nat.12H.O; 4SbL,3NH,L9LLO; SbL,BhL,2HLO. Nickles (C. R. 51, 1097) describes two series of compounds MLSbl,2HLO, and MLSbl,HLO where M = K, Na, or NH, obtained usually by the action of I on Sb in presence of saturated MIAq; these salts are isomorphous with corresponding double salts of Bi.

Antimony, livers of. This name is applied to the impure double sulphides obtained by heating Sb,S, with various metallic sulphides, more especially with the alkali and alkaline earth sulphides. These bodies are obtained either by fusing Sb₂S, with K₂S, &c., with K₂SO₂ &c. and C, or by dissolving Sb₂S, in K.SAq, &c. The behaviour of aqueous solutions varies according to the relative quantities of Sb₂S₃ and alkali sulphide employed; if not more than 2 parts Sb.S. are used to 1 part alkali sulphide, the product is wholly soluble it. water; if more Sb₂S₃ is used the product is partly, or wholly, insoluble (because of production of antimonate and Sb_iO₆ v. Antimonious sulphide, Reactions, No. 15). Solutions of these bodies dissolve Sb.S, on boiling; the Sb.S, pp3. again on cooling; they readily absorb O from the air, forming antimonate and thicantimonate (v. Antimonious sulphide, Reactions, No. 15). Addition of alkali bicarbonates pps. thioantimonite.

Antimony, oxides of. Three oxides are known: Sb₂O₄, Sb₂O₄, and Sb₂O₄; only the first of these has been gasified; the molscular weights of the others are not known. The pentoxide acts as an anhydride; the two others are feebly salt-forming whether they react with strong acids or strong alkalis (v. ANTIMONY, ACIDS OF). Marchand (J. pr. 34, 381) described an oxide, Sb₂O₂, said to be obtained by the electrolysis of a solution of eream of tartar; but l'Ottger (J. pr. 68, 372) failed to obtain anything except antimonic acid by repeating the experiments.

I. Antimonious oxide Sb₄O₆ (Antimonious acid). Mol. w. 575-76. S.G. trimetric 5-5 to 5-6, regular octahedra 5-1 to 5-2 (Terreil, C. R. 58, 1209). V.D. 286-5 (at abt. 1550°; Meyer, B. 12, 1284). S.H. 0927 (18° to 100°; Neumann, P. 126, 123). C.E. (40° cub., Senarmontite) 00005889 (Fizzan, A. Ch. [4] 8, 335).

Occurrence. - Native; as Antimony bloom in trimetrie prisms, as Senarmontite in octahedra.

Formation.—1. By heating Sb in a loosely covered crucible, and then raising the temperature, when Sb,O₈ mixed with a little Sb,O₄ sublines on to the crucible cover.—2. By treating Sb with dilute IINO₃Aq and washing thoroughly with water and then with vory dilute Na,CO₃Aq (Rose, P. 53, 161).—3. By fusing Sb with KNO₃ and KHSO₄ and boiling fused mass in water (Preuss, A. 31, 197).—4. By washing the white pp, obtained by adding H,O to SbCl₃ with dilute KOIIAq and then with H₂O.—5. By adding excess of NH₃Aq to hot KSb,C,H,O,Aq, heating pp. for a short time in contact with the liquid, collecting, and washing.

Preparation. -1. 3 parts finely powdered Sb are heated with 7 parts cone. II,SO,; the evide Sb sulphate is treated repeatedly with hot water, and then with very dilute Na,CO,Aq, and the oxide is collected and dried. -2. 1 part powdered Sb is heated, so long as an action occurs, with 4 parts IINO,Aq, S.G. 1.2, and 8 parts II,O; the nitrate of Sb is treated as the sulphato in 1.

I'roperties. - A white, more or less crystalline. powder (regular octuliedra); very slightly soluble in water, fairly soluble in glycerine (Köhler, D. P. J. 258, 520); becomes yellow when heated. but white again on cooling; melts at a dark red heat, and erystallises on cooling. Volatilises rapidly about 1550° (Meyer, B. 12, 1281). Insolublo in HNO, Aq and H, SO, Aq; dissolves casily in HClAq and H,C,H,O,Aq; also in KOHAq and NaOHAq, from these solutions Sb,O is ppd. on cooling (Mitscherlich, A. Cr. [2] 33, 394), but according to Terreil the pp. is an antimonite (A. Ch. [4] 7, 380). Sb.O. is formed in trimetric prisms (a.b.c. = 334:1:1-414) by burnather. ing Sb or Sb,S, in air, by heating oxychloride (obtained by adding H.O to SbCl3) with H.O to 150° (Dobray, C. R. 58, 1209), or by rapidly subliming the octahedral crystals (Terreil, C. R. 62, 302); Sb.O. is formed in regular octahedra by subliming at a dark red heat. Both forms are obtained by saturating hot Na CO Aq with Sb.O. or SbCl, and allowing to cool (Mitscherlich, P. 15, 453); or by passing a slow stream of dry air through a porcelain tube containing Sb, the tube being heated at first only where the Sb is, but after a few hours also at the point where the prisms might condense, after about 12 hours prismatic crystals are found near the Sb, prisms

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mixed with octahedra further on, and octahedra only near the end of the tube (Terreil, I.c.). Sb,O. is isodimorphous with As O. (q. v.). According to Guntz (C. R. 98, 303) the change of prismatic Sb.O. to octahedral is attended with production of 1200 gram units of heat per 576 grms. Sb.O. changed.

Reactions.—1. Heatod in air or oxygen, Sb.O. is formed. -2. Conc. hot nitric acid oxidises to

Sb2O, and Sb2O; it dissolves in cold furning IINO; and forms Sb,O, N,O, (Peligot, C. R. 23, 709). 3. Treated with furning sulphuric acid, small lustrous erystals are obtained, which, after drying for six months in contact with burnt clay have the composition Sb₄O₆.2SO₂; by treating these crystals with H₂O the salt Sb₄O₆.SO₂ is obtained (Peligot, Le.). Schultz-Schlac (B. 4, 13) describes the salt Sb. 3SO₄ (= Sb₄O₂.6SO₂) as long lustrous noedles obtained by evaporating solutions of Sb,O, in fairly conc. H,SO,Aq; this salt is unchanged in dry air, but gives off SO, on heating, and is decomposed by water. Sb₂(SO₄)₃ is also formed by dissolving Sb.S. in hot cone. H.SO, Aq (Hensgen, R. T. C. 4, 401) (v. Sul-PHATES).-4. Sb.O. dissolves in solution of potassium hydrogen tartrate, forming the salt C₁H₁KSbO₂, which is probably the K salt of the acid Sb.C.H.O. OH (v. Clarke and Stallo, B. 13, 1787). - 5. Sb O acts as a reducing agent towards salts of silver, gold, &c. (v. Antimony, Detec-Tion or, Antimonious compounds).-6. Sh,Oa does not directly combine with water, but two hydrates have been prepared: -(a) Sb.O.2II.O. by adding CuSO, Aq to Sb, S, dissolved in KOHAq until the filtered liquid gives a white pp. (Sb, O, 2H, O) on addition of an acid (Fresenius; v. also Schaffner, A. 51, 182); (b) Sb O_x.311₂O, a white powder which begins to lose water abovo 150°, obtained by the spontaneous decomposition of an aqueous solution of the acid H.C.H.SbO, obtained by decomposing (C₁H₄SbO₂). Ba by the proper quantity of H₂SO₄Aq (Clarke a. Stallo, B.13, 1793).—7. Dissolves in boiling ant monious chloride to form oxychlorides; ShOCl.7SbCl, is described by Schneider (P. 108, 407).

II. ANTIMONIC OXIDE Sb.O. (Antimonic acid). Mol. w. unknown. S.G. 3.78 (Playfair a. Joule,

C. S. Mem. 3, 83).

Preparation. - By dissolving powdered Sb in aqua regia, or conc. HNO, Aq, evaporating to dryness, and heating [not above 275°] (Cleuther, J. pr. [2] 4, 438; Dubrawa, A. 186, 110).

Properties.—Citron-yellow powder; insoluble in water, but reddens moist blue litmus paper; loses O at 300' (Geuther, Le.) giving Sb.O.; soluble in conc. HClAq, slightly soluble in conc. KOHAq.

Reactions .- 1. Heated with antimony or antimony sulphide, Sb.O. is formed .- 2. Heated in chlorine, SbCl, and Sb,O, are produced. - 3. Heated with ammonium chloride, is completely volatilised .- 4. Reacts with alkaline earbonutes on fusion, with evolution of CO, (v. further ANTIMONY, DETECTION OF, Antimonic compounds).-5. Hydrates are not produced by the direct action of water, but indirectly the three compounds, Sb₂O₅3H₂O₇ Sb₂O₅2H₂O₇ and Sb₂O₅.H₂O₇ have been obtained (v. Antimony, ACIDS OF, ANTIMONATES). Forms many compounds with WO₃ and MoO₃ (v. Gibbs, C. N. 48, 155; Am. 7, 209 and 313).

Antimonoso-Antimonic oxide Sb.O. (Antimony tetroxide). Mol. w. unknown. S.G. 4.074 (Playfair a. Joule, C. S. Mem. 3, 83); 6.5 (Boullay, A. Ch. [2] 43, 266). S.H. (23°-99°) 09535 (Regnault, A. Ch. [3] 1, 129).

Occurrence .- Native, as Antimony ochre. Preparation.-1. By heating Sb,Oe in air.-2. By oxidising Sb, Sb, Oa, or Sb, Sa, by cono. HNO, Aq, evaporating to dryness, and strongly

heating.

Properties.—White powder, becoming yellow on heating; has not been melted or volatilised: insoluble in water, but reddens moist blue litmus paper; very slightly acted on by acids.

Reactions .- 1. Heated with solution of ercam of tartar, $\operatorname{Sb}_{s}\operatorname{O}_{s}x\operatorname{H}_{s}\operatorname{O}$ remains and solution contains C.H.KShO, -2. Solution in HClAq dropped into water, is decomposed into Sb,O, and Sb,O, .-- 3. Heated with antimony Sb₁O₆ is formed.—4. With molten potash forms K₂O.Sb₂O₄ (v. ANTIMONY, ACIDS OF; ANTIMONOSO-ANTIMONATES); solution of this in water slowly reduces AgNO3Aq and AuCl₃Aq. Sb₂O₄ reacts as a compound of Sb₂O₃ and Sb₂O₅ (= Sb₄O₈); it is sometimes regarded as antimonyl antimonate (SbO)SbO₂₁ derived from HSbO₃.

Antimony, oxybromidos ef. Two oxybromides are obtained by the action of H.O on SbBra, viz. Sb₄O₅Br₂ and 10Sb₄O₅Br₂SbBr₃, SbOBr is probably formed by the action of smalight on SbBr.

in CS, (v. Antimony, bromide or).

Antimony, oxychlerides of. compounds are known; SbOCl, SbOCl.7SbCl, Sb₁O₂Cl, and 10Sb₄O₂Cl, SbCl, obtained from SbCl₃; SbO₂Cl, and SbOCl₄, from SbCl₅.

When SbCl, is added to a little water, SbOCl is obtained (sometimes mixed with SbCl₂). This oxychloride seems to exist either as a white amorphous powder, or as monoclime erystals isomorphous with SbOI (Cooke, P. Am. A. [2] 5, 72); the crystals are best obtained by using 10 pts. SbCl, and 17 pts. H.O, allowing to stand for a day or two, pressing, and washing with other (Sabanajew, Bt. [2] 16, 79); the amorphous powder is best prepared by adding 3 pts. H.O to 1 pt. SbCl₃, filtering at once, drying over H₂SO₄, and washing with ether. Crystalline ShOCl is also obtained by heating SbCl, with C2H6O (in ratio SbCl₃:C.H₂O) in a closed tube to 160° (Schäffer, A. 152, 314). By the action of much water on SbCl3 (5 to 50 parts to 1 part SbCl, according to Sabanajew, I.e.) the oxychloride Sb₁O₂Cl₂ is obtained as an amorphous powder, which becomes crystalline on standing. To prepare the erystalline forms it is best to use 30 parts of cold water, or 3 parts of water at 60° to 70° (in the latter case allowing the pp. to romain a few hours before collecting); there are some differences in the forms of the two sets of orystals (Sabanajew, l.c.). Crystals (trimetrio Schäffer, A. 152, 314; monoclinic, Cooke, P. Am A. [2] 5, 72) of Sb₁O₁Cl₂ are also obtained by heating SbCl₂ with C₂H₄O (in ratio SbCl₂:3C₂H₄O) to 140°-150° (Schäffer, Lc.). The compound Sb,O.Cl, is also produced by the action of alcohol on SbOCl (Schneider, P. 108, 407); and also by heating dry SbOCl (5SbOCl & SbOCl + SbCl; Sabanajew, l.e.).

The product of the action of much H2O on SbCl, is known as powder of Algaroth; the composition varies according to temperature, quantity of water, and quantity of HCl in the solution of SbCl, used (comp. Duflos, S. 67, 268; Johnston, J. pr. 6, 55; Malaguti, J. pr. 6, 253; Peligot, A. 64, 280).

According to Williams (C. N. 24, 224) the action of hot water on SbCl3 produces 10Sb,Cl2O3.SbCl3; Williams also describes two oxychlorides obtained by heating Sb.O. with SbCl₃ (Sb₂O₅:3SbCl₃) to 140° in a closed tube;

Sb₄OCl₁₃ M.P. = 85°, and Sb₄O₂Cl₁ M.P. = 97·5°. Cooke (P. Am. A. [2] 5, 72) describes another oxychloride Sb₀O₁₁Cl₂; and Schneider (P. 198, 407) two others, SbOCl.7SbCl₃ and 2SbOCl.Sb₂O₃.

According to Thomsen (Th. 2, 240) the heat

of formation of Sb,O,Cl, from SbCl, and Aq is 8910 gram-units (v. also Guntz, C. R. 98, 512). By dropping the calculated quantity of very cold water on to SbCl₂, Dubrawa (A. 184, 118) obtained SbCl₃ (SbCl₃ + II₄O = SbCCl₃ + 2IICl); this oxychloride is a yellowish, somewhat crystalline, mass, solublo in alcohol; it deliquesces to a yellow liquid, from which needle-shaped crystals separate in dry air. When heated it melts and decomposes (probably to SbOCl + Cl.). It is decomposed by $Na_{2}CO_{3}Aq$ ($2SbOCl_{3} + 3Na_{2}CO_{3}Aq =$ 6NaClAq + 2O + 3CO, + Sb,O₃).

Antimony, oxyfluoride of, 3SbOF.SbF₃ ob-

tained by deliquescence of SbF, (v. Antimony,

FLUORIDES OF).

Antimony, expieddes of. Sb,OsL, and SbOL Sb₄O₅I₂ is obtained as light yellow crystals by ovaporating a solution of SbCl₃ in KIAq, adding H.O and evaporating again. The composition of the oxylodide obtained by the action of H.O. on SbI, varies according to the conditions of its preparation; by pouring Sb1, in IIIAq into hot II,0, Sb,0,I, is obtained (v. Macivor, C. J. [2] 14, 328). By the action of air and smalight on SbI₃ in CS₃ both oxylodides are formed, production of SbOI proceeding rapidly (Cooke, P. Am. A. [2] 5, 72). When SbOI is heated in a current of an inert gas to 150°, SbI3 begins to sublime, and at 200° is given off rapidly; no further change occurs till 350° is reached, when Sb1. again sublimes and crystals of Sh₂O₃ remain (Cooke, P. Am. A. [2] 5, 72). By the action of HCIAq, HNO, Aq, or H, SO, Aq, on SbO1, I is separated.

Antimony, oxysulphides of. Various oxyeulphides of Sb, or more probably mixtures of Sb₂S₃ and Sh₂O₃, were formerly used in pharmacy. The compound Sb.O., 2Sb.S. occurs native as antimony blende (v. H. Rosc, P. 3, 452). The oxysulphide Sb₂OS₂ is obtained as a rodbrown powder by hoiling SbSI (v. Antimony, ioxide of) with ZnO and H₂O (Schneider, P. 110, 147); also by the action of Na.S.O.Aq on SbCl, in HClAq (Böttger, C. C. 1857, 333). A compound of Sb₂S₃ and Sb₂O₃ is much used as a brilliant crimson red pigment; it is probably Sb₂O₂2Sb₂S₃ (= 3Sb₂OS₂), (v. Böttger, l.c.; Wagner, J. 1858. 235; Kopp, C. C. 1859. 945). Antimony, phosphides of, v. Antimony, Com-

binations, No. 8.

Antimony, selenides of, v. Antimony, Combinations, No. 6.

Antimony, ssienc-acid of, v. Seleno-Antimonates, p. 286.

Antimony, sulphides of. Two sulphides are known, Sb₂S₃ and Sh₂S₃; neither has been gasified, and therefore mol. w. of neither is known. Unger (Ar. Ph. [2] 147, 193) supposed he had obtained a disulphide, Sb_sS_s , by the action of NaOHAq on Sb, Sa; but the existence of this Sb₂S₂ is very doubtful. Sb₂S₃ is a feebly marked salt-forming sulphide, c.g. it dissolves in NaOHAq to form Na Sb S, (v. ANTIMONY, THIO-ACIDS OF). Sb.S. is a distinctly salt-forming sulphide; the thie antimonates (q. v.) are well-marked salts. The trisulphide, Sb₂S₃, occurs native; the pentasulphide does not.

I. ANTIMONIOUS SULPHIDE (Antimony trisulphide, Mineral Kermes, &c.) Sh.S. [low red heat]. S.G. (stibnite) 4.51-4.75; (amorphous) 4.15; (fused, by direct union of Sb and S) 4.892 (Ditte, C. R. 102, 212). S.H. (23°-99°) 08433 (Regnault, A. Ch. [3] 1, 129; v. also Neumann, P. 23, 1). Two forms are known; crystallised (trimetrio;

a:b:c = 985:1:1.0117) and amorphous.

Occurrence. - Native as Stibnite or Antimony glance, crystallised in trimetric prisius, usually containing P, As, Fe, and Cu.

Preparation .- (a) Crystallised: by gradually heating to redness, in a covered crucible, a mixture of 13 parts of finely powdered Sb well mixed with 5 parts pure S; thou fusing for some time under a layer of NaCl; cooling, powdering, mixing with a little S, and again fosing under NaCl. (b) Amorphous: by boiling 4 parts KOllAq, S.G. 1.25, and 12 parts H.O, with one part crude Sb.S. out of contact with air for some time, adding 50 parts boiling H.O. filtering quickly, and decomposing the solution by dilute ILSO, Aq; the pp. is collected, boiled with very dilute H2SO, Aq, washed with cold water, digested with agneous tartaric acid (to remove any Sb.O.). again washed with cold water, pressed, and dried at a low temperature. Cooko (P. Am. A. [2] 5, 1) dissolves Sb in large excess of HNO₃Aq (S.G. 135), keeping the temperature as low as possible, neutralises with NaOllAq, dissolves in large excess of H₂C₄H₄O₆Aq, pps. by H₂S in an atmosphere of CO₂, collects and washes pp. and dries below 210°. The amorphous sulphide is also produced by melting crystalline Sb S, in a glass tube, and after a time throwing it into a large quantity of cold water (Fuchs, P. 31, 578). An impure Sb.S3, containing Sb.O., known as Kermes, is prepared for commercial purposes by heating crude antimony sulphide with aqueous alkalis or alkalino carbonates.

Properties. - (a) Crystallino: grey-black trimetric prisms; melts easily. (b) Amorphous: prepared by ppn., is a reddish-brown, loose, powder which marks paper with a brownish streak; prepared by melting and suddenly cooling the crystalline Sb.S3, it is a hard greyish mass; moltod and cooled slowly it yields the crystalline form; heated to 210 -220° it becomes grey (Cooke, P. Am. A. [2] 5, 1). Both forms of Sb.S. aro insoluble in water, and in NH, Aq, dissolve in KOHAq, in HClAq, and very slowly in tartaric acid. They may be distilled nu-ohanged in a stream of N.

Reactions .- The products of the reactions of orystalline and amorphous Sb.S, are, in almost every case, the same; the actions usually proceed more rapidly with the amorphous than with the crystalline form. 1. Boiled with water, is partially decomposed to Sb₁O₆ and H₂S (De Clermont a. Frommel, C. R. 87, 330; Lang, B. 18, 2714). -2. Heated in hydrogen, Sb is formed.-

B. Calcined in air, Sb.O., or Sb.O., and SO. are produced.—4. Heated in chlorine, SbCl, and S2Cl, result.—5. Aqueous hydrochloric acid forms SbCl, and H.S; after a time the action stops, but if the H.S is removed the whole of the Sh₂S₃ is decomposed (v. Lang, B. 18, 2714; also Berthelot, C. R. 102, 22).—6. Conc. nitric acid oxidises to nitrate and sulphate of Sb mixed with S.-7. Aqua regia forms SbCl, H.SO, and S .- 8. Dilute solutious of sulphuric acid have no action on crystalline Sb.S3, but slowly evolve H.S from amorphous Sb, S,; conc. H, SO, Aq evolves SO., separates S, and forms Sb. (SO.), (Hensgen, R. T. C. 4, 401) .- 9. Fused with at least 17 parts nitre, KSbO, is formed, with Sb sulphate; with less than 17 parts nitre, Sh.O. is sometimes formed in addition to the other products, or a part of the Sb₂S₃ remains moxidised and combines with K₂S formed to produce a thio-salt.— 10. Melted with excess of lead oxide, Sb2O2, (and Pb), are formed .- 11. Melted with potassium cyanide, Sb is produced, along with KCNS and a compound of Sb₂S₃ and K₂S.—12. Iron, zinc, and many other metals reduce Sb.Sa, when heated with it, forming Sb and a metallic sulphide; metals whose sulphides are basic (c.q. K) generally combine with part of the Sb2S, to form thio salts.—13. Many easily reduced metallic oxides when heated with Sb.S3 form Sb.O4 and SO, -14. Many metallic sulphides combine when heated with Sb.S. with production of double compounds; several of these double compounds occur native, e.g. Sb₂S₃.PbS; Sb₂S₃.Cu₂S; Sb₂S₃.3Ag₂S; &o.-15. Alkali sulphides combino with Sb.S, either when heated in the solid state or in solution; the compounds produced are generally known as livers of antimony, q. v. (v. also next reaction); aqueous solutions of these compounds absorb O forming Sb₁O₈, antimonates, and thio-antimouates.-16. Caustic alkalis react with Sb.S., when fused together, or when in aqueous solutions, to produce antimonito and thio-antimonite: thus, 4Sb.S. 4K.O= $\begin{array}{ll} 6KSbS_2 + 2KSbO_2 \; ; & \text{or} & 4Sb_2S_3 + 8KOHAq = \\ 3(Sb_2S_3 \; K_2SAq) \; \; [= 6KSbS_2Aq] \; + 8b_2O_3 \; K_2OAq \end{array}$ [=2KSbO2Aq]+4H2O. Addition of HClAq to this solution pps. Sb.S.; $(6KSbS_2Aq + 2KSbO_2Aq + 8HClAq =$

4Sb₂S₂ + 8KClAq + 4H₂O). If, however, much Sb₂S₃, relatively to the amount of KOH, is used, formation of KSbS, Aq proceeds, but the KSbO, being much less soluble pps. along with some oxysulphide and Sb.O. which has not combined with KOIL (crocus of antimony). The solution of KSbS, is acted on by air, giving finally antimonate and thio antimonate $(6KSbS_2Aq + 6O =$ 4KSbS₃Aq + 2KSbO₂Aq),—17. Solutions of carbonated alkalis, K₂CO₃ and Na₂CO₃Aq, dissolve Sb2S3 only on heating; the solutions behave similarly to those obtained by KOHAq and NaOHAq; on boiling in air a pp. of KSbO₃ (Sb₂O₃, K₂O) combined with Sb₂S₃ is obtained (Kermes), and KSbS, remains in solution. When 1 pt. Sb, S, is fused at a strong red heat with 3 pts. Na2CO3, and H2O is added, a solution containing antimonate and thio antimonate is obtained, and Sh is ppd. (probably, 10NaShO,Aq + 2H₂O = 6NaShO,Aq + 4NaOHAq + 4Sh). The action of alkalis on Sb₂S₂ has been chiefly investigated by Liebig (A. 7, 1).

Combinations .- With metallic sulphides to

form thio-antimonites (v. supra), q. v. under Anti-MONI, THIO-ACIDS OF.

Antimonious sulphide; hydrated. The orange-red pp. obtained by passing H₂S into a solution of SbCl₃ or C₄H₄KSbO, containing little acid is amorphous hydrated Sb₂S₃, which is fully dehydrated only at 200° (Fresenius; according to Wittstein the pp. contains no chemically combined H₂O, Fr. 1870. 262). It behaves towards acids, alkalis, &c. in the same way as amorphous Sb₂S₃. When this pp is treated with peroxide of hydrogen, in presence of NH₄Aq, a portion of it is oxidised to antimonic acid, some of which separates out and some remains in solution as NH₄SbO₂ (Raschig, B. 18, 2743).

II. ANTIMONIO SULPHIDE Sb.S. (Antimony peritasulphide or persulphide, Golden sulphine of antimony, &c.). Mol. w. unknown. Not found native. Sb₂S₂ does not directly combine with S; but by heating Sb₂S₃, S, and Na₂CO₃ together, Na₂SbS₄ is formed, from which Sb₂S₅ is obtained by the action of acids.

Formation.—1. By the action of H₂S on SbCl, in H₂C₄H₄O₄Aq, or on Sb₂O₃xH₄O suspended in water.—2. By decomposing solutions of thio-antimonates by dilute acid.

Preparation.—10 parts crystallised Na, SbS, 9H.O (q. v. under Antimony, thio-acids or) are dissolved in 60 parts H₂O; the solution is poured (with constant stirring) into a cold solution of 3.3 parts pure H₂SO₄ in 100 parts H₂O; the pp. is washed by decantation, then on a filter, with cold water, as quickly as possible; to remove all traces of acid, the pp. is now digested with a cold solution of 1 part NaHCO₃ in 20 parts H₂O for a few days; it is again washed then pressed, and dried in a dark place at a low temperature.

Properties.—A dark-orange powder; insoluble in water; completely soluble in aqueous alkalis; in absence of air, soluble in NH₂Aq, and in aqueous alkali sulphides; soluble in Na₂CO₃ or K₂CO₃Aq, not in (NH₄)₂CO₃Aq.

Reactions.—1. Heated out of contact with air, Sh₂S₃ and S are formed.—2. Decomposed by boiling with hydrochloric acid, giving SbCl₃ and H₂S₅.—3. Caustic alkalis dissolve Sh₂S₃, forming antimonate and thio-antimonate.—4. Carbon disulphide dissolves out a little S (about 5 p.e., Rammelsberg, P. 52, 193). That this is due to a decomposition of Sh₂S₃, and not to the action of CS₂ on a limited S (it has been supposed that the action of CS₂ proves the non-existence of Sh₂S₃), is shown by the fact that muoh less than S₂ is withdrawn from each Sh₂S₃, especially the solubility in NH₃Aq in which Sh₂S₃ is insoluble, and the insolubility in (NH₄)₂CO₂Aq which dissolves Sh₂S₃.

Combinations. - With alkali sulphides to form thio-antimonates, q.v. under Antimony, Thio-acids of.

Antimeny, sulphe-acids of, v. Antimony, Thio-ACIDS OF

Antimony, sulphe- (or thio)- chlorides of. SbSCl.7SbCl₃, and 2SbSCl.Sb₂S₃, obtained by action of Sb₂S₃ on SbCl₃; and SbSCl₃ obtained by the action of H₂S on SbCl₄ (v. Antimony, CHLOBIDES OY).

Antimony, salpho- (or this). iedides of. SbSI; by action of Sb₂S₂ on SbI₂, or of I on Sb₂S₂ (v. Antimony, iodide of).

Antimony, tellurides of, v. Antimony, Com-

binations, No. 7. Antimony, thio-acids of. No thio-acide of Sb are known, but a few thio antimonites, MSbS, and one M3SbS3, and a coneiderable number of well-marked thio-antimonates, M.ShS., have been prepared. The thio-antimonites may be regarded as derived from the hypothetical acid HSbS, (=SbS.SH); they correspond in composition with the antimonites MSbO, and with the meta-thioareonites MAsS. The thio-antimonates may be regarded as derived from the hypothetical acid H₃SbS₄ (=SbS(SH)₃); nc corresponding antimonates are known (MShO, and M,Sb,O, represent the antimonates); the thio-arsenites are represented by three series, one of which (the orthoseries) corresponds with the thie-antimonates.

Thio-antimonites. A very few of those salts have been prepared. Addition of absolute alcohol to a solution of Sb.S₃ in NaOHAq pps. amorphous NaShS₂, soluble in water. By heating to 30° equivalents of Sb.S₃ and NaOH (in cone. solution) copper-coloured 2NaShS₂.H₂O ie formed (Unger, J. 1871, 325). The silver ealt Ag₃SbS₃ is said to be obtained as a grey mase, reddish when powdered, by heating Ag₃SbS₃ out of contact with air (Rammelsberg, P. 52, 193). Several minerals may be regarded as thio-antimonites, e.g. PbS.Sb.S₃; Ag.S.Sb.S.: Ch.S.Sb.S.: FeS.Sb.S.S.

 $\begin{array}{lll} \textbf{Ag}_{s}\textbf{S}.\textbf{Sb}_{s}\textbf{S}_{s} ; \textbf{Cu}_{s}\textbf{S}.\textbf{Sb}_{s}\textbf{S}_{s} ; \textbf{FeS}.\textbf{Sb}_{s}\textbf{S}_{s} \text{ &c.} \\ \textbf{Time-antimonate} & \textbf{M}_{s}\textbf{SbS}_{s} & \textbf{Investigated} \\ \textbf{ehierly by Rammelsberg} & (P. 52, 193). & \textbf{Some of} \end{array}$ these salts are obtained by the action of alkali sulphides on Sb,S,; but they are better obtained by acting on Sb.S. with aqueous solutions of alkali sulphides in presence of oulphur, or with aqueous solutions of alkali polysulphides. They are also obtained by fusing Sb2S3 with alkali sulphides (or with sulphates and carbon) and sulphur. The decomposition, in air, of alkaline livers of antimony also ofton yields thio-antimonates (v. Antimonious suipnide, Reactions, No. 15 and 16). The thio-antimonates of the alkali metals are soluble in water, many of the others aro insoluble and are obtained from the alkali salts by ordinary double decompositione, the metallic salt solution being added in quantity less than sufficient to decompose the whole of the alkali thio-antimounte. Solutions of thio-antimonates are easily decomposed by acids, even by the CO2 of the air, with ppn. of Sb2S5, and, when exposed to air, of alkali thio-sulphate. The alkali thio antimonates are not decomposed by heating out of contact with air; the salts of the heavy metals lose S, and give thio-antimonites. The more important thio-autimonates are those of potaseium and sodinm.

Potassium thio antimonate. 2K, SbS, 9H, O; slightly yellow deliquescent crystals; prepared by boiling, for soveral houre, 1 part S, 6 parts K, CO,, 3 parte CaO, and 20 parts H, O, with 11 parts Sb, S,, filtering, and cooling out of contact with air. A calt K, SbS, KSbO, 6H, O ie obtained, in long white needles, by adding cold cone. KOHAQ to Sb, S,, filtering from 2KH(SbO_x), 6H, O which separates out, and evaporating (v. Schiff, A. 114, 202).

Sodium thio antimonate Na, SbS, 9H, 0

(known as Schlippe's salt). Prepared by fusing together 16 parts dry Na₈SO₄, 18 parte Sb₈B₃, and 4-5 parts wood charooal, diesolving in water, boiling with 2½ parte sulphnr, filtering and evaporating; the cryetale are washed with very dilute NaOHAq and then with water, and dried quickly at a low temperature. The ealt may also be prepared by boiling Na₅CO₅Aq with CaO, Sb₂S₃, and S. Sodium thioantimonate forms large yellowish monometric tetrahedra; it dissolves in 2.9 parts H₂O at 15°, the colution has an alkaline reaction. The crystale are book kept in contact with their mother liquor to which a little NaOHAq is added; they decompose in air, giving Sb₅B₃, Sb₅S₃, Na₂S, Na₂CO₄, when the mother liquor from Schlippe's salt is evaporated, crystals of a double salt Na₅Sb₅, Na₂S₀, 20H₄O are obtained. Solution of tartar emetic is decomposed by Na₅Sb₅Aq thue; \$G_1I_1KSbO,Aq + 2Na₅SbS,Aq

GC₄H₁KNaO₄ + Sb₄O₅ + Sb₅S₅ + Sb₅S₅.

The other thic antimonates are generally obtained from the codium salt: the best-marked are Ba₃(SbS₂)₂, GH₂O₅ Cu₃(SbS₃)₂; Pb₃(SbS₃)₂; Hg₃(SbS₃)₂; Ag₃SbS₃; a few others are known (e.g. of Cd, Co, Mn, Ni, V, Zn) but they are very easily decomposed, and not many have been obtained in definite forme.

M. M. P. M.

ANTIMONY, Compounde with organic radicles.

References.—Löwig a. Schweizer, A. 75, 315; Landolt, J. pr. 52, 385; 57, 134; 84, 330; A. 78, 91; 84, 44; Buckton, C. J. 13, 115; 16, 17; Löwig, A. 88, 323; 97, 322; C. J. 8, 261; Berlé, J. pr. 65, 385; Scheibler, J. pr. 64, 505; Friedlünder, J. pr. 70, 449; Cramer, Pharm. Cent. 1855, 465; Hofmann, A. 103, 357; Strecker, A. 105, 300; v. Rath, P. 110, 115; Jörgeseen, J. pr. [2] 3, 342; Le Bel, Bl. [2] 27, 444; Michaelis a. Roese, A. 233, 42.

Tri-methyl-stibine SbMe₃. Mol. w. 167. (81°), S.G. $\frac{15}{2}$; 1·523.

Preparation. - An alloy of antimony (4 pts.) and sodium (1 pt.) ie mixed with sand and MeI and distilled. MeI and SbMe₂ pass over but unito in the receiver to form SbMe₄I, which when distilled with an alloy of antimony and potaesium in a current of CO₂ gives SbMe₃ (Landolt).

Properties.—Liquid, smelling of onions, sl. eol. water; may take fire in air. Takes fire in chlorine. Reduces salts of silver and mercury.

Salte.—Unites directly with non-metals.—SbMe₂Cl₂: hexagonal crystals, al. soluble in water. Formed also from SbCl₂ am' HgMe₃.—SbMe₂Cl₂SbMe₂O: octahedra, sol. water.—SbMe₄Br₂—SbMe₄Br₃SbMe₂O: octahedra, sol. water.—SbMe₄L: formed by heating Sb with MeI at 140°.—SbMe₄L: SbMe₂O: octahedra.—SbMe₅S: ecales.—SbMe₃NO₂.—SbMe₅SO₄.

Tetra-methyl-etibonium saite. SbMe, I. S. 30 at 23°. From SbMe, and MeI: eix-sided plates. When distilled the vapour (SbMe, + MeI) takes on the obtained, in eolar with eolar wi

soluble green powder; oxidises rapidly.— 8bMe,NO, [1500]: orystals, v. sol. water, not decomposed by boiling concentrated H.SO,— SbMe,SO,H: soluble plates.—(SbMe,),SO, 5aq.

Di-methyl-stibine sulphides (SbMe.),S, [c. 100°] and (SbMe.),S are formed by passing H.S into an ethercal solution of SbMe, that has been oxidised by exposure to air.

Antimony penta-methide SbMe₅. (o. 98°). Formed together with antimony tetra-methide, (SbMo₄)₂ (c. 90'), by distilling trimethyl stibine iodide with ZnMo. Both are oils which do not fume in air.

Methyl-tri-ethyl-stibonium salts SbMeEt.1. S. 50 at 20°. From SbEt, and Mol. Glassy prisms, sol. alcohol, insol. ether; the solutions are lavorotatory. HgCl. gives a precipitate of SbEt, MeI 1, HgI, -SbMcEt, OH; from moist Ag2O and the iodide, or from the sulphate and baryta. Pps. metallio salts, the hydrates of zino and aluminim dissolving in excess .--SbMeEt, Cl: small needles. - SbMeEt, 1Hgl., -(SbMcEt₃), CO₃: amorphous. — (SbMcEt₄), SO₄: [100°], deliquescent, shining, bitter crystals. (SbMeEt₂).C₂O₄: glassy needles, m. sol. water.—SbMeEt₂C₂O₄H: needles, v. sol. water.—The acetate, formate, and butyrate are orystalline.

Tri-ethyl-stibine SbEt, Mol. w. 209, (159°). S.G. 16: 1 324. V.D. 7 44 (calc. 7 18).

Formation.—1. From SbCl₃ and ZnEt₄.— 2. From EtI and a mixture of sand with an alloy of Sb and potassium. -3. From SbCl, and 1IgEt4. By distilling SbEt 12 with Zn.

Properties.—Oil, smelling of garlic, v. sol. alcohol and other. Takes fire in air; hence it should be kept under water. Decomposes furning hydrochloric acid with evolution of hydrogen: $SbEt_2 + 2HCl = SbEt_3Cl_2 + H_2$ When slowly oxidised it forms SbEt,O and SbEt,(SbO.), Combines directly with S, Se, I, Br, and Cl. Dilute IINO, dissolves it, giving off NO and forming Sblet, (NO₃). In all these reactions tri-ethyl-stibino behaves like a metai. alcoholio solution of ShEt, shaken with HgO liberates Hg while SbEt, O remains in the solution.

Tri-ethyl-stibine-oxide SbEt.O.

Formation .- 1. From SbEt, L and Ag.O. 2. From SbEt₃SO₄ and baryta. -- 3. By slow oxidation of an alcoholic or ethercal solution of SbEt,; SbEt, (SbO.), is formed at the same time, but this differs from SbEt,O in being insol. ether .- 4. By shaking alcoholic SbEt, with HgO.

Properties .- A syrup, v. sol. water and alcohol, m. sol. ether; combines with acids and precipitates metals as hydrates from solutions of their salts. If its aqueous solution is free from SbEt₂(SbO₂)₂, no pp. is produced by H₂S; otherwise a pp. of SbEt₄(SbS₂)₂ is formed. Potassium converts it into SbEt ..

Tri-ethyl-stibine salts.

Chloride.—SbEt,Cl., S.G. 17: 1-540. Oil, soluble in alcohol. Ppd. by adding HCl to an aqueous solution of the iodide or sulphato. Cono. H.SO, decomposes it, giving off HCl.

Oxy-chloride. -SbPt₃Cl₂SbEt₃O. From the oxy-iodide and HgCl₂. Deliquescent solid.

Bromide. — SbEt, Br., S.G. 17: 1.953. Solidifies at -10°. Insol. water, sol. alcohol and ether. Behaves like a metallio bromide.

Iodide.—SbEt.I. [71°]. Formed by heating

Sb with EtI at 140°; or by adding fodine to an alcoholio solution of SbEt, at -15°. Needles (from ether). Unlike the chloride, it is soluble in water. Potassium removes the iodine thus:

SbEt₃l₂+ K₂ = 2KI + SbEt₃. $Oxy \cdot iodide$.—SbEt₃l₂, SbEt₃O. Formed by treating the iodido with NIIa; or by mixing the

iodide with the oxide SbEt, Ö.

Sulphide. -SbEt₃S. From the oxide and H₂S or from SbEt₃ and S. Soluble in water and alcohol. Its aqueous solutions pp. metals as sulphides from their salts.

Nitrate. -SbEt₃(NO₅)₂. [63°]. Formed by dissolving SbEt₃, or its oxide, in dilute HNO₃. Rhombohedra, soluble in water.

Oxy-nitrate. -SbEt,O, IINO, oxy-iodide and AgNO_s.

Sulphate.-SbEt, SO, [100°]. From the sulphide and CuSO. Small prisms, sol. water and alcohol.

O.cy-sulphate.—(SbEt,O),II,SO, the oxy-iodide and Ag.SO. Gummy mass.

Tetra-ethyl-stibonium salts.

Iodide. "SbEt,I, 1, aq (and 'aq). S. (anlydrous): 19 at 20°. From SbEt,, water, and EtI at 100°. Hexagonal prisms, sol. alcohol and other. $-SbEt_4I_{\frac{3}{4}}HgI_2$. $-SbEt_4I_{\frac{3}{4}}HgI_2$.

Hydrate.—SbEt OII. From moist Ag O and the above. Alkaline syrup. Pps. metallic hydrates from salts: stannio oxide and alumina dissolve in excess. Expels NH, from its salts.

Chloride.—SbEt,Cl. Hygroscopic needles.

Forms compounds with HgCl2 and with PtCl4.

Bromide.-ShEt, BrAq: needles.

Periodide.—Sblit,I,. Nitrate.—SbEt, NO.: deliquescent needles. Sulphate.—(SbEt₁)₂SO₁: deliquescent mass. Oxalate.—(SbEt₁)₂C₂O₁.

Antimony-penta-ethide SbEt, (?). (c. 165°). From SbEt, I, and ZnEt,

Tri-isoamyl-stibine Sb(C, II,1)3. From an alloy of Sb with K by C, II, I. Funning liquid, does not take fire in air.

Oxide. Sb(C, II,),O: insoluble resin, soluble in alcohol.

 $Salts. = Sb(C_sH_{11})_3Cl_a \colon \ oil, \ heavier \ than$ water, soluble in alcohol, ppd. by water.— Sb(C,H₁₁)₃Br₂: oil. -- Sb(C,H₁₁)₄I₂: oil. -- $\mathrm{Sb}(\mathrm{C}_{3}\mathrm{H}_{11})_{3}(\mathrm{NO}_{3})_{2}\colon [20^{\circ}]$ slender crystals, insoluble in water, soluble in alcohol.—Sb(C,H11)3SO1: oil.

Antimoay di-isoamyl Sb2(C3H11)4 (?). Formed by distilling Sh(C,H11)3. A heavy oil, soluble in alcohol. Does not fume in air, but explodes in oxygen. Its salts are amorphous.

Tri-phenyl-stibine Sbl'li₃. [48°] (above 360°). S.G. ¹² 1 500. From SbCl₃ (1 pt.), chloro-benzene (1 pt.) and Na; benzeno being used as diluent. Small quantities of SbPh,Cl, and SbPh, Cl2 are also formed. The benzene deposits crystalline SbPh; this is warmed with alcohol containing HCl, which dissolves ShPh,Cl,, and the residual SbPh₃ is converted into SbPhaCla by chlorine. The latter is reduced by alcoholio ammonia and hydrogen sulphide:

 $SbPh_3Cl_2 + H_2S = SbPh_3 + 2HCl + S.$ Properties.—Colourless triolinic tables a:b:o $\alpha = 100^{\circ} 38'.$ $\beta = 103^{\circ} 37'.$ = .697 : 1 : .889. $\gamma = 75^{\circ} 25'$; sl. sol. alcohol, v. e. sol. other, bonzene, glacial HOAc, CS., chloroform, and petro-leum; insol. water and aqueous IICl. It does not decompose HCl; but it combines directly with halogens; it reduces oupric, to cuprous, chloride. With merouric ohloride it reacts as follows: $SbPh_3 + 3HgCl_2 = SbCl_3 + 3HgPhCl_3$

Fuming HNO₃ forms SbPh₃(NO₃)₂.

Salts.-SbPh,Cl2 [143°]: long thin needles; not affected by water; insol. light petrolenm, sl. sol. ether and alcohol, v. sol. benzene and CS_n.

—SbPh₃Br₂. [216°]. — SbPh₃L. [153°]; white
tables. — SbPh₄(OH)... [212°). From the bromide and alcoholic KOII. Amorphous powder, sol. glacial HOAc and reppd. unaltered by water. Insol. ether, v. c. sol. alcohol. Converted by 11Cl, HBr, or HI into haloid salt.—SbPh. (NO.). [156°]. Insol. water, sol. alcohol.

Antimony di-phsnyl chloride Sbl'h, Cl, aq [180°]. Obtained as a by product in preparing SbPh. Needles, insol. water, sol. hot dilute HCl, v. e. sol. alcohol. Alcoholie NII, converts it into Ph.SbO(OII), a white powder, insol. water, ammonia, alcohol, ether, or Na.CO₃Aq, but sol. NaOHAq and glacial

HOAe.

ANTIPYRINE v. Oxy-di-methyl-quinizine.

APHRODÆSCIN. A substance contained in the cotyledons of the horse-chestnut (v. Æscinic acid).

APIÏN. When common parsley (Apinm petroselinum) is extracted with boiling water the filtrato gelatinises on cooling. The jelly is dried at 100° and extracted with alcohol, and the alcohol poured into water. The operation of dissolving in alcohol and ppg. with water is repeated several times, and the apiin finally crystallised from alcohol, with stirring. Apim also occurs in parsley seed; if this is boiled with water, apiol distils over, while apiin separates out from the residue.

Properties. - Needles; sl. sol. cold water, v. sol. hot water, separating again as a jelly; v. sol. alcohol; insol. ether. Its solution in boiling water gives a blood-red colour with FeSO. Gives pieric acid with HNO3; and phloroglucin by potash fusion.

Apigenin C₁₅II₁₀O₅. Boiling dilute II₂SO₄ splits up apiin into apigenin and glucose:

 $C_{21}H_{32}O_{16} + 11.0 = C_{15}H_{16}O_{5} + 2C_{6}H_{12}O_{6}$.

Crystallises in plates (from alcohol). It sublimes near 294°. Sl. sol. hot water, v. sol. alcohol, insol. ether. Potash fusion gives phloroglucin, protoentechnic acid, p-oxybenzoio acid, and oxalic acid.

References .- Rump, Buchner's Repert. f. Pharm. 6, 6; Braeonnot, A. Ch. [3] 9, 250; v. Planta a. Wallace, A. 74, 262; Lindenborn, B. 9, 1123; v. Gerichten, B. 9, 1121; Whitney, Ph. [8] 10, 585.

API0L C, H, O,. [30°]. (c. 300°). Extracted by alcohol from parsley seeds (v. Gerichten, B. 9, 1477). Needles; insol. water. Alcoholic KOII converts it into two crystalline bodies. [54°] and [114°]. 'The essential oil obtained by distilling parsley seeds with water contains apiol but consists chiefly of a terpene, (160°-164°), S.G. 12 865, [a] -30 8°. It has a strong smell of parsley. A small quantity of a hydrochloride, [116°], can be got from it.

References.-Löwig a. Weidmann, P. 46, 53; v. Gerichten, B. 9, 258, 1121, 1477; Pabitzky, Braunschw. Anzeiger, A.D. 1754; Blanchet a. Sell; A. 6, 301; Martins, A. 4, 267; Homollo a.

Joret, J. Ph. [8] 28, 212; and the references under Apiin.

APÓ-. Compounds beginning with this prefix are described under the words to which it is prefixed.

APOCYNIN. The root of Apocynum Cannabium contains amorphous resinous apocynin, sol. alcohol and ether, v. sl. sol. water, and a glneoside, apocynein (Schmiedeberg, Ph. [3]

APOPHYLENIC ACID. The methylohydroxide of einchomeronic acid; v. Prhiding DI-OARBOXYLIC ACID.

APPLES. The artificial essence of apples contains iso-amyl iso-valerate dissolved in

rectified spirit (Hofmann, A. 8I, 87).

APRICOTS. The artificial essence of apricots contains isoamyl butyrate and isoamyl alcohol.

AQUA REGIA v. CHLORHYDRIC ACID. AQUA VITÆ. Alcohol.

ARABIC ACID (Arabin) $C_{12}II_{12}O_{11}$; $nC_{14}II_{12}O_{14}$ (Neuhauer, J.~pr.~62,~193;~71,~255); $C_{12}II_{12}O_{14}$ (Seleibler, B.~6,~612); $C_{89}H_{112}O_{14}$ (O'Sullivan, C. J. 45, 41).

Occurrence. - It is a constituent of probably all lavorotatory gums, and has been isolated from Levantine, Senari, East Indian, Senegal, and Turkey, gum; these gums contain also, as a rule, other acids different from, but closely allied to arabic acid (O'Sullivan). It exists in sugar-beet (Scheibler), and in the extract of yeast obtained by boiling water (Schiltzenberger, BL [2] 21, 204; C. R. 78, 493). The beet gum is probably related to arabic acid, but there is no evidence that the yeast-extract body belongs to the arabin group. Many gum-like constituents. of seeds and roots are referred to as gums, but the great bulk of them obviously hold no relation to arabic acid. It is found in certain animals (Städeler, A. Ph. 111, 26).

Formation.-It is a product of the action of sulphuric acid on alga-mucilage (Brown, Ed. Ph. I. 26, 409); on quince, linseed, and flea-wort mucilages, cellulose being at the same time produced (Kirchner a. Tollens, A. 175, 205); and on metagummic acid (Frémy, C. R. 50, 125). Cellulose is transformed into gum in plants (Mercadante, G. 5, 408). In none of these cases. have we any information as to the character of the gum produced.

Preparation .- The leverotatory gums are principally potassium, magnesium, and calcium salts of arabic or allied acids; they contain from 12 to 18 p.c. water, and yield 2.7 to 3.0 p.o. ash consisting almost wholly of carbonates of these metals. Any one of these gums is dissolved in the least possible quantity of water, the solution is allowed to stand, and, when olear, decanted from any insoluble matter. To the clear liquid twice or thrice as much HCl as is sufficient to convert the bases into chlorides is added, and the gum-acid or acids are precipitated by a moderate excess of alcohol. If the guin contains only arabic acid, the whole pp. can he parified as is described below when dealing with one of the fractions; but if, as is very frequently the case, other allied acids are also present, it is necessary to have recourse to fractional pre-cipitation to isolate the arabic acid. It is found in the fructions least soluble in dilute alcohol, and may be obtained as follows :- The

ourdy pp., produced by excess of alcohol in presence of HCl, is well washed with spirit, and then pressed as free from it as possible. It is redissolved in warm water, care being taken to avoid heating for any length of time, because even the small quantity of HCl retained by the pp. has a tendency on heating even for a short time to decomposo the arabio acid, as will be described below. The solution is cooled, and alcohol gradually added with continual stirring. In this way, the liquid can be made milky without the formation of a pp.; from this 'milk' the acid or acids can be precipitated in successive fractions by the addition of HCl in carefully graduated quantities. If 4 or 5 fractions are obtained, one or more of them is arabic acid. Each fraction is freed from ash by repeated precipitation from aqueous solution with alcohol in the least possible oxcess in presence of HCl, and from IICl by repeated precipitation from concentrated solution by the rapid addition of strong alcohol, whereby the production of a 'milk' is obviated. The fractions thus purified are troated with alcohol (S.G. 81) to render them friable, rubbed down to a powder, filtered out, pressed, and dried over sulphuric acid. Thus prepared they are white, friable bodies, easily soluble in water. If in this state they are exposed for any length of time to a temperature of 100°, they are converted into the meta modifications (meta acids) which are insoluble in water and only swell up to jelly-liko masses whon treated with it. If, however, they are previously dried in a vacuum over sulphuric acid until the weight becomes constant, they can then be dried at 100° without becoming insoluble. The fraction or fractions which are found to have an optical activity, $[a]_i = -26^\circ$ to -28° , and which, when again divided, yield fractions each of which has the same activity, consist of arabic acid.

Properties .- Arabio acid, when slowly dried out of syrupy solutions, on glass plates, is a brittle, transparent, colourless, glassy body, solnble in water. During the drying process, especially if a little mineral acid is present, the acid is frequently converted into the meta modification. Solutions of the body are strongly acid to litmus paper, and have a sharp acid taste; they completely neutralise solutions of the alkalis and alkalino carths, and decompose oarbonates. The salts of the alkaline earths are procipitated out of solution by alcohol; those of the alkalis are not precipitated under the same conditions, but yield peculiar milky or opalescent solutions from which arabic acid, with some of the alkalino salt, is precipitated on the addition of stronger acids. BaSO, PbS, and other sulphides, and some hydrates precipitated in solutions of arabic acid, cannot be filtered out, but pass, in greater part, through the filtor. Fine animal charcoal is carried through in the same way (C.O'S.). Gum arabio provents the precipitation of the alkaloids by phosphonolybdio acid, potassium-mercury iodide, and tannin (Lefort a. Thibault, J. Ph. [5] 6, 169). Theso are properties common to all the gum acids. The defining characters of arabic acid are its optical activity, viz. $[a]_1 = -26^{\circ}$ to -28° , for solutions containing 5 to 6 grams dry substance in 100 c.o., and the composition of its neutral

barium and calcium salts; in the dry state, the former contains 6.0 p.o. BaO and the latter 2.28 p.o. of CaO (O'S.). Solid gum roasted with oxalic acid yields metagummio acid (Frémy), this is dissolved by solutions of the alkalis and alkaline earths with the reproduction of arabic acid (v. Rhem. D. P. J. 216, 539). Gum arabio and tragacanth are rendered insoluble by potassium bichromato and light (Eder. J. pr. 19, 299). Gum, oven in small quantities, injected into the blood diminishes the climination of urine, largo doses completely stop the secretion, with a marked increase of blood pressure (Richet a. Montard Martin, C. R. 90, 88).

Reactions .- 1. Heated with moderately strong nitric acid, arabic acid yields mucic (v. Kiliani, B. 15, 34), saccharic, oxalic, and tartario (Liebig) acids; with fuming nitric acid it yields substitution products.-2. Gum heated in scaled tubes with broming yields a colourless or yellowish liquid, probably C12H211O10Br,, which, when treated with silver oxido, lead oxide, or caustic soda, is converted into isodiglycolethylenio acid, C₁₂H₁₀O₁₂ (Barth a. Hlasiwetz, A. Ch. Pharm, 122, 96). It is possible some of the decomposition products of arabic acid would yield the same results. 3. Gum arabic or arabin when heated to 150° with 2 parts acetic anhydride yields tetracetyl-diarabin (?) $C_{12}H_{16}(C_2H_3O)_1O_{16}$; and, when heated to 180 with 6 to 8 parts of the anhydride, a body having the composition C12H13(C2H3O)3O10; these acetyl derivatives are white amorphous powders (Schützenberger a. Naudin, A. Ch. [4] 21, 235). These bodies are certainly not derived from arabic acid as a whole, but from some one of its decomposition products (C.O'S.) .- 4. Pepsin, in dilute HCl solution, acts on dextrorotatory gum acid, arabinose being amongst the products; pancreatin has no action (Fudakowski, B. 11, 1072). 5. (a) Gum arabic left for some time in contact with sulphuric acid is converted into dextrin (!) and, on boiling, yields a sugar probably identical with galactosa (Berthelot, C. O. 21, 219). (b) Strong sulphuric acid converts a strong solution of gum in a few hours into metagummic acid; but gum arabic freed from lime by oxalic acid is not transformed in the same way (Frémy). (c) Pulverised gum arabic, triturated with strong sulphuric acid, yields sulphoguminic acid and a peculiar gum resembling that produced from linen by the action of sulphuric acid (Braconnot) and not capable of fermenting with yeast (Guérin-Varry). (d) Arabic acid, digested with dilute sulphuric acid, yields a crystallisable sugar, a non-crystallisable one, and an acid the barium salt of which is insoluble in alcohol: gums from different sources yield these bodies in various proportions, somo varieties yielding searcely any of the crystallisable sugar (Scheibler, B. 6, 612). • (e) A solution containing 30 grams arabio acid (pure), 100 o.o. water, and 2 grams sulphuric acid, yields, on digestion at 100° for 15 minntes, at least two sugars and a new soid the Ba salt of which is insoluble in alcohol: the following equations ropresent the change :-

| I. C₉₀H₁₄₂O₇₄ + H₂O = C₈₅H₁₂₂O₈₆ + C₈H₁₂O₈₆ arabin cold | A-arabin cold

Continued digestion produces changes that may be represented as follows:—

III. IV. V. VI. VII. and VIII. $C_{77}H_{122}O_{64} + 9H_2O = C_{41}H_{68}O_{57} + 6C_{5}H_{12}O_{6} \\ -6-\text{rabinosio acid}, \beta_{77} + 6C_{5}H_{12}O_{6} \\ -6-\text{rabinosio acid}, \beta_{77} + 6-\text{rabinose}. \\ On further digestion we get: \\ IX. C_{41}H_{68}O_{37} + H_2O = C_{35}H_{36}O_{37} + C_{4}H_{12}O_{6} \\ -6-\text{rabinosic acid} & 3-\text{rabinose} (?) \\ X. C_{23}H_{36}O_{27} + H_{2}O = C_{23}H_{36}O_{27} + C_{6}H_{12}O_{6} \\ -6-\text{rabinosic acid} & 3-\text{rabinose} (?) \\ XI. C_{29}H_{18}O_{27} + H_{2}O = C_{23}H_{38}O_{27} + C_{6}H_{12}O_{6} \\ -6-\text{rabinosic acid} & 3-\text{rabinose} (?) \\ \end{cases}$

This last acid is very stable, resisting the action of a boiling 3 to 4 p.c. solution of sulphuric acid for a considerable time (O'Sullivan, C. J. 45, 41). a-arabinose is not yet fully described; β-arabinose' is Scheibler's arabinose, y-arabinose is probably identical with galactose, and the remaining sugar or sugars are imper-fectly described. These reactions convey some idea of the constitution of the gums of tho arabin group, i.e. of those which are salts of a gum-acid with alkalino or alkaline-cartly bases, and enable us to understand some of the differonces observed in their properties. Those bodies vary considerably in optical activity; this is due (a) to the varying proportions of different closely related acids they contain. O'Sullivan found the acid of some samples of gum arabic to consist almost wholly of arabic acid, whilst others contained also a arabinosic acid, $[a]_j = -36^{\circ}$, and an acid $(C_{05}H_{152}O_{79})$, $[a]_j = -23^{\circ}$, with a C.H. O. group more than arabic acid. (b) To the character and position of the C.H.10O. group in the acid from the C23H18O22 body upwards. Kiliani (l.c.) shows that different varieties of gum yield, when oxidised with HNO, proportions of mucic acid varying between 143 and 383 p.c., thus indicating a difference in the number of galactose (γ-arabinose)-yielding groups they contain (see also Scheibler). And (c) possibly to the structure of the C₂₃H₃₈O₂₂ group itself. The guins, too, vary much in the character of tho solutions they yield (Gm. 15, 194); some give a thin syrupy solution, others a thick and jellylike one; this is due to the varying proportion of the acid naturally converted into the meta modification-the gums which yield the thinnest solutions are those which contain the greatest amount of ash. Gums from the same source have not always the same optical activity; Scheibler found the beet gum of one season +, and of another -; Kiliani states that East Indian gum, and gum arabic elect. are +, the samples of these gums examined by O'Sullivan were -; the sample of Australian gum examined by the former was +, that by the latter, inactive. From this it would appear that the same plant does not produce the same gum in every season, but, on the whole, it is fairly certain that the acids of all the guns are constituted in the same way as arabic acid described above.

Combinations.—The arabic salts of the alkaiine earths are prepared by exactly neutralising moderately strong solutions of the pure acid with clear solutions of the earth the salt of which is required, precipitating with alcohol,

treating the pp, with strong alcohol until it admits of being powdered, collecting the powder on a filter, pressing, and drying over sulphurio acid. The Ba salt contains C_{s} . $H_{1,2}O_{s}$,BaO (6.0 p.c. BaO) and the CaO salt, C_{s} , $H_{1,2}O_{s}$,CaO (2.28 p.c. CaO). When the syrupy solutions of these salts are allowed to dry spontaneously they yield glassy, transparent masses, like natural gums: the salts of the arabinosic acids yield similar boliles. C. O'S.

ARABITE C.H.O. which is probably CH₂(OH).CH(OH).C

ARABONIC ACID v. Trtra-oxy-valeric acid. ARACHIC ACID C₂₆H₄₆O₂. Mol. w. 312. [75·5°]. Arachidic acid.

Occurrence.—1. In the oil from the ground nut Arachis hypogoza, from which it is obtained after saponification (Gössmann, A. 89, 1).—2. In butter (Heintz, P. 90, 146).—3. As glycerylether in the fruit of Nephelium lappaceum (Oudemans, J. pr. 99, 407).

Formation.—1. By potash-fusion from brassidic acid (Goldschmiedt, J. 1877, 728).—2. From stearic acid by converting it into the corresponding alcohol, C., II., Cll_OH, and iodide C., H., CH_1. The latter gives octadecyl aceto-acotic ether when treated with sodio-aceto-acetic ether, and alcoholic KOII thence produces arachic acid (Schweizer, Ar. Ph. [3] 22, 753).

Properties.—Small shining plates; v. sol. boiling alcohol, and ether; insol. water. Gives a nitro-, [70°], and an amido- [59°], arachic soid.

Salts.—(Scheven a. Gössmann. A. 97, 257). AgA': prisms (from alcohol)—CuA';: needles (from alcohol)—BaA';: hardly soluble in alcohol.—SrA',—MgA';: crystalline powder (from alcohol).—KA': usually gelatinous.

Methyl ether. [55°]. Scales (Caldwell, A. 101, 97); [53°] (Schweizer).

Ethyl ether. [49.5]. (296°) at 100 mm. Iso-amyl ether. [45]. Scales. Arachins. Prepared artificially by heating

Arachins. Prepared artificially by heating arachic acid with glycerin (Berthelot, A. Ch. [3] 47, 355); they are insol. ether. Di-arachin C.H. (OlH)(OC.,H.,O), melts at 775°].

 $C_3H_3(OH)(OC_{20}H_{30}O)_2$ melts at [75°]. Arachyl chloride $C_{20}H_3O.Cl.$ [67°] Unstable (Tassinari, B. 11, 2031).

Arachamide C₂₂H₃₉O.NH₂. [99°]. Prisms grouped in stars; insol. water, sol. hot alcohol (Gössmann a. Scheven, A. 97, 262).

Acetic-arachic anhydride C.,H.,O.O.Ao. [60°]. From AcCl and potassium arachato. Scales (from ether) (T.).

Valeric-arachic anhydride

C_xH_{xy}O.O.C.II,O [68°] (T.).

ARALEÏN. The hark of Aralia spinosa contains a tannin which gives a green colour with Fe_xCl_x and a glucoside called aralein. The latter is a neutral, light-yellow substance, sol. water and alcehol, insol. ether, henzene, and CHCl_x. Gives no pp. with lead acetate, HgCl_x or PtCl_x. Boiling dilnte HGl converts it into white araliating insol. water (Holden, Ph. [3] I1, 210).

According to Kiliani (B. 20, 339a, 1933) the formula of arabinose is $O_aH_{i,i}O_a$; should this prove to be so, the formula of arabin seid and of the arabinosis acids must be diminished by a CH₂O group for every molecule of arabinose-yielding group they contain.

10, 199). Baup (J. pr. 55, 83) finds four orystalline substances in it: Amyrin [174°]; Breidin, S. 39 at 20°; Brein [187°]; and Bryoidin [185°].

ABBUTIN C₁₂H₁₆O, §aq. [166°].

Occurrence.—Together with methyl-arbutin in the leaves of the red bearberry, Arctostaphylos Uva Ursi (Kawalier, A. 82, 241; 84, 356), and in the leaves of a species of winter-green, Pyrola umbellata (Zwenger a. Himmelmann, A. 129, 203).

Preparation.-The aqueous infusion of the leaves is ppd. by lead acetate, excess of lead romoved by H.S, the filtrate evaporated and the arbutin extracted and crystallised by a mixture of ether (8 pts.) and alcohol (1 pt.). The product is a mixture of arbutin and methylarbutin, which can be separated by orystallisation from water (H. Schiff, G. 11, 99; 13, 538;

A. 221, 365; cf. Habermann, M. 4, 753). Properties.—Long glistening needles which melt at 165°, but, on second fusion, at 187° (H. Schiff, B. 11, 304; A. 206, 159). V. sol. aleohol and boiling water, v. sl. sol. ether. Its aqueous solution gives no pp. with lead acetato or subacetate. Does not reduce alkaline cupric solution. Dilute Fe Cl, gives a blue colour.

Reactions.-1. Split up by emulsin or by boiling dilute II SO, into hydroquinone and glucose (Strecker, A. 107, 229).-2. H.SO, and MnO, form quinone .- 3. Converted by Ag.O into water

and diarbutin, an extremely soluble syrup, whence arbutin can be recovered by reduction with Zn and H SO₁ (Schiff, A. 154, 244).-4. Chlorine passed into an aqueous solution forms di-, and tri-chloro-quinones (Strecker, A. 118, 295).

Acctyl derivative C.H.Ac.O. or needles (from alcohol), insol. water.

Benzoyl derivative C.H.Bz.O. Crystallino powder, sl. sol. alcohol.

Di-nitro-arbutin C12H14(NO2)2O2. Golden needles (from water); insol. ether (Hlasiwetz, a. Habermann, A. 177, 313). Gives an orange pp. with lead subacetate. Boiling dilute H.SO gives di-nitro-hydroquinone. Converted by Ac.O into C₁₂H₄Ac₂(NO₂)₂O₁.

Methyl arbutin C₁₂H₁₂O₂ i.e.

 $C_eH_{2}O(OH)_{1}O.C_{0}H_{2}OMe.[169^{\circ}]$ (Michael); [175°] (Schiff). Occurs in nature associated with arbutin. Formed synthetically by the action of acetochlorhydrosc upon potassium hydroquinone methyl ether, KO.C.H.OMe (Michael, Am. 5, 178; B. 14, 2097). Also from crude arbutin by converting the free arbutin into methyl-arbutin (by Mcl and KOH) or into benzyl-arbutin (H. Schiff, G. 12, 464; A. 221, 366).

Properties .- Colourless silky needles, with bitter taste. Contain ag and melt at 169° (Michael); contain aq and melt at 175° (Schift). Sol. water and alcohol, v. sl. sol. ether. Gives no blue colour with Fe,Cl,

Benzyl-arbutin C₁₀II₂₂O₇ i.e. PhCH_O.C₆II₁O.C₆II₁₁O₈ aq. [161°]. S. ·19 at 23°. From PhCIL Br, commercial arbutin (containing methyl-arbutin), and KOH in boiling alcohol. Excess of KOH is removed by CO2, and after evaporating and adding water, benzyl-arbutin

ARBOL ARBOL ARBOL. The product of a lig ppd. while methyl-arbutiu remains in solution, tree (Canarium album) growing in the Philippine Arborescent needles. Soluble in boiling water, Islands (Maujean, J. Ph. 9, 45; Bonastre, J. Ph. very soluble in alcohol. Does not reduce Fehling's solution, except after short boiling with H.SO, which splits it up into glucose and benzyl-hydroquinono. Gives on nitration yellow needles which are benzyl-nitro-arbutin, C₁₉H₂₁(NO₂)O₇, [143°], split up by dilute H₂SO₄ into glucose and benzyl-nitro-hydroquinone (Schiff a. Pellizzari, A. 221, 355).

Isoamyl-arbutin. From the mixture of arbutin and methyl-arbutin by amyl bromide and NaOH (S. a. P.). Necdles. On decomposition gives iso-amyl-hydroquinone and glucose.

ARCHIL or Orscille is a purple dyo obtained from various lichens (Roccella, Lecanora, and Variolaria) containing acids (erythric, lecanoric, &c.), which on decomposition yield ore in (q. v.) which is converted by air and ammonia into red orcein. Whon K.CO. or Na.CO. as well as ammonia is added to the lichens litmus is produced.

ARGININE C.H., N.O. Easily soluble in water; reacts alkaline. Occurs to the extent of about 3-4 p.c. in the young shoots of the lupine (lupinus lulcus). The shoots are extracted with water; tannin and lead acctate are added to the extract: the filtrate is acidified with H.SO4, again filtered, and ppd, with phosphomolybdic acid; the pp. is washed and treated with cold milk of lime, and the solution of the base finally neutralised with HNO, and evaporated to crystallisation.

Salts.-B'HNO, laq: slender white soluble needles; with phosphomolybdic acid it gives a white pp. soluble in hot water; with pieric acid a yellow crystalline pp. is formed on standing. B'HCl: large crystals. - B',Cu(NO,), 3aq: formed by heating a solution of the nitrate with cupric hydrate; dark-blue prisms, sl. sol. cold water (Schulze a. Steiger, B. 19, 1177).

ARGOL. Crude acid potassinm tartrate deposited from wine.

ARGYRÆSCIN C2,1142O12. A glucoside in the cotyledons of the horse-chestnut. Minute tables (from dilute alcohol). Split up by diluto HCl into argyrescetin C. Ha,O, and glucose. Potash produces propionic acid and ascinic acid (q. v.) (Rochleder, J. pr. 87, 1; 101, 415).

ARIBINE C. H. N., S. 12.9 at 23°. A base extracted by dilute H.SO₄ from the bark of Arariba rubra, the solution being treated with lead acetate and the base dissolved in other, from which it separates as anhydrous pyramids or (with 8 aq) as four-sided prisms. V. sol. water and alcohol, m. sol. ether (Rieth a. Wöhler, A. 120, 217). Salts. - B" 211Cl. - B"H.PtCl. -B"H.SO., -B" 21L.SO.,

ARICINE C₂₂H₂₆N₂O₄. [188°]. S. (ether) 5 at 18°. 'Cusconine,' Cinchovatine. Yellow Cusco bark contains '24 p.c. aricine and '37 p.o. ensconine. Occurs also in bark of cinchona cuprata (Hesse, Ph. [3] 12, 517). Prisms (from dilute alcohol); insol. water, v. e. sol. chloroform, m. sol. ether, v. sl. sol. alcohol. Solutions are not fluorescent. Leevorotatory in alcoholio or ethereal solutions; its solution in dilute HCl is inactive. In a 1 p.c. othereal solution $[a]^{\text{b}} = -94^{\circ}8^{\circ}$; in a 1 p.c. alooholic (97 p.e.) solution $[a]_{\text{b}} = -54^{\circ}$. Aricino is turned dark green by cono. HNO₃. Bleaching powder and NH₃ only give a yellowish colour.

Salts. — B'HC'saq. — B'HRPHR faq. — 3 HI. — B'HNO. — B'H-SO.: slender needles, n. sol. cold water.—B'H-SO.: small prisme, v. sl. col. cold water.—B'HOAc Saq: graine, v. sl. ol. cold water.—B'HOAc Saq: graine, v. sl. ol. cold water.—B'H-C₂O. 2aq: white prisms, wickly changing to rhombohedra, S. 049, sl. ol. hot alcohol. — B'HSCy. — Salioylate VC,H₂O. 2aq.

References.—Pelletisr, A. Ch. [2] 42, 330; 1, 185; Pelletier a. Corriol, J. Ph. [2] 15, 575; everköhn, Repert. f. Pharm. 33, 357; Manzini, Ph. [3] 2, 95; Howard, Ph. [3] 5, 908; Hesse, 166, 259; 181, 58; 185, 321; 200, 303.

ARNICIN C₂₀H₂₀O₄ (?). An amorphous subsance present in the root, leaves, and blossoms ! Arnica montana (Walz, N. Jahrb. Pharm. 3, 175; 14, 79; 15, 329).

AROMATIC SERIES. Substances whose molecules contain a benzenc nucleus are said to belong to the aromatic series.

Elements attached to a carbon atom belonging to the benzene nucleus are more firnly fixed than when attached to a carbon atom not in that nucleus: in the former case the derivative (called an eso derivative) has the character of a derivative of benzene, in the latter case the derivative (called an eso derivative) behaves like a fatty compound. Thus exo-chloro-toluene (benzyl chloride) C_oH_oCH_oCH_oCl behaves like ethyl chloride, being readily converted into an alcohol, amine, or cyanide, by treatment with KOH, NII, or KCN, respectively; while eso-chloro-toluene, C_oH_oCl.Ch_o, is not affected by these reagents.

It must, however, be added that the ease with which a given atom or radicle in the molecule of an aromatic compound may be displaced depends not only upon its position in relation to the carbon atoms but also upon the existence and position of other elements or radicles in the molecule. Thus o- and p-, but not m-, chloro-itro-benzene are converted by hot aqueous totash into nitro-phenols, and by NII₃ into itro-anilines; while chloro-phenols and chloro-enzene sulphonic acids are converted into discrepenzene by potash-fusion.

Halogens acting upon cold hydrocarbons in he presence of carriers (such as I) enter the enzeno nucleus, but when acting alone npon ydrocarbons at 100° or upwards they enter a ide chain (exo position). Direct sunlight has the same effect as elevation of temperature, but its effect is entirely counteracted by the presence of iodine (Schramm, B. 18, 606). Halogens attack a benzene nucleus that already contains hydroxyl, amidogen, or SO₂H, with much greater vigour than when its carbon atoms are united only to hydrogen and carbon.

Cone. HNO, and cone. II SO, attack aromatic compounds, NO, and SO, H displacing H in the nucleus; they do not act upon fatty occupounds in this way.

The constitution of the molecule of benzene, and the methods by which the relative position of elements or radicles in the molecules of its lerivatives have been determined will be discussed in another article, v. Benzene.

Laws of Substitution.—I. When one of the ollowing radicles has displaced one of the atoms if hydrogen in the benzene molecule, forming he compound C₂H₂A, any new group on enter-

ing will take up a position meta to A. Here A may be CO.H. SO.H., or NO., and probably also CN, CHO, SO.Ph, and CO.CH., (Hubner, B. 8, 875; Nolting, B. 9, 1797; cf. Armstrong, C. J. 51, 259; Morley, C. J. 51, 579).

II. If in a substituted benzene, C₄H₃B, the eubstituting element or radicle be not one of the "g, then any new group on entering will take up an ortho or a para position: usually chiefly p with a little c. Examplee of B are NH₂, NiIAc, OH, Cl, Br, I, CH₃ and all chains of carbon atoms except such as begin with CO.

These laws tell the chief product of the substitute of isomerides at variance with.

The radicles that have meta substitution are all composed of an element (N, C, or S) united to a chlorous group; CCl₃ is also a radicle of this kind, and it gives a m-nitro-derivative, but it also gives a p-chloro-derivative.

The radieles that induce p or c substitution are either single elements, or elements united to basylous elements or groups. The radieles CH_Cl and CHCl₂ are intermediate in character; the latter appearing to resemble CCl₃, the former resembling Cll₃. Armstrong points out that the radieles producing m derivatives are unsaturated, and might form additive compounds before substitution takes place.

Amido compounds in presence of excess of H.SO, (20 pts.) when treated with the calculated quantity of HNO, dissolved in H.SO,, added at 0°, give chiefly meta-nitro derivatives, some of the p-nitro derivative being also formed. Examples: aniline, acctanilide, toluidine, xylidine, p-bromo-anilino (Nölting a. Collin, B. 17, 261), di methyl-aniline, di-ethyl-aniline (Groll, L. 19, 198), ethyl-aniline (Nölting a. Stricker, B. 19. The amount of meta-nitro derivative formed is increased by increasing the quantity of sulphuric acid present. In all these cases nitrogen is attached to a chlorous radicle, anilino sulpliate being Call, N(O.SOall) II, and might therefore be expected to produce a meta derivative. It is, however, curious that a solution of aniline sulphate in a small quantity of sulphuric

acid gives very little m-nitraniline on nitration.

When a new element or radicle enters a benzenc nucleus in which more than one H is already displaced if it can satisfy the requirements of each of the substituents already present it will do so; if not it obeys the most powerful substituent precent. The following appears to be the order of priority, beginning with the strongest:

HO; NH₂; halogens; CH₃; otheralkyls; NO₄; CO₂H and SO₂H. T.ac conversion of p-nitrophenol into C₂H₂(OH)Br(NO₂) [1:2:4], and that of o-nitro-phenol into C₂H₂(OH)Br(NO₂) [1:5:2], hy the action of Br are instances where both NO₂ and OH are obeyed; but in the action of Br upon p-brono-phenol and of HNO₃ upon C₂H₂(OH)Cl₂ [1:2:4] the new substituent obeys the stronger radicle, producing C₃H₂(OH)Br₃ [1:2:4:6] and C₄H₂(OH)Cl₂(NO₂) [1:2:4:6] respectively.

When a hydrocarbon radicle is introduced by the agency of AlCl, it does not always follow the foregoing rule; thus m-xylene is the chief product of the action of McCl upon beuzene impresence of AlCl.

Differences between o, m, and p compounds. Ortho, meta, and para compounds usually boil at about the same temperature, but the para compounds have the highest melting-points. The ortho compounde are usually the most, and the para compounds the least, volatile with eteam. In the oxidation of ortho compounds the benzene ring is liable to be broken up, while in the meta and para compounds this is not the case (v. Hydrocarbons). Ortho compounds readily give rise to products of condensation in which the side chains may be supposed to be joined in the form of a ring; this tendency is observed to some extent in the para series but not at all in the meta series. Thus by loss of H₂O o-amido-phenyl-glyoxylic acid gives isatin; o-amido-cimamic acid gives carbostyril, o-oxycinnamic acid gives coumarin; phthalic acid gives phthalic anhydride.

o-nitraniline hydrochloride is readily decomposed by water, p-nitraniline hydrochloride less so, and m-nitraniline hydrochloride is hardly decomposed by water (Lellmann, B. 17, 2719). In general, the introduction of a radicle into the m-position produces less change in the properties of a compound than the introduction of the same radicle into the o and p position, and of the resulting derivatives the meta are the most stable. Thus m-xylene is oxidised with difficulty, while dilute HNO, readily converts o-, and p-, xyleuc into toluic acids. Ortho-, and para-, oxybenzoic acids are converted into phenol by heating with aqueous HCl in sealed tubes. while m-oxybenzoic acid, like benzoic acid itself, is unaffected. Ammonia converts o- and p- nitroanisols into nitranilines, but does not affect mnitro-anisol or anisol itself; similarly ammonia converts o-, and p-, bromo-nitro-benzenes into nitranilines, but does not affect m-bromo-nitrobenzene or bromo-benzene itself. Boiling alkalis convert o-, and p-, nitraniline into nitrophenols, but do not affect m-nitraniline or aniline. Anilino and m-nitraniline resemble one another in readily uniting with phenyl thio-carbimide (forming diphenyl-thio-urea and nitro-di-phenyl-thiourea respectively), while p-, and o-, nitraniline require to be heated for some time with phenyl thio-carbimide before they will combine. Benzoic acid and m-oxy-benzoic acid are readily reduced by sodium-amalgam to benzyl alcohol and m-oxy-benzyl alcohol respectively, while o-, and p-, oxybenzoic acids are not attacked. Orthoand para-nitro-acetanilide dissolve in conc. potash; the former is readily saponified by the potash, giving potassium acetate and o-nitroaniline; a similar decomposition occurs with the latter, but with greater difficulty. Mcta-nitroacetanilide is insolublo in conc. potash, and is scarcely affected by it (Kleemann, B. 19, 336). The substitution of an atom of hydrogen in the nucleus by an atom of bromine is accompanied by absorption of heat; in an actual experiment this is not observed, being more than counterbalanced by the heat developed in the eimultaneous formation of HBr (Werner, Bl. 46, 282).

Occasional reactions.

1. Acetyl bromide not only displaces H by Ac but sometimes even turns ont an alkyl; thus it converts di-methyl-aniline and di-ethyl-aniline

into methyl acetanilide, and ethyl-acetanilide respectively (Staedel, B. 19, 1947).—2. Bensoyl chloride sometimee behavee in the same way, converting di-methyl-aniline and di-ethyl-aniline into mothyl-benzanilide and ethyl-benzanilide respectively.—3. Nitric acid in nitration sometimos turns out acetyl, converting ethyl acetanilide into $C_0H_3(NO_y)_2NEtH$ [1:3:4], and behaving similarly towards methyl acetanilide (Norton, B. 18, 1997). Nitric acid somotimes turns out bromine; thus it converts p-bromoanilino into tri-nitro-aniline (pieramide) (Hager, B. 18, 2578).—4. Potash converts di-nitro-di-mothyl-anilino, $C_0H_3(NO_y)_2NMc_y$ [4:2:1] into di-nitro-phenol, di-methyl-amine being given off.

Molecular changes.

At high temperatures o-compounde may change to p-, and both o- and p- to m-. At 100° o-phenol-sulphonic acid changes to p-phonol-sulphonic acid. At 220° potassium salicylate changes to p-oxybenzoate, while sodium salicylate is not affected at that temperature. Resorcin is obtained by potash-fusion from benzeno p-disulphonic acid, p-chloro-benzene sulphonic acid, and o- and p-bromo-pbenol.

Methyl can pass from combination with nitrogen into the nucleus; thus dimethylanilino methylo-iodide at 220° gives o- and p. dimethyl toluidino, methyl-xylidine, and di-methyl-xylidine; while at 335° it gives ψ-cumidine, C_oH₁Me₂NH₂ (Hofmann, Pr. 21, 47).

Ethyl aniline hydrochloride at 320° changes similarly to ethyl-phenyl-annine, C.H.EthH., while isoamyl-aniline hydrochloride becomes isoamyl-phenyl-amine, C.H.(C.H.,)NH. (Hofmann, B. 7, 526). In these cases it may be supposed that Mel, MeCl, EtCl, and C.H., Cl respectively are split off and then attack the nucleus; this action of MeCl is seen in the conversion of xylidine hydrochloride into \(\psi\$-cumidine by heating with methyl alcohol at 280° (Hofmann, B. 13, 1730).

Conversion of fatty compounds into aromatic.

1. By passing acetylenc (q. v.) through a red hot tube.-2. By heating acctone or allylene with sulphuric acid mesitylene is formed. -3. Uvitic acid, Coll. Mc(CO.H) [1:3:5] is formed by boiling pyruvic acid with baryta.-4. Oxy-uvitic ether is formed by the action of chloroform on sodium aceto-acetic ether. - 5. Succinylo-succinic ether from succinyl chloride, sodium, and succinic ether is di-oxy-di-hydro-terephthalic ether; when heated with KOH it gives hydroquinone. Hydroquinone is also formed when succinates are subjected to dry distillation .- 6. Phloroglucin tri-carboxylic ether is formed by the action of sodium or of ZnEt, on malonio cther (Baeyer, B. 18, 3457; Lang, B. 19, 2937). 7. Tri-mesic ether is formed by the action of sodium on a mixture of formic and acetio ether (Piutti, B. 20, 537).—8. Hexyl iodide and bromine at 200° gives hexa-bromo-benzene (Krafft, B. 9, 1085; 10, 801).—9. K and CO combine, forming C₆(OK)₆ (Nietzki a. Benckiser, B. 18, 1833).

Conversion of aromatic compounds into fatty.

Carbonic, oxalic, and formic acids are products of oxidation of aromatic compounds.

2. Benzene is converted by KOlO, and HOl into C.H.Cl.O., which is converted by baryta into fumario acid.—3. Nitrous acid converts pyrocatechin in ethereal solution into dioxytartaric scid.—4. HCl and KClO, convert gallic acid into tri-chloro-glyceric acid.—5. Chlorine passed into a cold aquecus solution of phloroglucin gives dichloro-acetic acid (Hlasiwetz a. Habermann, A. 155, 132).

Aromatic acids. The principal aromatic acids are those containing SO₃H and those containing CO₂H. The former will be discussed as SUL-PHONIC ACIDS, the latter will be briefly characterised here (v. also ACIDS, AMIDO., BROMO., OHLORO., IODO., and NITRO-acids).

Formation .- 1. By oxidation of homologues of benzene or derivatives of such homologues. K2Cr2O, (2 pts.), H.SO, (3 pts.), and water (3 to 5 pts.), is a convenient mixture for the purpose; but it converts all side chains into carboxyls, thus m- and p- xylene become iso-, and tere-, phthalie acids, while mesitylone becomes trimeiic acid. Dilute HNO, (S.G. 1.2) and aqueous KMnO, oxidiso more gradually, attacking one sido chain at a time, thus converting o- and ptylene into o- and p- toluic acid and mesityleno nto mesitylenic acid .- 2. By passing CO, over mixture of a bromo derivative and sodium: $J_0H_0Br + CO_2 + Na_2 = C_0H_0CO_2Na + NaBr(Kekulé,$ 4. 137, 178). -3. By heating a bromo-derivaive with chloroformic ether and sodium, thus: C₈H₈Br + ClCO₂Et + Na₂ = C₆H₈CO₂Et + NaCl + VaBr (Wurtz, A. Suppl. 7, 125). - 4. By the etion of COCl, or CO, on aromatic hydrocarbons a presence of AlCl, (Friedel a Crafts, v. Alu-INIUM CILORIDE, p. 147) the product being reated with water. The amides may be prouced in a similar way by using Cl.CO.NH, istead of COCl. (Gattermann a. Schmidt, B. 20, 58).-5. By heating sulphonates with sodium fornate: $PhSO_3K + HCO_2Na = Ph.CO_2Na + HSO_3K$ V. Meyer, A. 156, 273). - 6. By saponification of itriles. The nitriles may be obtained either by eating sulphonates or exo-chloro derivatives ith potassium cyanide or by heating thio-carimides with copper (Weith, B. 6, 212). Nitriles ay also be obtained by distilling the formyl erivatives of amines with zine-dust, c.g.: hNH.CHO = H₂O + PhCN.—7. By exidation of cohols or aldehydes .- 8. Aromatic acids couining carboxyl in the side chain can be preared by synthesis with aid of aceto-acetic etlier .v.) or of malonic ether (q.v.).—9. Oxy-acids are formed by boiling diazo-acids with water, or by potash-fusion from chloro-, bromo-, iodo-, or sulpho-, acids.—10. By passing CO, into sodium phonols at 180° or potassium phenol; in the former case CO.H takes up a position orthe to the hydroxyl. Potassium phenol at 140° gives malicylic acid, but at 170°-200° it gives p.oxy-ben-

poic acid. The reaction takes place in two stages: C_aH_3 , $O.Na + CO_2 = C_aH_3$, $O.CO_2Na$; C_aH_3 , $O.CO_2Na = C_aH_3$ (OH), CO_2Na

v. OXY-BENZOIC ACIDS).—11. By heating phonols with dilute alcoholic solution of CCl₄ and NaOH at 100°: C₂H₄OH + CCl₄+6NaOH = Z₄H₄(ONa)CO₂Na+4NaCl+4H₂O (Tiemenn a. deimer, B. 9, 1285). The carboxyl takes up exitions pars and ortho to the hydroxyl.—

12. Perkin's synthesis of cinnamic acid and its hemologues is described and discussed in the article on ALDEHYDES.—13. Resorcin and its homologues are converted into (1, 3, 4) and 1, 3, 2) di-oxy-benzoic acids and their homologues by heating with ammonium carbonate and water; while hydroquinone and its homologues heated with potassium bicarbonate, water, and a little K₂SO₃ give (1, 4, 2) di-oxy-benzoic acid and its homologues (Senhofer, Sitz. B. 80, 504; 81, 430, 1044; M. 2, 448).

Reactions.—1. The aromatic acids are subject to the general laws governing substitution in the benzene nucleus.—2. They are usually sl. sol. water but v. sol. alcohol and ether. The homologues of benzoic, and of salicylic, acid are volatile with steam, m., and p., axy-benzoic acids are not volatile with steam. Salicylic acid and its homologues are soluble in chloroform, p-oxy-benzoic acid and its homologues are not. Ortho-oxy-acids are also characterised by giving a violet colouration with Fo₂Cl₁.—3. Ortho-oxy-acids of the form C₀H₁(OH).CH₂CO₂H or C₀H₁(OH).CH₂CI₂Cl₂H. have a tendency to produce anhydrides or lactones; ortho-amido acids of the form C₀H₁(NH₂).CH₂CO₂H or C₀H₁(NH₂).CH₂CO₂H. readily form anhydrides, similarly called lactams: C₀H₁(NH₂)CH₂CO₂H.

or lactims: $C_kH_k < \frac{CH_x}{N} > C.OH.-4$. Benzene is produced by fusion with NaOH from benzoio acid (75 p.c.), trimellitic acid, hydrocinnamic acid, and cinnamic acid (50 p.c.); a little diphenyl is also formed. Fusion with NaOH converts o_x , and p_x , oxy-benzoic acids into phenol (50 to 60 p.c.); protecatechuic acid into resorcin (50 to 60 p.c.); $\{1,3,5\}$ -di-oxy-benzoic acid into resorcin (60 p.c.), phloretic acid and p-counaric acid into p-oxy-benzoic acid and finally into phenol; oxy-benzoic acid, and finally into phenol (Barth a. Schreder, B. 12, 1255).

Aromatic bases. The preparation and preperties of the aromatic bases have been discussed in the article on Amers. They may be divided into two classes according as the nitrogen is attached to carbon in a benzene nucleus or in a side chain; bases of the latter form resemble fatty amines. Amines containing audiogen attached to the beazene nucleus ars weakened in basic power by introduction of nitroxyl or halogens into the nucleus, more especially if these radicles do not occupy a position meta to the amidogen. Trichloraniline, dinitraniline, and trinitraniline do not combine with acids; the lefer is even saponified by potash with formation of trinitrophenol.

ARSENATES. Salts of arsenic acid, v. Arsenic, Acids of, p. 305.

ARSENIC. As (Arsenicum, Regulus arsenici, λροενικών. By the term σανδαράκη Aristotle seems to mean a compound of arsenic and sulphur, called άρλενικόν by Theophrastus). At. w. 74-9, Mol. w. 299-6; 149-8 at c. 1700° (Biltz a. Meyer, B. 22, 725). Melts only under great pressnre (Landolt; also Mallet, C. N. 26, 97). S.G. 5-23 to 5-76; pure, crystalline 16/6 5-726-5-723; grey, pearly crystals 16/14 4-71 (Bettendorff, A.

144. 110); amorphous $\frac{140}{14}$ 4710-4716 (ibid. l.c.); fused $\frac{19}{19}$ 5.709 (Mallet, C. N. 26, 97). V.D. 147-2 at 860° (Deville a. Troost, C. R. 56, 871); 153-7 at 640-670° (Mitscherlich, A. 12, 159). S.H. crystallised, 093; black, amorphous, 0758 (Bettendorff a. Wüllner, P. 133, 293). C.E. (linear at 40°) 00000559 (Fizeanx, C. R. 68, 1125). E.C. (Hg at 0°=1) 2-679 at 0°, 1×73 at 100° (Matthiessen a. Bose, T. 152, 1). S.V.S. cryst. 13·1; amorph. 15·9. $\frac{\mu_{\Lambda}-1}{d} \times$ at, wt. 15·4 (Gladstone, Pr. 18, 49). Chief lines in emission spectrum, v. Huntington, P. Am. A. [2] 9, 34; Hartley a. Adeney, T

1884. 121. Occurrence .- Found native, but more frequently associated with other metals and sulphur, in widely distributed ores. Obtained as a principal product chiefly from native arsenic, arsenical iron FeAs, and FeAs, and arsenical pyrites FeAsFeS2; obtained as a secondary product from smaltine, cobalt giance, arsenical cobalt, nickel glance, many fahl-ores, &c. Occurs also in ferruginous deposits of certain mineral waters (Will, A. 61, 192); in nearly all iron ores (Walchner, A. 61, 205); in soils, from the weathering of iron pyrites (Sonnenschein, Ar. Ph. [2] 113, 215); in the residue obtained by evaporating sea water (Danbrée, Ann. M. [1] 19, 669); frequently found in metallic bismuth (Schneider, J. pr. [2] 20, 418); in various kinds of pyrites, and hence in most samples of commercial sulphuric acid, and in many substances in the manufacture of which this acid is used (v. H. A. Smith, P. M. [4] 41, 370).

Preparation .- On the large scale by heating to redness, out of contact with air, arsenical iron or arsenical pyrites; arsenic sublimes; iron, or ferrous sulphide, remains. Prepared in small quantities at a time by heating As O6 with powdered charcoal, or with 'black flux,' in crneibles covered with conical iron caps. Also by heating As, S₃ with charcoal, an alkaline carbonate, and KCN. Purified by resublimation after mixing with powdered charcoal; or by heating with a little 1 (Ludwig, Ar. Ph. [2] 97, 23); or by boiling with moderately conc. K_CrO, Aq acidified with H.SO, (Böttger, J. pr. [2] 2, 181). Arsenic was first prepared from arsenious acid in 1694 by Schröder; its chemical nature was further investigated by Brand (1733), Macquer (1746), Manuet (1773), and others. Scheele discovered arsenic acid and arsenuretted hydrogen in 1775.

Properties.—Very brittle, steel grey, lustrous; crystallises by sublimation in hexagonal rhombohedra isomorphous with Sb and To; a: c=1:1·4025.H=3·5. When As is sublimed in a rapid H stream in a glass tube the sublimato nearest the heated part of the tube consists chiefly of rhombohedra, that farther from the hottest part but still on a warm portion of the tube (210°-220°) of black amorphous As, while the coolest part of the tube is filled with yellow tumes which condense to grey crystals (Bettendorff, A. 141, 110). Black amorphons As is also obtauned by condensing As vapour at a fairly Ingh temperature; by decomposing As sompounds by heating in glass tubes to

moderately high temperatures (e.g. AsH₃), or by heating with roducing agents (e.g. As, O, with C); or by reduction of As compounds in the wet way (Engel, C. R. 96, 497). As can be obtained in regular octahedra by heating a mixture of much H with a little AsH, (Cooke, Am. S. [2] 31, 91). Amorphous As is changed to crystalline by heating for some time at 310° (Engel, C. R. 96, 1314); by heating to 358°-360° (Bettendorff, A. 144, 110). Amorphous As when subjected to a pressure of 6500 atmospheres acquires metallic lustre and its S.G. increases (Spring, B. 16, 326). The vapour of As is citron-yellow (Le Roux, C. R. 51, 171). The spectrum of As shows lines in the orange (6169-5), yellow, and green (5331) (Thalen, A. Ch. [4] 18, 244); also many more refrangible lines (v. Hartley, T. 1884, 124). As combines with Cl and O with production of heat; [As, Cl'] = 71,390; $[As^2, O^2] = 154,670$; $[As^2, O', Aq] = 147,120$; $[As^2, O'] = 219,380$; $[As^2, O', Aq] = 225,380$ (Thomsen). As volatilises at a dark red heat without previous fusion at ordinary pressures. The molecule of As is tetratomic (As,); the atom is trivalent in gascons molecules (AsII, AsCl, &c.). The atomic weight has been determined (1) by analysing, and determining V.D. of, various gaseons compounds, AsII, AsCl, AsI, As O., &c.; (2) by determining S.H. of As; (3) by comparing isomorphous compounds of As, Sb, and Bi, arsenates with phosphates and vanadates, &c. (Wallace, P. M. [4] 18, 279; Dumas, A. Ch. [3] 55, 174; Kessler, P. 95,

As is insoluble in alcohol and ether, but is said to be dissolved by certain oils. It oxidises fairly rapidly in air at ordinary temperatures; heated in air, it burns to As,O₆ with a bluish flame; is oxidised by nitrie and sulphuric acids, and by fusion with alkalis. As forms two series of compounds, of which As,O₆ and As₂O₅ are representatives.

In many of its physical properties As is metallie, but in its chemical relations it is decidedly non-metallic or negative. Exhibits allotropy; oxides are acid-forming (v. Arsenic, OXIDES OF; also ABSENIC, ACIDS OF); at the same time As Os appears to react with SO3 to form a salt, and with KH.C.H.O. to form a compound analogous with tartar emetic, and with conc. HCIAq to form AsCl, (v. Arsenious oxide, under Arsenio, oxides of). Arsenious acid is unknown, and an aqueous solution of the oxide behaves towards alkalis as a very feeble salt-forming compound; but arsenie acid is as strong an acid as phosphoric, their relative affinities are nearly equal (v. Afrinity, p. 67). The haloid compounds of As do not show any marked tendencies to form double salts. The hydride AsH, does not combine with acids, as NII, and PH, do; but at the same time compounds belonging to the form AsR X, where R is an alcoholic radicle C_nH_{2n+1}, and X is a halogen or even OH, are known (v. Arsenic compounds, organic). For a fuller discussion of the chemical relations of arsenio v. arts. BISMUTH, CHEMICAL RELATIONS OF; and NITROGEN GROUP OF ELEMENTS.

Reactions.—1. Hydrochloric acid, no action in absence of air; in presence of air a little AsCl₄ is formed.—2. Nitric acid and aqua regia react with production of much heat; exides of N,

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As, O_a, and H₂AsO_a ars formed.—3. Hot. sono. sulphuric acid evolves SO₂ and forms As₁O_a—4. Molten potash or soda produces an arsenite and H.—5. Molten nitre or potassium chlorate produces potassium arsenate; the action is more or less explosive.—6. Solution of sulphur dioxide reacts, when heated with As in a closed tube to 200°, to produce As₁O_a, S, and H₂SO₂Aq, but no sulphide of As (Geittner, J. 1861, 113).—7. Ammonia solution is without action on As.

Combinations .- I. With nascent hydrogen AsII₃ and AsII are formed (q. v.).—2. With chlorine, bromine, or iodine, AsCla, AsBra, or AsI, (q. v.) is produced. -3. With fluoring (action of HF on As Og) AsF3 is formed (q. v.). -4. With oxygen As combines to form As O. (q. v.); As O. (q. v.) is produced by heating one of its hydrates. -5. The sulphides As.S. and arsenic and sulphur; the sulphide As, S, is, however, best obtained by decomposing solutions of alkaline sulpharsenates by acid. 6. Tellurium combines with arsenic to form As Te₂ and As Te₃, when the two elements are melted together in the required proportions (Oppenheim, J. pr. 71, 266).—7. When arsenic is melted with sulphur and selenion in the proportions represented by the formule As, SeS, and As, SSe,, two bodies having the compositions indicated are obtained. The first is a lustrous red semi-transparent mass from which the whole of the Se separates out after some days. This body is easily soluble (when powdered) in NH, HSAq. The body As, SSe, is a crystalline opaque solid which may be distilled nuchanged; it is less easily soluble in NII, HSAq than As, SeS, (v. Gerichten, B. 7, 29) -8. Arsenic appears to be incapable of combining with phosphorus directly [older experiments by Landgrebe (S. 60, 181) probably yielded only a mixture of P and As]; but if AsH, is led into PCl3, or PII3 into AsCl3, a red-brown solid is obtained (after drying it appears as a darker powder without lustre) which is insoluble in alcohol, ether, and CIICl_s, but fairly soluble in CS... This solid is PAs; it is changed by water into PAS₂O₂ with which chlorine reacts to produce AsCl, and POCl_x. The compound PAs is rapidly oxidised concentrated HNO, less rapidly by dilute INO, giving II, AsO, and H,PO,; solutions of OH, NH,OH, or Ba(OH)2, easily decompose PAs rapidly when warm) producing PH₂, AslI₄, I₂PO₂, II₂AsO₃, and As. Heated in air PAs burns o As.O. and P.O.; heated in absence of air, or n CO, phosphorus sublines and then arsenic. the reactions of P.As,O. are very similar to hose of PAs (v. Janowsky, B. 6, 216; 8, 1636). -Arsenic forms alloys with many metals. Some f these are produced by very strongly compressing the constituents (Spring, B. 16, 324). These alloys are generally brittle, they are only parfally, in many cases not at all, separated into peir constituents by the action of heat out of ontact with air; they are generally oxidised to resenates, and oxides of the metals, by fusion with nitre; fused with alkalino carbonates and ulphur, thio-arsenite or thio-arsenate of the alkali netal is generally produced, and the metals fornerly alloyed with the arsenic are completely eparated as sulphides. Arsenides of heavy octals are scarcely if at all attacked by nitric cid or aqua regia. Many alloys of arsenio are

definite compounds; several of them occur native as minerals (v. Winkler, J. pr. 91, 193; Scharmont, A. Ch. 80, 221; Rammelsberg, P. 128, 441). The alloys with cobalt, which are brittle and iron-grey in colour, are formed, with production of heat, by melting the elements together. CoAs occurs native as Smaltine; it always contains more or less iron and nickel replacing part of the arsenic. Co.As,, generally containing more or less iron, also occurs native as Skutterudite, or Modumite. Arsenio alloys with copper to form white solids which tarnish in the air. According to Lippert (J. pr. 81, 168) the grey deposit obtained by heating coppor in an HCl solution of arsenious oxide is Cu,As,; when this body is heated in hydrogen Cu As remains. The compounds Cu₃As, Cu₄As, and Cu₄As, occur native as Domeykite, Algodonite, and Darwinite, respectively. The alloys of arsenic and iron are brittle solids formed by melting the elements together; FeAs, and Fe, As, ocenr native as Arsenical iron, sometimes containing Ag, Au, and Cu. Arsenie alloys with lead to form brittle solids. With nickel, arsenie alloys easily; NigAs is obtained by melting the clements together. The minerals Copper-nickel NiAs, and Clounthite NiAs., occur native; they contain varying quantities of Sb. Fe, Pb. Co, and Co. A lustrons crystallino alloy Ni3As2 is obtained by reducing arsenate of nickel by charcoal at a high temperature; Ni₃As is said to be formed when KCN, As, and NiO are fused together (Descamps, C. R. 86, 1965). Arsenic also alloys, with production of much heat, with potassium and sodium; the products are decomposed by water with formation of KOH (or NaOH)Aq, AsH3, and As. An alloy of 15 parts tin and 1 part arsenic forms large leaf-like crystals. Arsenie is not much used in technical enemistry; the alloy with lead is employed in making shot. Arsenic appears to form alloys with several other metals, especially Hg (?) Pd and Pt. By strongly compressing (6500 atmos.) As with various metals, Spring (B. 16, 326) obtained several well-defined alloys, e.g. Zn, As, Cu3As, &c. &c.

Detection. - In dry way. Arsenic heated with slight access of air volatilises with a garliclike odour (probably due to a little As O and condenses on cooling as a lustrous black deposit, which is easily converted into a white crystalline sublimato (As,Oe) by heating in presence of plenty of air. Sulphides or oxides of arsenio, and the salls of arsenious and arsenic acid, yield sublimates of black amorphous arsenic when heated with an alkaline carbonate alone or mixed with chargoal or KCN (v. Fresenius, A. 49, 801; Rose, P. 90, 193). Oxide of arsenic heated with much NaC₂H₃O₂ in a tube closed at one end yields eacodyl oxide, recognised by its foul smell. In wet way. l. Arsenious compounds. (i.) Sulphuretted hydrogen passed into a solution of As,Os, or an arsenite, acidified with HC!, forms a bright yellow pp. of As.S., soluble in NH,OH, NH,SH, and (NH,) CO, solutions, and reprecipitated by HCl. As Sa is said to be soluble in a considerable quantity of boiling water and in boiling dilute 11Cl (Odling, Guy's Hosp. Rep. [3] I, 239). (ii.) Neutral solution of silver nitrate produces a canary-yellow pp. of Ag. AsO, easily soluble in most acids and in

ammonia. (iii.) Neutral solution of copper sulphate precipitates green CuHAsO, (Scheele's green) easily soluble in acids and in ammonia. (iv.) Reinsch's test (J. pr. 24, 244). A warm solution of an arsenious compound acidified with HCl deposits a film of arsenic on a piece of bright copper kept in contact with it for some timo; this deposit may be oxidieed to As O. by heating in air (v. supra). The deposit consists of. Ae,Cu, according to Lippert (J. pr. 81, 168). Areenic compounds may be detected by Reinsch's test, if present in considerable quantity (Werther, J. pr. 82, 286), even in small quantity if hoated for some time with HCl (Reinsch, N. J. P. 16, 135); the HCl used must not be weaker than S.G. 1.1 (Bettendorff, Z. [2] 5, 492). According to J. M. Scherer (Fr. 3, 200) the delicacies of the wet tests are as follows; ΛgNO_3 detects $\frac{1}{200,000}$ part of As, CuSO₄ detects 1 part, H₂S detects 1 part, and Reinsch'e test detects $\frac{1}{120,000}$ part, or after boiling for $\frac{1}{4}$ hour $\frac{1}{250,000}$ part of Ae. (v.) Marsh's test. When hydrogen is evolved in contact with an acidified colution of an arsenious compound, whether by the action of zine (March, B. J. 17, 191; 20, 190; 22, 175), magnesium (Roussin, J. 1866, 801), or the electric current (Bloxam, C. J. 13, 14), arsenic trihydrido ie evolved; arsenio may be separated by passing the gas through a glass tube heated at one part, or by burning the gas in a limited supply of air and presenting a cold surface of porcelain, or thick platinum (Davy, J. 1858. 609), on which the arsenic condenses. The deposit of arsenic is easily coluble in HNO3, S.G. 1.2 to 1.3; the solution contains arsenious acid (which on boiling for some time is changed to arsenic acid), it gives the characteristic yellow pp. of AgaAsO, (v. supra); the deposit of arsenic ie also easily soluble in aqueous NaClO free from Cl; if the deposit is warmed in dry 11,8 yellow As, S, is produced which is not changed when warmed in a current of dry HCl gas. Arsonious hydride passed into aqueous AgNO3 precipitates Ag, but the whole of the As remains in solution along with HNO3 formed in the reaction, and may be detected by filtering and carefully neutralising the filtrate with dilute ammonia, when yellow Ag, AsO, is produced. [SbII, precipitates Ag from AgNO, but the whole of the Sb is at the same time thrown down; the deposit of Sb obtained by heating, or burning, SbH, is insoluble in NaClO colution free from Cl, and is much less soluble than As in HNO,Aq of S.G. 1.2 to 1.3; moreover, the solution in HNO,Aq gives no reaction with AgNO,Aq and anunonia. The reactione of the two gases towards AgNO, colution affords a means for separating them (Husson, C. R. 83, 199).] The electrolytic method of preparing AsH, has the advantages (a) of avoiding the use of zinc which generally contains arsenic; (b) of not interforing with the sub-sequent testing for other motaie; (c) of allowing the separation of antimony if present-this ie done by adding a little H.S to the liquid, whereby As 2S, and Sb S, are formed; the former is easily, the latter not at all, decomposed by the current (Bloxam, C. J. 13, 14, 338). The presence of HNO, prevente the formation of AeH, (Blondlet,

J. 1868. 681). (vi.) Bettendorf's test (Z. [2] 5, 492). Stannous chloride in fuming HCl; added to a solution of As,O, or As,O, in the same acid, precipitates metallic As mixed with a little SnO,. This test is said to be extremely delicate; it may he need to detect As in presence of Sb, also to remove As from HClAq (v. also Hagor, J. 1870. 966). II. Arsenic compounds. (i.) Neutral silver nitrate precipitates rod-brown Ag₄AsO₄ solublo in NH₄OHAq and in HNO₃Aq, but less soluble in HNO, Aq than Ag, AsO,, so that if AgNO, ie mixed with a colution of an arsenite and an arsenate in HNO, Aq, and NH, OHAq is then added drop by drop, Ag, AeO, is precipitated before AgaAsOa; a solution of arsenio acid in nitric acid is precipitated by AgNO, if a few drope of a concentrated solution of an alkaline acetato are added (Avery, Am. S. [2] 47, 25). (ii.) Neutral copper sulphate precipitates bluish green CuHAsO, soluble in HNO Aq and in NH,0HAq. (iii.) An alkaline solution of magnesium sulphate, containing enough NII,Cl to prevent precipitation of magnesia, precipitates white Mg(NH,)AsO,; delicacy 1 part of As in 56,000 (Lovol, B. J. 28, 130) (cf. Estimation of Arsenic). (iv.) Ammonium molybdate in excess precipitates bright yellow arseno-molybdate of ammonium from warm (not cold) solutions of arsenates containing HNO₃. (v.) Sulphuretted nydrogen elowly reducee are enic to arsenioue acid and then (in presence of HCi) precipitates yellow As, S, mixed with S. (vi) Marsh's test. Arsonic compounds in solution are reduced to AsH, by hydrogen evolved in contact with the solution; the reduction takes place under the same conditions as, but more slowly than, the reduction of arsenioue compounds (v. supra).

Arsenic may be reduced to arsenious compounds by such deoxidising agents as sulphurous or phosphorous acid; the reverse change may be accomplished by heating with HNO₂Aq, HClAq

and KClO₃, &c.

Detection in cases of poisoning.—Arsenious oxide or white arsenie ie the usual form in which the poison is administered. Because of the insolubility of this compound, small solid particles of it may sometimes be picked out of the food or contents of the intestine; these should be tested by reduction to metallic argenic, &c., in the dry way. The poison is, however, usually mixed with large quantities of organic matter which must be removed or destroyed, after which the arsenic may be ppd. as As, S, by long-continued passage of H.S. the pp. may then be dissolved in warm HClAq with a cryetal of KClO3, free chlorine removed by warming, or the pp. may be dissolved in warm H2SO,Aq, and Reinsch's, or better March's, test used for detecting the arsenic in solution. The organio matter may be removed by diffusion (Graham); or it may be destroyed by (a) treatment with warm concentrated H2SO,Aq, (b) warming with HClAq and crystale of KClO, added from time to time, (c) warming with HClAq and a little HNO, Aq, (d) passing Cl into the liquid instead of adding KClO₂, (c) heating with HNO, Aq till the roeidue is semi-pasty and yellow, adding NaOHAq till the acid is neutralised, mixing with powdered Na.CO, and a little NaNO, drying in a orucible and gradually heating until the mass fuser (Wöhler; v. also Meyer, A. 66, 237). All re

agents used must be carefully freed from arsenio; a blank experiment should be conducted with the reagents alone. ' (Buiz a. Schulz find that oertain parts of the animal organism reduce As, O, and also oxidise As₁O₆; they think that the poisonous effects of As₄O₆ are due to rapid deoxidation and reoxidation, v. B. 12, 2199; 14, 2400; 15, 1388.)

Estimation. - I. Gravimetric methods. i.) As magnesium-ammonium arsonate. Solution of MgSO, is mixed with excess of NH,OHAq and allowed to stand for 12 hours; this liquid is added to the solution containing arsenic acid to which an excess of NH,OHAq has previously been added. After some time the pp. is collected on a weighed filter, washed with ammonia-water, and either (a) dried in wacuo over H.SO, when it has the composition Ig.NH, AsO, GH.O, or (b) dried at 100° to 110° hereby 2(Mg.NH, AsO, H.O is produced, or (c) grongly heated over a Bunsen lamp (temperatre being gradually raised) whereby Mg.As.O, formed. Method c is recommended by Puller 7. 10, 41), who states that the best means of everting As.S. into arsenic acid is treatment th rel funing 11NO, (v. also Rammels-g, B. 7, 544; Wood, Am. S. [3] 6, 368; Leivor, C. N. 32, 283).—(ii.) As arsenious In lphide. Arsenates are reduced to arsenites SO₂ solution; the arsenious solution is acidi-ted with HClAq, and As S₄ is ppd. by long conhaued passage of H.S; the pp. is washed with S, to remove any S which it may contain, collected on a weighed filter, and dried at 10°-110° (Pullor, Fr. 10, 41). — (iii.) As ranyl pyroarsenate. Uranio acetato in resence of ammonium salts pps. ammoniumranyl arsenate NH, UO, AsO, + Aq, insoluble water and acetic acid, but soluble in mineral kids. When this pp. is washed, dried, and adually heated to bright redness, the pyrosenate (UO.), As.O, is obtained. This method rocommended by Puller (l.c.); As.S, may ter adding NH, C.II, O, and H.C.II, O, and II. Volumetric methods. — (i.) By

odine. Arsenious, is converted into arsenie, cid, by iodine in presence of alkali; a solution ! NaHCO, saturated in the cold and used in excess is the best alkali (Waitz, Fr. 10, 158) .-(ii.) By potassium dichromate. Arsenites are converted into ursenates by the action of 1 K₂Cr₂O₇Aq in acid solutions; the residual K₂Cr₂O₇ is determined by a solution of FeSO₁; xcess of HCl should be avoided (Kessler, Fr. 0, 205).--(iii.) By potassium permangaate. Arsenites are oxidised to arsenates by Mn_O,Aq in solutions containing H.SO ; an cess of K.Mn.O. is added, and the residual Mn.O. is determined by FeSO, solution

aitz, *i.c.*).

Minute attention to detailed precautions must be obd in testing for arsenic in poisoning cases. These will be found in the following memotirs and les:—Marsh, B. J. 17, 191; 20, 190; 22, 175; Reg. A. Ch. (3) 2, 169; Freenius a. V. Babo, A. 49, 287; F. A. 69, 384; Schuedder, P. 85, 433; Fyfe, P. M. French, V. Wilher, a. V. Siebold, Dusty-ericht. Verfahrm better Arsenit. Fergiting (1817); Sp. Aumittelung der Gifte [English ed. On Poisons]; Sp. Jumos Prafama chom. Gifte (1807); Taylor's lical Jurisprudence; Taylor On Poisons. Minute attention to detailed precautions must be ob-Vol. I.

References .- Besides the papers referred to in the article, the following older memoirs on arsenio and its compounds may be consulted:-Soheele, Opus. 2, 28; Bergmann, Opus. 2, 272; Buchholz, S. 15, 537; Laugier, A. Ch. 85, 26; Fischer, S. 6, 236; 12, 155; 39, 364; Thomson, S. 17, 422; 29, 430; Berzelius, A. Ch. [2] 5, 179; 11, 225; S. 34, 46; P. 7, 1, 137; Gehlen, S. 15, 501; Gay Lussac, A. Ch. [2] 3, 136; Pfaff, S. 45, 95; Buchner, S. 45, 419; Soubeiran, P. 19, 991; Mitscherlich, A. Ch. [2] 19.

Arsenic acid and Arsenates v. Arsenio,

Arsenic, acids of .- (In connection with these compounds v. arts. Acros; Acros, Basicity of; Hydroxides.) Arsenious oxide, As O., dissolves in alkalis, forming salts from which other arsenites may be obtained; no hydrato of As,O, is, however, known. The heat of solution of the oxide is negative; $[As^1O^6, Aq] = -15,100.$ senio oxide, As,O., dissolves in water with formation of the hydrate H,AsO4, which crystallises from concentrated solutions as 2II, AsO, H,O; these crystals heated to 100° lose water, and arsenio acid, H₃AsO₄, remains. By the action of heat on this acid, two other acids are produced, viz. at 140°-180' pyroarsenic acid HASO, and at 200° metarsenic acid HASO,; these acids dissolve in water with reproduction of HyAsO4. Each arsenic acid yields a series of salts; the arsenates (or ortho-arsenates), divisible into three classes of the forms MH, AsO., M.HAsO., and M, AsO.; the pyroursenates M, As.O.; and the metarsenates MAsO.. The pyro- and metarsenates have not been much investigated. they appear to exist only as solids and to yield arsenates when brought into contact with water. Thomsen's thermal examination of the behaviour of aqueous solutions of the two oxides of arsenic towards soda shows that these solutions differ much as regards reactions and hence also as regards composition. In each case the thermal behaviour of the oxide of arsenio is compared with that of the corresponding exide of phosphorus (the formula As O3 is used as being directly comparable with P.O3). x [As $\overset{\circ}{O}$ $\overset{\circ}{A}$ $\overset{\circ}{Q}$ $\overset{\circ}{A}$ $\overset{\circ}{A$

1	7.300		٠	14,800
	.,	6,500	13,700	,
2	13,800	•		28,500
		1,200	400	
8	15,000			28,900
		600		
6	15,600			•
x[I	As ² O ⁵ Aq, <i>x</i> N	aOHAq] p	iff, [POAc	,xNaOHAq į.
1	14,800		<u> </u>	15,000
		12,300	12,600	
2	27,100			27,600
		6,900	8,400	•
8	34,000			36,000
		1,300	1,400	
6	35,300		-	37,400
. 111		1 1	. 4 . 4 3	

The mean thermal value of the reaction which occurs when one formula-weight of soda is added to an acid is 13,500; this value is reached when 2NaOHAq is added to As OaAq, but a little more heat is produced when a third formula-weight of soda is added. The values when baryta solution is used are [As2O3Aq, BaOAq] = 14,000; [As'O'Aq, 2BaOAq] = 15,600. It seems probable that the sods reacts with

water and arsenious oxide, not with arsenious soid, to produce an arsenite NaAsO,HAsO, analogous to potassium arsenite, and that addition of more sods changes this either into the normal, or some other, arsenite.

I. Arsenioue acid and Arsenites (Pasteur, 1. ARSENTOGE ACID and ARSENTIES (FASCULT, A. 68, 308; Filhol, A. 68, 308; Kithn, Ar. Ph. [2] 69, 267; Streng, A. 129, 238; Stoin, A. 74, 218; Reynoso, C. R. 31, 68; Girard, C. R. 34, 918; 36, 973; Bloxam, C. J. 15, 281). No arsenioue acid has been obtained in separate form; but an aqueous solution of the oxide As O reacts with bases to form unstable salts the compositions of which may be well represented by regarding them as derived from one or other of the three hypothetical arsenious acids, H₂As₂O₄(= As(OH)_a), HAsO₄(= As.OOH), H₁As₂O₄(= As.O(OH)_b). The arsenites as a class are very easily decomposed; the ammonium salt gives off ammonia at ordinary temperatures and pressures, its aqueous solution yields pure As, O, on evaporation; the potassium and sodium salts are decomposed in solution by carbon dioxide with separation of As,O,. CaO, BaO, and SrO, dissolve when boiled with water and As O., addition of lime-baryta- or strontia-water precipitates arsenites insoluble in water, but solublo in acids and in ammonia. Solutions of arsenites of metals which form sulphides soluble in water are decomposed by H.S with precipitation of As Sz; if the metal of the arsenite forms a sulphide insoluble in water then H.S precipitates thie sulphide along with As, S,. Many arsenites are not, however, decomposed by metals even when the oxide of the metal of the arsenite is insoluble in potaslı. Insoluble arsenites are obtained by adding a soluble salt of the metal to a solution of K or Na arsenite. All arsenites, with the exception of those of the alkali metals, are partially or wholly insoluble in water; when formed they usually retain some arsenious oxide, so that it is difficult, and sometimes impossible, to obtain definite compounds of fixed composition. Most arsenites are decomposed by heat with formation of an arsenate and arsenie; heated with carbon, or with carbon and borax, the whole of the arsenic is usually separated in the metallie state. Heated with ammonium chloride, most arsenites yield AsCl, and chloride of the metal of the arsenite. Solutions of the alkaline arsenites exposed to the air absorb oxygen and produce arsenates (Fresenius, J. 1855. 382).

Arsenites. Ammonium arsenites NH, AsO2 (Pasteur, Bloxam); (NH4), As2Os (Stein). By dissolving As,O, in cone. aqueous NH,; white lustrous prisms, very soluble in water.

Barium arsenites Ba(AsO₂)₂; white gelatinons pp. by adding BaCl₂Aq to KAsO₂Aq (Filhol). BaH₄(AsO₂)₂; by mixing BaCl₂Aq with As₄O₆Aq and NH₂Aq (Bloxam). Ba₂As₄O₆AH₂O; by dissolving As₄O₆ in BaOAq (Stein).

Calcium arsenites. Solutions of the various potassium arsenites mixed with CaCl.Aq yield potassium arsentes mixed with Caclady pied pps. of varying composition (Filhol, Stoin). A boiling solution of As,O, added to CaOAq precipitates Ca₂(AsO₃)₂(Kühn). CaCl₂Aq mixed with As,O,Aq and NH₃Aq precipitates Ca(AsO₂)₂ (Simon, P. 40, 417). As,O,Aq with oxcess of CaOAq precipitates Ca2As2Os (Stein).

pp. produced by reaction of KAsO, Aq with CoCl. Aq in presence of NH, OlAq.

Copper arsenites. CuSO, Aq + KAsO, Aq, or ammoniacal CuSO,Aq + As,O,Aq, yields a light green pp. (Scheelo's green) of CuHAsO,; soluble in NH, Aq with formation of H, AsO, and Cu2O; eoluble in KOHAq with formation of KASO, and Cu.O, which precipitates; when heated CullAsO, evolves As and H.O, and a mixture of CuO and copper arsenide remains. By digesting CuCO, with H.O and As, O, and evaporating the eolution, a yellow-green salt, probably Cu(AsO2)20 is produced.

Iron arsenites. Ferrous arsenite Fc, As, O. is a greenish white pp., soluble in NH,Aq, obtained by mixing FcSO,Aq with As,O,Aq and a little NH,3Aq. Various ferric arsenites appear to exist. Freshly precipitated Fe (OII) digested with conc. As OcAq containing not more than 14 as much As Oca as there is Fc2(OH) completely converts all the As,O, into an insoluble salt (Bunsen); with less Fe (OH), the whole of the As O is not removed from solution, a basic salt, FeAsO3.Fe.O3, is produced from which water removes some As,O₆. A basic salt (rusty-red, hard, soluble in NaOHAq), 2FeAsO₃, Fe₂O₃, 7H₂O₄ is produced (a) by adding Fc,3SO,Aq or Fc,Cl,Ac to KAsO.Aq, (b) by oxidising FeSO.Aq by aqui-regia, nentralising by NH,Aq, adding a saterated solution of As,O, in NaOHAq, and allowing to stand for twelve hours. Another stlt, 2FeAsO₃.Fe₂O₃.5H₂O₄ is obtained, as an ochrayellow pp. drying to a brown mass, on mixing As O.Aq or KAsO.Aq with Fc. (C.H.O.), Aq; water withdraws part of the acid from this salt; it is decomposed by heat, losing all its arsenious acid (Simon); only a part of its acid (Bunsen; v. also Grimaux, Bl. [2] 42, 211).

Lead arsenites. Pb(AsO₂)₂ is a white pp.

(Filhol), melting to a yellow glass (Berzelius), obtained by adding KASO Aq to Pb(C.H.O.)2Aq. Other salts are known: Pb.As.O(?) by precipitaother sants are known: D₂, ks_0(t) by precipitating Ph(C₂H₃O₂), Aq by As₁O₈Aq and NH₂Aq (Filhol, Bloxam); Pb₃(AsO₃), by precipitating Ph(C₂H₃O₂), Aq by (a) hoiling As₁O₈Aq (Kühn, Bloxam), or (b) As₁O₈ in NaOHAq (Streng, A. 190, 202). 129, 238).

Magnesium arsenites. A solution of As O. in excess of NH3Aq mixed with MgSO,Aq with NII, ClAq gives a pp. which is Mg, (AsO_a)₂ (Stein) MgHAsO3.H2O (Bloxam); heated to above 250° Mg.As.O, remains. Other salts appear to exist but their composition is vague and uncertain (v. Filhol, I.c.).

Manganese arsenite. Mn3Ha(AsO3)4.2H2O is a light rose-red pp. obtained by addin NH,AsO,Aq to MnSO,Aq. Mercury arsenites. Composition undecide obtained by adding

Hg2NO3Aq mixed with As,O6Aq givos a white pp. soluble in KAsO₂Aq; if excess of KOH is present Hg is precipitated. An arsenite of mer oury seems to be formed by mixing HgNO. with KAsO2Aq, or by digesting AsO.

with Hg.
Nickel arsenite. Ni₃H₆(AsO₃)₄H₂O is of tained as a greenish-white pp. on adding KAsO, Aq to NiCl, Aq containing much NH, Cl; heated in air, this salt loses H,O, then As,O,, and yellow infusible Ni_s(AsO₄), remains.

Potassium arsenites. The acid salt Cobalt arsenite. Co₂H₄(AsO₂).H₂O; rose-red! KAsO₂:HAsO₄:H₂O is produced by adding alcolul

to a solution of much As O, in KOHAq (Pasteur, Bloxam). By digesting this salt with K₂CO₂Aq the neutral salt KAsO, is produced (l'asteur, Filhol); by treating this with KOHAq and preoipitating by alcohol the salt K₁As₂O₃ is formed (Bloxam). Two double salts, KASO₂.IIASO₂.ASOI and K.H., 3As,O., 2KI are described by Emmet (Am. S. [2] 18, 583), and Harms (A. 91, 371), obtained by adding KIAq to As,O., Aq, or KIAq to KAsO Aq containing so much IIC Hao, that no brown colour is produced with turmeric

Silver arsenites. Ag₄AsO₃, a yellow pp. obtained by adding As₄O₄Aq neutralised by NH₃Aq to AgNO, Aq (Külin, Filliol, Bloxam); solublo iu HNO3, II.C.H.O., NII, Aq, and solutions of ammoninm salts, also in KOHAq; solutions are not precipitated by KClAq, but dissolve freshly pre-pared AgCl. Heated alone or with alkalis it pared AgCl. I

 $4\Lambda g_3\Lambda sO_3 = \Lambda g_1O + 2\Lambda g + 2\Lambda g_2\Lambda sO_4 + \Lambda s_2O_3$ (Wöhler, A. 101, 363). Other silver arsenites are described by Harms (l.c.).

None obtained pure; Sodium arsenites. scem to be all very soluble in water (Pasteur,

Filhol, Bloxam).

Strontium arsenite. Sr(AsO₂),4II.O. By mixing SrCl₂Aq with KAsO₂Aq and allowing to doposit crystals slowly (Stein).

Tin arsenites. Scarcely known; both stannons and stannic arsenites seem to exist.

II. ARSENICACID and ARSENATES. - (Setterberg, B. J. 26, 206; Baumann, Ar. Ph. 36, 36; Kodschonbey, J. pr. 49, 182; Field, C. J. 11, 6; Uelsmann, Zeits. f. d. ges. Naturwiss. 23, 347; Schiff, A. 112, 88; Maumené, C. R. 58, 250; Debray, A. Ch. [3] 61, 419, also C. R. 59, 40; Leohartier, C. R. 65, 172; Salkowski, J. pr. 104, 129). The conditions of formation of the three arsenio acids, H2AsO,, HAsO,, and H4As2O, have been already described (p. 305). The following thermal data are given by Thomsen (Th. 2, 236) $[As, O^4, H^2] = 215,630; [As, O^4, H^3, Aq] = 215,230; [As^2O^4, 3H^2O] = 6,800; [AsO^4H^2, Aq] =$ [As'0', 3H''0]=6,800; [As'0', Aq]=2,000; [As'0', Aq]=400; [As'0', H']=360,830; [As'0', Aq]=6,000; [As'0', O']=64,710; [As'0'Aq, O']=78,260. The following heats of neutralisation are also given by Thomsen (Th. 1, 196):

n [Na	$\mathrm{OHAq}, n\mathrm{H}^{2}\mathrm{AsO}$	'Aq]:
j .	6,233	
j.	11,972	
Ä	13,790	
5 5 1 2	14,994	
2	14,724	
n [n]	TaOHAq, H3AsO	Aq].
∯ _	7,362	_
i	14,994	Diff.
Ω	27,580	12,586
8	35,916	7,236
•	•	1,484
6	37,400	

The acid H, AsO, is therefore tribasic; but as the reaction of the third formula-weight of soda is accompanied by the production of not much more than half the quantity of heat which accompanies the reaction of the first or second formula-weight, it is probable that this acid belongs to the same class of tribasic acids as

phosphoric acid, which class is represented by Thomsen by the typical formula HR(OH)₂H (v. Acros, basicity of). The relative affinity of arsenic soid (v. AFFINITY) is 21 (that of HNO. = 100) which is a little less than the value for phosphoric acid.

The acid H.AsO, forms translacent crystals; the acid H,As,O, clear Instrous orystals; and the meta-acid HAsO, a white somewhat iridescent solid; the ortho-acid dissolves in water with disappearance of heat (v. supra), the other acids dissolve with production of heat and formation of the ortho-acid; [H'As2O7, Aq] = 1,300 (Thomsen, B. 7, 1003).

The following facts rofer to the ortho-acid H₃AsO₁.

Formation .- By action of HNO Aq on As or

Preparation.—Arsenious oxido is suspended in water and chlorine is passed in; the solution is ovaporated to dryness in a platinum dish, the temperature not being allowed to rise much above 100° the residuo is dissolved in water and evaporated slowly at 100°; after a time small needle-shaped crystals of H₂AsO₄ separate out (E. Kopp, J. pr. 69, 270).

Properties.-Action of heat already desoribed (v. p. 305). Aqueous solution tastes sour and metallic; it burns the skin. The most concentrated solution has S.G. 2.5; the following table gives some data regarding S.C. of aquoous solutions (Schiff, A. 113, 183).

S.G. at 15° 1.7346 Petge. of H, AsO, 67.4 45.0 1.3973 ,, 1.235030.0 ,, " 1.1606 22.5 ** 1.1052 15.0 1.0495

Reactions .- 1. Heated with carbon, many metals, or potassium cyanide, it yields arsenic .-2. Heated with conc. hydrochloric acid AsCl, and Cl are produced; if the S.G. of the aqueous HCl used is less than 1.04 no AsCl, is produced (Fresenius and Souchay, Fr. 1, 448); with acid of S.G. 1.1 traces of AsCl₄ are formed (Mayrhofer, A. 158, 326).—3. Sulphurous acid produces arsenious oxide and sulphuric acid (Wöhler, A. 30, 224) .- 4. Zinc and iron dissolve in aqueous II, AsO, with evolution of H and AsII, and produotion of gelatinous solids (? arsenates of Zn or Fe); in presence of H2SO4, these metals precipitate As, and ovolve H and AsII, -5. Sulphydric acid (H.S) reduces H.AsO, in solution to As.O., with precipitation of S, and then throws down As S; this reaction takes place slowly at 10°-15°, but more rapidly at 60°-70°. 6. Sodium thiosulphate solution, in presence of HCl, produces a pp. of As₂S₃ (q. v.).—7. Tung-states of potassium, ammonium, and silver when states of potassium, animonium, and situer when evaporated with KII_AsO₄ solution yield complex compounds, viz., 6WO₂.As₂O₃8K₂O.3II₂O; 6WO₂.As₂O₄(NH₄)₂O.5H₂O; and 16WO₂.As₂O₅Gag₂O.11H₂O (Gibbs, P. Am. A. 15, 1; 16, 109; v. also Debray, C. R. 78, 1408; and 15 and and Seyberth, B. 7; also Sonnenschein, J. pr. 53, 339, 391). Dorivatives of arsenio acid in which OH is replaced by Cl, or Br, or I, are not known.

Arsenates. Arsenates are isomorphous with corresponding phosphates. The arsenates of the alkali metals, and the acid arsenates of the alkaline earth metals, are scluble in water; they lose all their arsenic as chloride by strongly heating with sal-ammoniao (Rose, P. 116, 453). The neutral and basic arsenates are easily soluble in mineral acids, including arsenio acid. Debray (l.c.) describes a series of amorphous arsenates, MH_AsO, H_O, obtained by precipitating alkaline arsenates by salts of the metals M; these arsenates become crystalline, according to Debray, by long-continued digestion with the liquids in which they are produced. By fusing chlorides and fluorides with arsenates, Lechartier (l.c.) obtained a series of crystalline salts analogous in composition to Wagnerite and Apatite; thus:

Corresponding to Wagnerite [(PO.F.Mg2)].

AsO, Cl.Ca, AsO₄.Cl(F).Mg₂ AsO₄.Cl.Mg₂ AsO, Cl.Mn2

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Corresponding to Apatite [(PO,)3.Cl(F).Ca,]. (AsO₄)₃.Cl.Sr₅ (AsO₄)₃.Cl.Ba₅. (AsO₄)_s.Cl.Ca₅ (AsO₄)₃.Cl(F).Ca₅ (AsO₄)₃.Cl.Pb₅

Debray also obtained the following crystallised insoluble arsenates by digesting arsenio acid solution with carbonates of the various metals: 2A₈O₄CaH.H.O; A₈O₄CuH; 2A₈O₄CuH.3H₂O; (A₈O₄)₂Cu₃.4H₂O; and A₈O₄Cu.(CuOH). Arsenates are usually prepared by adding Na2HAsO, Aq to solutions of salts of the various metals, using the calculated masses of the reacting bodies.

Aluminium arsenate. White pp. by adding a soluble arsenate to a solution of an Al salt.

Ammonium arsenates. (NH₁)₂.H.AsO₄; by adding NH₃Aq to conc. H₃AsO₄ until tho pp. which formal discourse (Schloracki). (NH₁) 4.00 SH O forms is dissolved (Salkowski). (NH₄)₂AsO₄.3H₂O; hy adding considerable excess of NH, Aq to H₂AsO₄Aq. NH₃II,AsO₄; hy adding one formula-weight II₃AsO₄ to one formula-weight (NH.),HAsO,.

Barium arsenates. BaIIAsO,; crystalline, obtained by adding Na_HIAsO,Aq to BaCl_Aq. Ba,2AsO,; crystalline, obtained by adding Va_AsO,Aq to BaCl_Aq; acted on by water it forms the salt BaIIAsO, (Field). The salt BaH (AsO.), is obtained by dissolving BaHAsO. in warm H3AsOAq and allowing to crystallise (Setterberg). Two barium-ammonium arsenates 2Ba.NH₁.AsO₄.11₂O₅ and BaH₂(NH₄)₂(AsO₄)₂ are described (Baumann, Mitscherlich).

Bismuth, arsenate. 2BiAsO, H,O; yellowish white pp. insoluble in water and HNO, Aq, soluble in HClAq; insoluble in a cone. solution of Bi3NO, containing a little free acid (Schneider, J. pr. [2] 20, 418).

Cadmium arsenates. White crystallino pps. obtained by precipitating solution of Cd salts by Na.HAsO,Aq; two are known, Cd3(AsO4)2.3H2O and Cd, II, (AsO,) .4II,O (Salkowski).

Calcium arsenates. Ca.II.AsO,; prepared like BaHAsO,; occurs nativo as Pharmacolite. By treatment with NH, Aq this salt yields Ca, (AsO), (Kotschonboy). Two calcium ammonium arsenates are described, CaH2(NH4)2(AsO1)2 and CaNH. AsO. 6 or 7H.O (Baumann, Kotschonbey; also Bloxam, C. N. 54, 163). The salt Ca. As. O. is obtained hystrongly heating Ca.NH, AsO, 7H,O (Bloxam).

Cerium arsenate. Ce.H.AsO,; insoluble in water, soluble in excess of H, AsO, Aq.

Cobalt arsenates. CoHAsO, is unknown; CoH₄(AsO₄), obtained by ovaporating in vacuo solution of CoO.H₄O in excess of H₄AsO₄Aq. Cobaltio arsenate, Co,2AsO,8H,O (Karston, P. 60, 266) is a reddish powder obtained by precipitating the solution of a Co salt by Na, AsO, Aq; occurs nativo as Cobalt-bloom. An impuro basic arsenato of oobalt is known in commerce as Chaux metallique.

Copper arsenates. Cu, H2(AsO1), 2H2O; obtained as a blue pp. by adding (NII,), HAsO, Aq to CuSO Aq and drying at 130° (Salkowski). Cu, (AsO,) is a green powder obtained by preonjutating CuSO_AQ with Na₂HAsO_AQ; this salt dissolves in NH₂Aq, and on long standing, clear blue crystals having the composition ([NH₂]₂Cu]₃(AsO₁), separato out (Damour, B. J. 26, 247; and 27, 181). Various basio arsenates of copper occur native

ninerals (v. Coloriano, Bl. [2] 45, 707).

Chromium arsenates. Chromio arsenate is a green pp. obtained by mixing solutions of a chromic salt and K.AsO.; composition uncertain. By mixing solutions of K.CrO. and H.AsO. Schweizer (J. pr. 39, 267) obtained a gelatinous pp. having the composition (dried at 100°) (CrO), AsO, K,H(AsO₄), 4H₂O.

Didymium arsenate $Di_2(AsO_4)_2$. $DiH(AsO_4)_2$. H_2O ; by boiling H_3AsO_4Aq with Di_2O_3 (Marignac, A. Ch. 88, 185).

Iridium arsenate is a brown pp.

Iron arsenates. The ferrous salt is obtained by double decomposition; it oxidises easily; after drying, its composition is Fe.OAsO, 53H, O (Wittstein, Viertel-jahrsschr. pr. Pharm. 15, 185). The ferric salt (?FeH_d(AsO)_) is obtained by mixing Fe₂Cl₆Aq with Na₂HAsO₁Aq; it is a white powder which loses water on heating and becomes reddish, soluble in H, AsO, Aq, but this solution soon decomposes, unless very con-centrated, depositing Fe. (OII), (Lunge, Fr. 6, 185). Various other ferric arsenates of uncertain composition have been described by Berzelius (v. also Grimanx, Bl. [2] 42, 211).

Lead arsenates. The salt PbHAsO, is obtained by double decomposition (Salkowski); when treated with NH₃Aq it yields Ph₃(ASO₃).

Lithium arsenates. 2Li, AsO, HO is a white powdery pp. obtained by dissolving Li₂CO₃ in H₂AsO₄Aq and adding NH₃Aq; when this salt is dissolved in H3AsO1Aq, trimetrio prisms of 2LiII, AsO, 3H, O crystalliso out (Rammelsberg, P. 128, 311).

2MgHAsO4.13H2O; Magnesium arsenates. white pp. by mixing 5 parts Na, HAsO, in solution with a dilute solution of 3 parts MgSO, (Graham, A. 29, 21). Mg, (AsO,), is obtained by boiling the preceding salt for some time with come. Na,HASO,Aq. A magnesium-ammenium arsenate MgNH,AsO,6H,O is obtained by adding H₃ASO, Aq with excess of NH₄Aq to MgSO₄Aq containing NH₄Cl₁; it is slightly soluble in NH₄ClAq, strongly heated it loses NH₃ and Mg₂As₂O, remains [H. Rose; Field; Wittstein; Puller; Fresenius (Fr. 3, 206); Levol (A. Ch. [3] 17, 50)]. A magnesium-potassium arsenate MgKAsO,, and a corresponding sodium salt, are obtained by fusing MgHAsO, with K2CO, and KOH (or Na CO, and NaOH).

Manganese arsenates. MnHAsO, is produced by saturating H₂AsO,Aq with MnCO₂; using excess of H₂AsO,Aq the salt MnH₁ (AsO₂); is formed (Schiefer). Themanganeso-ammonium salt MnNH,AsO₂,6H₂O is obtained like the corresponding Mg salt as a gummy pp. which becomes crystallino. Some basic Mn arsonates are described by Coloriano (Bl. [2] 45, 709).

Mercury arsenates. The mercurous salt Hg_HAsO, is a yellowish white pp., which becomes orange red, produced by adding H_AsO, Aq to HgNO_Aq; it is insoluble in water and H.C.H._9_Aq, soluble in INO_Aq. When heated, this salt loses H_O and Hg, and a mercuric salt (probably Hg_As_O,) remains. By adding an excess of H_AsO, Aq to the mercurous salt, and evaporating, the uncta-salt Hg_(AsO_)_r is obtained as a white powder, somewhat soluble in INO_Aq, decomposed by IlClAq with withdrawal of half its arsenic as H_3AsO_ (Simon, P. 41, 424).

Molybdenum arsenate is a grey pp. (for combinations of As₂O₂, with MoO₃ and K₂O₂. Arsenic Acid., Reactions, No. 7).

Nickel arsenates. Analogous to cobalt arsenates (q. v.). The salt Ni₃(AsO₄)₂.8H₂O occurs native as Nickel-bloom.

Palladium arsenate is a clear yellow pp. obtained by heating Na₂HAsO₄Aq with I'd.2NO₄Aq.

Platinum arsenate is a light-brown pp. obtained like the Pd salt.

Potassium arsenates. The salts K₂AsO₄ and K₂HAsO₄ are obtained by adding K₂CO₃ or KOII to H₂AsO₄Aq; the former crystallises from vory cono. solutions (Graham, P. 32, 47). By heating together equal parts of KNO₂ and As₄O₆ dissolving in water and ovaporating, or by adding KOIIAq to H₂AsO₄Aq until neutral to litmus and crystallising, the salt KH₂AsO₄ is obtained. The crystals of this salt are fairly soluble in water (S. 6° = 19), insoluble in alcohol. Heated above 288° the salt melts to a glassy mass.

Silver arsenates. Ag₈AsO₄ is a dark redbrown pp. produced by mixing H₂AsO₄Aq, or selution of an alkaline arsenate, with AgNO₂Aq; it melts to a brownish-red glass, it is decomposed by IICIAq to AgCl and H₂AsO₄Aq, it is soluble in IIC₂II₃O₂Aq, NII₄Aq, and solutions of many NII₄ salts. By digesting this salt for some time at a gentle heat with H₂AsO₄Aq a white salt, 2AgAsO₂AsO₃, is obtained.

Sodium arsenates. Na.HASO, 12H_O, S.G. 1.67 (Schiff), is obtained in large crystals, isomorphous with common sodium phosphate, by adding a slight excess of Na₂CO₂Aq to H₄ASO₄Aq, and evaporating at a temperature not higher than 18° (Fresenius, J. pr. 56, 30). The crystals which form at 20 and upwards centain 7H₂O, S.G. 1.87 (Schiff); the crystals with 12H₂O effloresce quickly, those with 7H₂O do not. The salt melts when heated forming a clear transparent liquid. At 0° a salt crystallises with 13½ H₂O (Setterbeg). The salt NaH₂ASO₂H₂O is formed in large crystals by adding Na₂CO₂Aq to H₂ASO₂AQ until the solution is no longer ppd. by BaCl₂Aq; isomorphous with the corresponding phosphate; S.G. 2.535 (Schiff). A solution of S.G. about 1.7 gives crystals of NaH₂ASO₂PH₂O₃(Joly a.

Dufet, C. R. 102, 1891). A conc. solution of H₁AsO, decomposes NaClAq and NaNO, on heating. Two double salts, viz. NaKHASO, TheO (Schiff), or 9H₂O (Kotsolionbey), and NaNH,HASO, 4H₂O, are described (Uclsmann). The former is obtained by neutralising NaH₂AsO, Aq by K₂CO₂Aq, the latter by mixing solutions of Na₂HASO, and (NH₂)IIAsO. The compound Na₃AsO₄,NaF-12H₂O crystallises in monometric octahedra; it resembles common alum in appearance; S.G. 2·849; S. (25°) = 10·5; S. (75°) = 50. This salt is obtained by fusing together, with special precautions, 1 part As,O₉ 4 parts Na₂CO₃, 3 parts NaNO₃, and 1 part CrF₂ (Briegleb, A. 97, 95). Two compounds of sodium arsenates with sulphates seem to exist Na₂As₃O₄ (Setterborg).

Strontium arsenates. By adding Na. IIAsO, Aq to SrCl, Aq a pp. of SrNaAsO, H₂O is obtained; the filtrato from this when evaporated gives a

white pp. of SrHAsO, (Salkowski).

Tin arsenates. 2SniHAsO, H.O; a white pp. produced by adding excess of K.HAsO, Aq to SnCl.Aq; by the reverse process the salt Sn(SnCl)AsO, H.O is said to be formed (Lennson, A. 114, 113).

Zinc arsenate Zn₂(AsO₄)₂-3H₂O is a gelatinous substance (Köttig, J. pr. 48, 182 and 256); the salts ZnHAsO₄, ZnH₄(AsO₄)₂, and Zn₂(AsO₄)₂,NH₂, are also known (v. also Friedel, Bt. [2] 5, 433).

The compositions of the remaining arsenates, which have been very slightly investigated, are expressed by the following formule:

expressed by the following formule: Thallium, Tl.AsO, 2H.O. Thorium, Th.HASO, Uranium, (UO),HASO, 4H.O; the salts (UO),NASO, and (UO),Ou(AsO,),2H,O are also known (Werther, A. 68, 312). Vanadum, VII_(AsO,), Yttrium, YIIAsO, Arsenates of titanium and zirconium are said to have been obtained.

Arsenic, alloys of, v. Arsenic, Combinations of, No. 9.

Arsenic, ammonic-chloride of, v. Arsenic culonide, Combinations of, No. 2.

Arsenic, bromide of Ashr, (Arsenious bromide, Tribromide of arsenic). Mol. w. 314-25. [20°-25°]. (220°). S.G. 3-66 (Boedekor, J. 1860. 17). V.D. 157-5. II.F. 59,100 solid As, gaseous Br (Guntz, C. R. 101, 161). S.V.S. 85°S.

Preparation.—1. Powdered arsenie is shaken into a retort filled with Br vapour, and the bromide is distilled from the excess of As.—2. Powdered arsenie is added to a mixture of 1 part Br with 2 parts CS, until the liquid is colourless; more Br is then added, followed by more As, until the colour is no longer removed on the addition of As; the liquid is then filtered and evaporated; on standing, the bromide is deposited in crystals (Nicklès, C. R. 68, 837; J. Ph. [3] 41, 142).

Properties.—Colourless deliquescent prisms with strong arsenical odour, melting to a pale yellow liquid which fumes but slightly in the air. Volatilises unchanged and yields a crystalline sublimate.

Reactions.—1. Water, added in small quantity, produces AsOBr and HBr (? with a little arsenious oxide) (Serullas, S. 55, 345); added in large quantity, decomposes it into HBr and

As.O. (Serullas, Lc.). An aqueous solution containing HBr deposits the oxy-salts 2AsOBr.3As.O.12H.O and 2AsOBr.3H.O (Wallace, P. M. [4] 17, 261).—2. Ammonium bromide added to a cold concentrated aqueous solution pps. six-sided tables consisting chicfly of AsBr₃ (Wallace, P. M. [4] 17, 261). B. Sodium thiosulphate at first produces AsOBr. and then As₂S₃ (Nicklès, J. Ph. [3] 41, 112).-4. Arsenious oxide dissolves in molten AsBr3; on cooling to 150° the liquid separates into two layers, the upper of which is AsOBr, and the lower probably 3AsOBr.As.O. (Wallace, P. M. [4] 17, 261).

Combinations .- With the alkaline bromides to form rather nustable crystallisable com-

pounds (Nicklès, l.c.).

Arsenie, chloride cf. AsCl. (Arsenious chloride, Trichloride of arsenic, Butter of arsenic, Caustic oil of arsenic). Mol. w. 18111. [below 29°]. (130°·2) (Thorpe, C. J. 37, 352). S.G. \(\frac{90}{4} \) 2.205 (Thorpe, \(l.c. \)). V.D. 90.75. Volume at $t^{\circ} = \text{vol. at } 0^{\circ}$

 $+ \cdot 000991338t + \cdot 000000849t^2 + \cdot 0000000027551t^3$ (Thorpe, *l.c.*). S.H.p. 1122; S.H.v. 7034; H.V. 69-74 (Regnault, *J.* 1863, 77). H.F. 71,460 [solid As and gaseous Cl] (*Th.* 2, 326). R. 49:59 (Haagen, P. 131, 122). S.V. 94:37

(Thorpe, l.c.).

Formation .- 1. By distilling As O, with cone. H.SO, and NaCl (Glauber in 1618; Dumas, P. 9, 308).—2. By distilling As O, with conc. HCl. 3. By leading dry HCl gas over powdered As O . - 4. By passing dry HCl gas over gently heated powdered As,O3: As,O3 + 10HCl = $2AsCl_1 + 2Cl_2 + 5H_2O$ (Mayrhofer, A. 158, 326).

5. By adding 11.80, to a solution of As O, in cono. HCl.—6. By heating As S, with 3HgCl, in a retort (Ludwig, Ar. Ph. 97, 23).-7. By the action of S₂Cl₂ on As (Wöhler, A. 73, 374).—8. By the action of PCl₃ on As₄O₅ or on As₂O₅ (POCl₃, and in the case of As₂O₅ free Cl, is also formed) (Hintzig and Geuther, A. 111, 171).

Preparation.—Coarsely powdered arsenio is heated in a retort, dry chlorine is then led in, and the contents of the retort are heated; the distillate is collected in a dry receiver, and freed from oxcess of chlorine by redistillation from arsenio, or by sliaking with mercury, pouring off from the black solid which is formed, and redistilling. Or the product may be distilled npwards for some time in a current of dry CO, (Thorpe, l.c.).

Properties. - A colourless, oily, funning, very poisonous, liquid; mixes with alcohol, ether, and liquid oils; it is decomposed by water (v. infra). Distilled with aqueous IICl, AsCl, is partly volatilised; HCl prepared from arsenic containing H2SO, always therefore contains AsCl.; arsenious ohloride when hot dissolves phosphorus and sulphur, but they crystallise out again on cool-

ing.

Reactions.-1. With a little water the solid oxychloride AsOCl.H₂O(=As(OII),Cl) is produced (Wallace, P. M. [4] 16, 358). With more water, especially if hot, AsCl3 is decomposed into HCl and As O a great portion of the latter separating in the solid form.—2. With arsenic hydride it reacts to form HCl and As.-3. By the action of NO2 it is converted into As2O2, NOCI being

simultaneously produced (Geuther, J. pr. [2] 8,

Combinations.—1. Arsenious oxide dissolves in boiling AsCl, and on distillation the oxychloride AsOCl (q. v.) is obtained (Wallace, P. M. [4] 16, 358).—2. Ammonia gas is absorbed by AsCl, with production of a white solid, 2AsCl, 7NH, according to H. Rose (P. 52, 62), 2AsClN11.4NH, cl.NH, according to Pasteur (A. Ch. 68, 307), Michaelis suggests the formula 2As(NH₃Cl)₃,NH₃ (Lehrbuch der anorg. Chem. ii. 459, [1881]); this compound is decomposed by heat, giving off N11, and then a white sublimate containing NH,Cl; it is soluble in alcohol, but is decomposed by cold water with production of heat and ammonia, from the solution six-sided plates crystallise out, having the composition As₁Cl₂N₂H₁₀O₈, these are decomposed by cone. ammonia with production of (NH,)AsO2 which soon decomposes (Pasteur, l.c.). -3. Sulphur dichloride, SCl2, is said to form a compound 2AsCla 3SCla, but according to Nilson, the product of the action is merely a mixture (J. pr. [2] 12, 295).-4. With alcohol forms easily decomposed crystals of AsCla CaH O (Do Luynes, C. R. 50, 831).

Arsenic, fluoride of. AsF, (Arsenious fluoride). Mol. w. 181-9. (60°-4) (Thorpe, C. J. 37, 352). S.G. 4°; 2·666 (Thorpe, l.c.). V.D. 66·1. S.V.S. 49·5.

Formation .- By heating sodium or ammonium fluoride with arsenious bromide or chloride (Macivor, C. N. 30, 169).

Preparation. - Equal parts of dry powdered fluorspar and arsenious oxide are heated in a leaden vessel with 5 parts cone. H.SO,; the distillate is collected in a dry glass receiver.

Properties.-A transparent, very volatile, furning, liquid; it acts slowly on glass in a closed vessel, but exposed to the air it absorbs moisture with production of As₄O₆, and HF which acts on the glass. Dropped on the skin it evaporates at once but leaves a painful wound (Dumas, A. Ch. [2] 31, 434). 1t absorbs dry NH, in large quantities; miscible with alcohol and ether (Macivor, l.c.).

Reaction. -With water it forms a clear liquid (heat is produced) which soon decomposes

to As O, and HF.

Arsenic, haloid compounds of. AsF, AsCl,; AsBr,; AsI, and AsI, (? AsI,). All gasified, ercept the di- and pont-iodide, and molecular weights known; v. the arts. Arsenic, fluoride of, chloride of, browide of, lodides of, v. also art. Haloid compounds.

Arsenic, hydrides of. (In connection with these compounds v. art. Hydrides.) hydrides are known; gaseous AsH3, and solid

nAs11.

I. Arsenio trihydride. AsH. (Arsenuretted hydrogen, Arsine). Mol. w. 77.9. [-113°.5]. (-54°.8) (Ölszewski, Sitz. W. 5, 127) (about -40°). V.D. 39 1 (Dumas, A. Ch. 33, 355). H.F. -361,700 H.F. - 361,700 (solid As) (Ogier, A. Ch. [5] 20, 5). Discovered by Scheele in 1775.

Formation.-1. By dissolving zino or iron in dilute HClAq or H2SO,Aq containing As,O or As,O,.-2. By electrolysis of solution of As O Aq or As O Aq .- 3. By the action of organio matter on many arsenio compounds; e.g. the action of paper on Scheele's green (the air of rooms the paper on the walls of which is coloured with Scheele's green usually contains a little AsH.).

Preparation .- 1. By the action of water, or better very dilute HCl or HSO, on the solid alloy of As and Na which is obtained by heating Na in AsH, obtained by the action of acids on Zu oontaining As (Janowsky, B. 6, 216).—2. By the action of H.SO, diluted with 3 times its weight of water on the alloy of As and Zn obtained by heating equal parts of finely granulated Zn and powdered As in a covered crucible (Soubciran, A. Ch. [2] 23, 307; 43, 207). The gae is collected over beiled water.

Properties .- A colourless gas with repulsive odour; excessively poisonous (in working with this gas it is impossible to be too careful; more than one chemist has been killed by it, e.g. Gehlen in 1815). Easily decomposed by heat into its elements even when mixed with much hydrogen. By the action of the silent electric discharge it yields solid As hydride (Ogier, A. Oh. [5] 20, 5). Slightly soluble in water. Dry oxygen has no action at ordinary temperatures.

Reactions.-1. Heated in air it burns to As O and HO; in a limited supply of air, to As and H.O .- 2. Mixed with oxygen and subjected to electric discharge, complete decomposition to As O6 and H2O occurs explosively. - 3. Concentrated acids decompose it into its elements; conc. H.SO, Aq forms also As, S, (Humpert, J. pr. 94, 392); conc. HClAq acting for some time forms also AsCl, (Napoli, J. pr. 64, 93) .-4. Decomposed by chlorine, bromine, or indine, with production of much heat and formation of haloid compounds of As.—5. Sulphuretted hydrogen at about 300° forms As₂S₃ and H.— 6. The haloid acids IICI, HBr, HI, are without action on AsH₃ at ordinary temperatures.-7. Heated with sulphur, As S, and H S are produced (Jones, C. J. [2] 11, 648). -8. Reacts with I'Cl, to produce AsP and IICl (v. Arsenic, Combinations of, No. 8). -9. Many metals, e.g. K, Na, Sn, heated in AsH, form alloys with As and set free H.-10. Many metallic oxides, e.g. CuO, decompose AsH, when heated with it, forming arsenides and water (the quantity of AsHa in a gaseous mixture may be thus determined).-11. Absorbed and slowly decomposed by alkalis, and by water containing oxygen or air .- 12. Many metallic salts in aqueous solutions absorb and decompose AsH,; salts of metals whose oxides are easily deoxidised produce water and arsenious oxide; salts of metals whose oxides are not so easily deoxidised produce water and arsenie which is precipitated with the metal. Thus with (1) AgNO, and (2) CuSO, solution, the reactions are (1) 24AgNO3Aq + 4AsH3 + 6H.O -

As O Aq + 211INO Aq + 24Ag; (2) $2AsH_3 + 3CuSO_1Aq = As_2Cu_3 + 3H_2SO_1Aq$. Gold chloride in solution is reduced to gold, while As, O, remains in solution.

References. — (Besides those in the text) Marsh, B. J. 17, 191; 20, 190; 22, 175; Stromeyer, Comment. Soc. Gott. 16, 141; Proust, Scher. J. 8, 285; Fischer, P. 9, 261; Myere, A. 159, 127; Parsons, C. N. 35, 235.

II. Solid arsenio hydride, nash. Mol. w. Philipp, B. 14, 2643). Mol. w. unknown. unknown. Janowsky (B. 6, 220) states that Preparation.—One part As is heated with 3 when potassium or sodium arsenide is decomparts I to 230° in a sealed tube, the product is

posed by water, a solid, brown, velvet-like, compound of As and H, in the ratio As:H, separates out. When dilute soids are used in place of water, AsH, is evolved and As deposited. According to Ogier (A. Ch. [5] 20, 5), solid nAsH is produced by the action of the silent discharge on AsH₃. Older observations recorded the production of a solid compound of As and II by the action of dilute acids on arsenide of zino (v. especially Wiederhold, P. 118, 615), but these are contradicted by Janowsky (*l.c.*). According to Blondlot (*A. Ch.* [3] 68, 186), a solid hydride of arsenic is produced (1) in a Marsh's upparatus when a little IINO, is present, and (2) when AsII, is passed into aqueous AgNO3, excess of Ag is removed by adding NaCl, a drop of HNO,Aq is added, and a piece of Zn is placed in the liquid.

Arsenic, hydrexides cf. The compounds of As, O, and H, are acids; v. ARSENIC, ACIDS OF (v. also arts. Acros, and Hydroxides).

Arsenic, iodides of.

I. ARSENIOUS TODIDE, AsIa (Tritodide of arsenic). Mol. w. 454.59. (394° to 414°) (Carnelley a. Williams, C. J. 33, 283). S.G. 4:39. V.D. 227:3. S. 30:12 at 100°. H.F. solid As, gaseous 1 [As, I^3] = 28,800 (Guntz, C. R. 101, 161). S.V.S.

Formation .- I. By subliming together 1 part As and 3 parts 1 in a retort arranged so that the sublimate condenses in the neck; the sublimate is treated with hot alcohol from which the AsI, erystallises on cooling (Bette, A. 33, 349).—2. By boiling 3 parts powdered As and 10 parts I with water, filtering, and evaporating (Plisson, S. 55, 335).-3. By acting on AsCl, with conc. HI solution [11Cl is evolved] (Hautefeuille, Bl. [2] 7, 198).-4. By adding a concentrated solution of K1 to a hot solution of As,O. in HClAq (Bainberger a. Philipp, B. 14, 2643).

Preparation. - A saturated solution of iodine in ether is heated to boiling with excess of powdered As in a flask with an upright condenser; the liquid is filtered while hot; on cooling, well-formed crystals of AsI, are obtained (Wiggers, Michaelis's Lehrbuch der anorgan. Chem. [1881] 2, 462).

Properties and Reactions .- Lustrous, red, hexagonal, tables, a:c = 1:2.998 (details of cryst. form, v. Friedländer, Z. K. 3, 214). Soluble in alcohol, other, benzene, &c.; soluble in mooh water; soluble without change in alcohol; decomposed by a little water with formation of 1AsOI.3As O.24II.O; action of boiling il_O produces AsOI.Ar, O_d which deposits on cooling (Wallace, P. M. [4] 17, 122).

Combinations .- 1. With iodides of the alkali metals to form very unstable compounds (Nicklès, C. R. 48, 237).-2. When ammonia is passed into a solution of AsI, in beuzene, a bulky white pp. of 2Asl, 9NH, is produced (Bamberger a. Philipp, B. 14, 2613).

Reactions .- I. Heated with oxygen, iodine is evolved and As O is produced .- 2. Heated with alcohol, ethylic iodide is formed (Bamberger a.

Philipp, B. 14, 2643).

II. ARSENIO DIIODIDE. AsI. (Bamberger a. Philipp, B. 14, 2643). Mol. w. unknown.

dissolved in CS, in an atmosphere of CO, and the liquid is allowed to deposit orystals. Properties .- Thin priems of a dark oherry-

red colour.

Reactions .- Very easily oxidised. Decomposed by water to AsI, and Ae; (3As I2= 2AsI, + As).

III. Arsenio Pentiodide. According to Sloan (C. N. 46, 194) a brown crystallino solid, containing As and I in the ratio As:5I, is obtained by heating As with a slight excess of I to 100° for some time in an atmosphere of CO,. The body is easily decomposed to AsI, and I, by heat

or by solution in ether or CS...
Arssnic, oxidos of. (In connection with these compounds, v. art. Oxides.) Arsonic forms two well-defined oxides, arsenious oxide As O., which has been gasified; and arsenic oxide, As2O3, which has not been gasified: both act as anhydrides; the acid corresponding to the former is not known, but many arsenites, MaAsOa, have been prepared. Three arsenic acids are known corresponding in composition to the three phosphoric acids, two of those exist only ac solids, they all readily lose water yielding the anhydride As2O2. The greyish film which forms on the surface of arsenic exposed to the air has been regarded as a cuboxide, but no definite proof of the existence of an oxide with less O than As,O, has been

I. ARSENIOUS OXIDE. As,O., Mol. w. 395.36. (Arsenic trioxide, White arsenic, Arsenious anhydride, Arsenious acid.) Vitreous As,O, melts under pressure, crystallino vaporises without melting (Wöhler, Gm. 4, 255). S.G. vitreous 3.698 to 3.738; oryst. 3.85 to 4.15 (v. Claudet, C. J. [2] 6,179, and Groth, P. 137, 426). V.D. 198 (at white heat, V. Meyer, B. 12, 1117). C.E. (cubic at 40°) ·00012378 (Fizcau, C. R. 62, 1133). S. (13°) vitreous, 4; cryst. 1·2 to 1·3. S. (100°) 11 (Bussy, A. 64, 286). S. (15° standing for somo days) cryst. 28; vitreous 92, S. (saturated at ays) cryst. '28; vitreous '92, S. (saturated at 100° and then cooled to 15°) cryst. 2'18; vitreous, 3'33 (Buchner, J. Ph. [3] 1, 421). S. (alcohol, 15°) cryst. '25; vitreous, 1'06 (Girardin, J. Ph. [3] 46, 269). S. (ether) 0. H.F. 309,3'10; H.F. in aqueous solution, 294,240; [As'O',Aq] = -15,100 (Th. 2, 236). S.H. cryst. '1279. S.V.S. vitreous, 106'3, cryst. 98'9. Occurrence.-Native, as the mineral Arsenite

(or Arsenolite); whenever arsenic volatilises in contact with air, or arsenio-containing minerals are heated in air.

Preparaten.-Obtained as a principal produot in the roasting of arsenical pyrites, and as a eccondary product in the reasting of arsenical oree of Sn. Co. Ni. or Ag. The exide ie condensed in chambers, and purified by resublimation. Extremely poisonous; doses of 0.6 grain are usually fatal (but v. Roscoe 'On the alleged Practice of Arsenic-caling in Styria,' Mem. of Lit. and Phil. Soc. of Manchester, 1860).

Properties.—Exists in three forms, amor-

phous, regular octahedra, and trimetric prisms (a:b:c:=3758:1:35); the first passes slowly into the second form on keeping; the third is obtained under special conditions (v. infra). The change from amorphous to octahedral arsenious oxide is attended with production of 5,300 gramunits of heat, and that from amorphous to prismatic with 24,950 units per As,O grams (Favre;

Troost and Hautefeuille, C. R. 69, 48). Amorphous arseuious oxide is produced by con-densing the vapour ou a surface slightly cooler than the temporature of volatilisation of the oxide; it is a transparent glass-liko eolid which gradually becomes opaque bocause of formation of octahedral crystals. The octahedral oxide is produced by cooling the vapour quickly; by crystallising either of the other forms from water; by treating the amorphous form with ammonia solution and washing with water. This form is obtained pure by fusing commercial arsenious oxide with carbonate of sodium and nitre, dissolving the arsenate of sodium so formed iu water, filtering from sodium antimonate which remains, and reducing with SO, solution. According to H. Roso (P. 35, 481) a solution of 2-3 parts amorphous As Os in 12 parts boiling conc. HClAq deposits crystals of the octahedral oxido when very slowly cooled, the formation of each crystal being accompanied by a flash of light; a similar solution of the crystalline oxide does not behave in this way. The trimotric prismatic variety of As,Os is obtained by saturating with As,O,, and then allowing to cool, a boiling solution of potash (Pastenr, C. R. 21, 474); Wöhler found this oxide in an oven in which arsenical ores had been reasted (P. 26, 177); Claudet found it native at San Domingos, in Portugal (C. J. [2] 6, 179); Kühn obtained it from a solution of Ag, AsO, in HNO, (J. 1852. 378; v. also Uhrich, J. 1858. 173); Scheurer-Kestner found it in pipes leading from the pyrites burners to the chambers of a sulphuric acid works (Bl. [2] 10, 414). The three forms of As,O, may be obtained, according to Debray (C. R. 58, 1209), by heating the oxide in a closed glass tube half immersed in sand, in an apright position, the lower part being at about 400?; on cooling, the lowest part of the tube contains amorphous, the middle part trimetric crystals, and the uppermost part octahedral crystals, of As Os. Arsenious oxide is iso-dimorphous with antimonious oxide. The vitreous (amorphous) variety may be fused before volatilising; the crystalline sublines without melting, even under pressure (Wöhler, Gm. 4, 255). The vapour is colourless and inodorous. The solubilities in water of the three forms are different (v. supra); long-continued contact with hot water decreases the solubility of the vitreous form inasmuch as it is thus changed to the octahedral form. Many acids dissolve As₄O_a, the vitreous more rapidly than the crystalline varieties (Bacaloglo, J. pr. 83, 111); from these solutions the oxide crystallises on cooling; in the case of HClAq some AsCl₃ remains in solution, tartaric acid is, however, said to form a salt analogous to tartar emetic. An aqueous colution of As O. slightly reddens litmus, but no acid has been obtained in definite form. The oxide must be regarded as a feebly acid-forming oxide possessing at the same time salt-forming tendencies (v. Arsenious acid and Arsenires; and also, infra, Reactions, especially Nos. 2, 10, and 11; and Combinations, No. 2).

Reactions. - Arsenioue oxide acts both as a roducing and an oxidising agent; it deoxidises nitrio, manganic, chromic, hypochlorous, acids, &c., with formation of arsenic acid; it oxidises carbon, sulpliur, phosphorus, hydrogen, sodium, ARSENIC.

potassium, carbon monoxide, potassium oyanide &o., when heated with these bodies. The oxide in solution is exidised to arsenic exide by chlorine, bromins, or iodine, in presence of alkaline bicarbonates; on this fact is based the use of As,O,Aq in volumetric analysis. 1. Chlorine passed over the dry oxide at a moderate temperature forms AsCl, (Weber, P. 112, 619); in solution chlorine produces arsenie acid and HClAq; iodine and bromine act similarly in presence of alkaline bicarbonates.—2. Hydrochloric acid forms seme AsCl₃, but in presence of HNO₃ or KClO₃ only As, O, is produced. -3. Free oxygen does not oxidise As O at ordinary temperatures, but if a plate of Pt is partly immersed in solution of As O in HClAq the exygen coming off from the Pt produces As.O. (Berthelot, C. R. 84, 1408).—4. An aqueous solution of As₄O₃ heated to 200° with phosphorus yields phosphide of arsenic (Oppenheim, Bl. [2] I, I63).—5. Na amalgam appears to reduce As O Aq with production of a selution which acts as an energetic reducer (Fremy, C.R. 70,61).-6. Phosphorus trichloride reacts at I10°-130° according to the equation $5As_1O_0Aq + 12PCl_3 = 2As_1 + 6P_2O_5Aq + 12AsCl_3Aq$ (Michaelis, J. Z. 6, 239). -7. Phosphorus penta-chloride produces AsCl, and POCl, (Hurtzig a. Genther, A. 111, 159). -8. When As O Aq is shaken with ammonia solution, an unstable compound is formed (De Luynes, C. R. 44, 1353).—9. Phosphorous and hypophosphorous acids precipitate arsenic and produce phosphoric acid.-10. Sulphydric acid (II.S) passed into an aqueous solution of As O, forms As S, which is ppd. on addition of an acid (v. Arsentous aut-PHIDE),-11. Ammonium hydrogen tartrate solution is said to dissolve As O,; on cooling, crystals of a double salt isomorphous with tartar emetic separate out (Marignac, Ann. M. [5] 15, 288). Pelouze (A. Ch. [3] 6, 63) describes a somewhat similar salt containing K in place of NH, Neither turtrate has, however, been satis-factorily examined.—12. Heated in a tube with a dry acctate, caeodyl oxide, As (CH1),O, is produced. -13. Heated with dry alkaline earth oxides or with dry carbonates of the fixed alkalis, an arsenate of the metal is produced along with arsenic which sublimes .- 14. The oxide dissolves in hot solutions of the alkalis but most of it pps. again on cooling in the air (v. Arsenious acid and arsenites).—15. The higher oxidised compounds of many metals are reduced by As,O.; thus CuO is reduced to Cu.O, in presence of alkali .- 16. Many metals, e.g. zinc, reduce acid solutions of As,O, with ppu. of As and formation of AslIn-17. As O Aq is reduced by a Cu-Zn couple with formation of AsH, (Gladstone a. Tribe, C. J. 33, 306).—18. Palladium or platinum charged with hydrogen separates As from As O Aq without fermation of AsH, (Gladstone, L.c.) -19. Some metallic salts which act as reducing agents convert the exide into arsenie; e.g. SnCl₂ Aq produces SnCl₄, As, and AsH₃ (Kessler, J. 1861, 265).—20. Freshly ppd. ferric hydrate reacts with As O Aq or with alkaline arsenites to form an insoluble compound, pro-bably arsenite of iron; on this fact is based the use of ferric hydrate as an antidete in eases of arsenie poisoning (v. Bunsen and Berthold, Das Eisenoxydhydrat, ein Gegengift der arsenigen Saure, Göttingen, 1834).

Combinations .- 1. Fused with arsenic oxide the body As,O. As.O. is probably produced (Bloxam, C. J. 18, 62). Other compounds of As,O, and As,O, are obtained by oxidising As,O. by warm HNO, Aq (v. Jely, C. R. 100, 1221).-2. Dissolves in fuming sulphuric acid; on evaporation yields needlo-shaped crystals of As,O. 4SO. which are decomposed by water (Schultz-Shellae, B. 4, 109, gives the formula As₂(SO₁)₃SO₃). A compound of As₂O₃ and SO₃ was obtained by Schafhäutl (B. J. 22, II3), in the fumes from copper-smelting works in Wales; and by Reich in a canal which carried off the sulphurous acid from a pyrites work near Freiberg (J. pr. 90, 176).—3. By cooling mixed hot aqueous solutions of KI, KBr, or KCl, and KASO, the compounds As,O_e,KI, As,O_e,KBr, and As,O_e,KCl, are obtained (Rüdorff, B. 18, 1441; v. also Schiff a. Sestini, A. 228, 72). Rüdorff (B. I9, 2678) also describes NII,LAs,O_e, NH,Br.As,O_e and 2NH Cl.As.O. Forms a complex series of compounds with MeO, and WO, and various bases (v. Gibbs, Am. 7, 209 a. 313; C. N. 48,

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Il. Arsenio oxide. As,O,. Mol. w. unknown; not less than that represented by formula. (Arsenic pentoxide, Arsenic anhydride.) S.G. 3.734 (fused oxide). S.V.S. 61 6. H.F. 219,100; H.F. in aqueons solution, 225,400. [As O'Aq,O'] = 78,350 (Th. 2, 236).

Preparation.—Not produced by heating arsenie in air or oxygen. If arsenic or arsenious oxide is digested with HNO3Aq, or with a mixture of 1 part HClAq and 12 parts HNO, Aq in a retert, or if chlorine is led into a warm solution of As O, arsenic acid, H, AsO, is produced and may be obtained as crystals by cooling a con-centrated solution. When this acid is heated to

low reduess the anhydrido As₂O₅ is produced.

Properties.—A white solid which slowly absorbs moisture from the air with formation of H3AsO4Aq. Slowly but completely dissolves in water forming H₂AsO₄Aq. Heated above low redness it yields As₁O₆ and O.

Reactions .- I. Heated with charcoal, many metals, or potassium cyanide, it yields As.-2. Heated with cone. hydrochloric acid it yields AsCl,; with HCl gas even in the cold the same product is obtained (Souchay, Fr. I, 189; Mayrhofer, A. 158, 326).—3. Reacts with phosphorus pentachloride thus: As.O., +5PCl₂ = 5POCl₃ + 2Cl₂ + 2AsCl₃ (Hurtzig a. Genthor, A. 3, 159) .- 4. According to Michaelis (J. Z. 6, 239), the exide is not acted on by POCl, even at 200° .- 5. Reduced in aqueous solution by nuscent hydrogen with formation of AsH2; but if chlorides are present only a trace of AsH, is produced according to Bloxam (C. J. 15, 56) .- 6. Stannous chloride, in the cold, produces stannous pyrearsenate and arsenite, in warm solutions produces arsenie and AsH, (Schiff, J. 1861. 278; Kessler, ibid. 265).—7. With water it reacts to produce arsenic acid, H₃AsO₄Aq. Joly (C. R. 106, 1262) describes a hydrate As₂O₄AH₂O. Arsenie oxide reacts as a strongly acid-forming oxide and exhibits no tendency to form corresponding salts by reactions with acids (v. e.g. reaction with HClAq).—8. Forms a large series of compounds with MoO, or WO. and bases (v. Gibbs, Am. 7, 209 a. 313; O. N. 48, 155).

Arsenia, exybromides of. AsOBr (Bromarservious acid); and ?As,O.Br. Mol. w. unknown, not less than represented by above formulæ.

Formation.—AsOBr is produced by the action

of H₂O in limited quantity on AsBr₃.

Preparation.—Arsenious oxide is dissolved in molten AsBr₃; the dark viscid liquid which results is distilled till it becomes rather thick, and is then cooled to 150° whereat it separates into two layers, the upper of which contains the oxybromide AsOBr, and the lower probably contains the other oxybromido A.O.Br. (Wallace, P. M. [4] 17, 261).

Properties. - Brown, waxy, solid. Decom-

posed by heat to AsBr, and As,O.

Combinations. - With water; a hydrate of arsenie oxybromide, 2AsOBr.3H.O is obtained as thin white pearly crystals by placing a cold concentrated aqueous solution of AsBr, containing HBr, ever sulphuric acid (Wallace, Lc.). If the solution of AsBr, in HBr is boiled, another compound, said to have the composition 4AsBr, 11As, Og. 2411,O, separates out (Wallace, l.c.).

Arsenic, oxychlorides of. AsOCl (Chlorarsenious acid); and As,O,Cl. Mol. ws. unknown, not less than represented by above formulæ.

Formation. - When AsCl, is mixed with less than sufficient water to completely decompose it,

AsOCI is formed.

Preparation of AsOCl.—By distilling until frothing begins the liquid obtained (a) by dissolving As O, in boiling AsCl, in the proportion As4O6: 2AsCla, or (b) by leading dry IlCl gas over dry warm As, O, until almost the whole of the latter has been changed to AsCla, and allow-

Properties of AsOCL.—Obtained as above, it is a hard, translucent, slightly-funning solid which slowly absorbs oxygen from the air (Wallace, P. M. [4] 16, 358; Hurtzig a. Geuther,

A. 111, 172).

Combinations.-1. A selution of AsCl₃ in cone. HClAq mixed with solid ammonium chloride, and allowed to stand, deposits crystals of AsOCl.11,O, but after some days white fibrous needles are formed, which, when dried over H.SO4, have the composition AsOCl.2NH4Cl.-2. With water; a hydrate of AsOCI, having the composition AsOC!.11,O(=As(O11),Cl) is obtained by adding water to AsCl₃ in about the proportion 8H₂O:AsCl₃ and allowing to stand for somo days. The hydrate forms small star-like crystals (Wallace, L.c.)

As,O,Cl is said to be obtained, as a hard, glass-like solid, when AsOCl is heated until As Oa begins to sublime from it (about 218°)

(Wallace, I.c.).

Arssnio, oxylodids of. AsOI.As,O. Mol. w. unknown. Produced in thin pearly lamine, according to Wallace (P. M. [4] 17, 122) by slowly cooling a hot cone. solution of AsI, in H.O. drying between filter paper, and then over

Arsenic, pantafinorids of ; double compounds containing. No gaseous compound of arsenic of the type AsX,, where X is a monovalent atom or atomic group, has yet been obtained. Selid compounds are, however, known, one of the constituents of which seems to be the group AsF. The following are described by Marignac to As and S by a powerful battery (Lapschin-

(A. 145, 287):—1. Potassic-arrenic fluoride, 2(KF.AsF.).H.O; formed in well-developed rhombic prisms by dissolving potassium arrecnate in much hydrofluorio acid .- 2. Potassicarsenic oxyfluoride, KF.AsOF, H.O; formed in acute rhonibic plates by repeated evaporation of the solution from which compound No. 1 is obtained, or by dissolving potassium arsenate in a small quantity of HFAq. - 3. Dipotassic-arsenic fluoride. 2KF.AsF., II.O; large, lustrous, rhombio prisms, obtained by adding KFAq to a solution in HFAq of either of the preceding salts, and evaporating .- 4. The double salt 4KF.AsF₃,AsOF₃.3H₂O is said to be produced when a solution in HFAq of salt No. 3 is repeatedly evaporated.

Arssnic, phosphide of, v. Arsenio, Combina.

tions. No. 8.

Arsenic, salanidos of, and Saleno-anlphidas

of, v. Arsenic, Combinations, No. 7.

Arsenic, snlphides of. (In connection with these compounds v. art. Sulphides.) Three sulphides of arsenie are known; As, S, As, S, and As₂S₅. None of these has been gasified, hence the formula do not necessarily represent molecules of the compounds. As S and As, S, occur native as Realgar and Orpiment respectively. The two sulphides As, S, and As S, are salt-forming; they dissolve in alkali sulphides with production of thio-arsonites M3AsS3 &c., or thio-arsenates M3AsS, &c. (v. infra). The disulphide, As, S,, is not salt-forming; Berzelius's statement that it combines with various metallie sulphides has been shown to be erroneous (Nilson, B. 4, 989).

I. ARSENIC DISULPHIDE. As₂S₂ (Realgar. Red orpiment, Ruby sulphur). S.G. 34-3-6. H. 15-2. Mol. w. unknown. S.V.S. 61-1.

Occurrence .- Native, as Realgar, accompany-

ing ores of silver and lead, &c.

Preparation .- 1. By heating tegether As and S, or As, S, with As, in the proper prepertions. -2. By heating As O, with S, in the proportion As Os:78, repeatedly subliming the mass from end to end of a glass tube in a stroam of CO, (Nilson, J. pr. [2] 8, 89).—3. By heating As S₃ with NallCO₃Aq in a closed tube to 150°; crystals are thus obtained | mm. long (Sénarmont, A. Ch. [3] 82, 129) .- 4. On the large scale, impure, containing As O. (Hausmann, A. 74, 196), by subliming a mixture of arsenical pyrites and iron pyrites.

Properties. - Occurs native in monoclinio prisms, a:b:c=1.32:1:4866; $o=85^{\circ}16'$; orangs. red, more or less translucent, resineus lustre, cenchoidal fracture. Pure As, S, is transparent, ruby colour, easily fusible, and crystalline after fusion; it burns in the air with a blue flame forming SO. and As,O6. It is used as a pigment, also in pyro-

techny.

Reactions.—1. Nitric acid oxidises As, S, to H, AsO, H, SO, Aq, and S.—2. Heated in a current of chlorine, S.Cl, and AsCl, are produced (Nilson, J. pr. [2] 12, 295; 13, 1).—3. Heated in hydrogen, As and H.S are formed (N.) .- 4. Solution of potash partially dissolves As.S., with formation of As.S. which then forms KAsS.Aq, and production of As (N.).— 5. It is slightly soluble in solutions of the alkali-metal sulphides .- 6. It is electrolysed Tichanowitsch, O. C. [2] 6, 613).—7. Heated with iodine, the compound As₂S₂.AsI₃(-AsSI) is produced (Schneider, J. pr. [2] 28, 486).

II. Arsenio trisulphine. As, S, (Arsenious sulphide, Sulpharsenious anhydride, Orpiment, Yellow sulphide of arsenie). Mol. w. unknown. S.G. 346-348. S.V.S. 709.

Occurrence.-Native as Orpiment.

Formation.-1. By heating to 70°-80° a selution of Na₂CO₃ saturated with As₂S₃ (Nilson, J. pr. [2] 12, 295; 13, 1).—2. Impure, commercial, by subliming together 7 parts powdered As O with 1 part S.

Preparation.-1. By subliming together As and S in the proper proportions. - 2. By saturating As,O.Aq with H.S a little HClAq being added. If no mineral acid is added the As, S, produced remains in solution in a colloidal form

(Schneider, J. pr. [2] 25, 431).

Properties. - Occurs native in trimetric prisms (a:b:c = 0.03:1:0.074) translucent, lemon-or slightly orange-yellow. Prepared in the wet way it forms a lemon-yellow powder which becomes darker when heated. Melts easily and volatilises at a higher temperature. When H.S is passed into As O, Aq, As S, is formed but remains in solution in colloidal form; a saturated solution contains 34 16 p.c. As S; it is slowly decomposed on standing, but may be boiled without precipitation of As,S,; bone char removes all the As.S3 from solution; most acids and many salts precipitate As S3 (Schneider, J. pr. [2] 25, 431). Used as a pigment, also as a reducing agent in dycing, also as a depi-

latory. Reactions .- 1. Long-continued action of hot water produces H.S and As, O, Aq according to Field (C. N. 3, 114) .- - 2. Dilute acids do not act on As,S3; conc. HClAq produces AsCla; conc. HNO₃Aq produces H₂SO₄Aq, S, and H₃AsO₄Aq. 3. Fused with potassium-hydrogen sulphate, SO₂ is evolved, and KHAsO₂ and K₂SO₄ remain. Chlorine acts readily, a brown liquid is formed said to be a chlorosulphide of As (II. Rose); heated with chlorine, AsCl, is produced (Ludwig, Ar. Ph. 97, 23) .- 5. Passed over hot iron, silver, &o., sulphide of the metal is formed, and arsenic which partially alloys with the metal. - 6. Passed over red-hot lime, sulphide and arsenate of calcium, and arsenie are produced. - 7. Heated with sodium or potassium carbonate, a mirror of As is obtained, along with arsenate and thioarsenato of the alkali metal; if the mixture is heated in hydrogen the arsenate is reduced (Rose, P. 90, 565). -8. Heated with an alkaline carbonate and charcoal or potassium cyanide, a mirror of As is obtained; according to Fresenius (4. 49, 287), the whole of the As in the As S, is thus obtained; according to Rose (Ph. C. 1853. 594), some of the As forms thio arsenate (KCNS being also produced) which is not reduced. mirror of As is obtained (Rose) if As S, is mixed with excess of S and heated with KCN; the presence of an easily reduced metal is also said to prevent the formation of As, because the As alloys with the metal. If the mixture of As, S, with Na₂CO₃ (or K₂CO₃) and KCN is heated in hydrogen, the whole of the arsenic is obtained as metal (comp. Rose, P. 90, 565, with Nilson, A. 49, 287) .- 9. As S. readily dissolves in cold

aqueous potash, soda, or ammonia, forming an arsenite and a thio-arsenite; thus :

 $2As_2S_3 + 4KOHAq = KAsO_2Aq + 3KAsS_2Aq + 2H_2O;$

on adding an acid to the solution the whole of the As is precipitated as As, S,; thus:

 $\begin{aligned} \mathbf{KAsO.Aq} + 3\mathbf{KAsS.Aq} + 41\mathbf{IClAq} = \\ 4\mathbf{KClAq} + 2\mathbf{As.S.} + 211.0. \end{aligned}$

If oxide of Ag or Pb is added to a solution of As,S, in NII,Aq and the solution is boiled, the whole of the S is precipitated as Ag.S or PhS, and Ag or Pb arsenite remains in solution. 10. When As, S3 is boiled with a solution of sodium or potassium carbonate, As S, is precipitated, CO2 and H2S are evolved, and the solution contains the following salts, Na,S,3As,S₃; Na,O,2As,S₃O₂, Na,AsS,; Na,HASO,; NaHCO₃; (Nilson, J. pr. [2] 14, 1, 145).—11. As,S₃ is easily soluble in a hot solution of polassium-hydrogen sulphite; thus, 2As,S₃ + 16KHSO,Aq = $4KAsO_{2}Aq + 6K_{2}S_{2}O_{3}Aq + 7SO_{2}Aq + 5S + 8H_{2}O_{3}$

Combinations .- As S acts as a salt-forming sulphide, or anhydride of a thio-acid; it combines with the sulphides of the alkali and alkaline earth metals, and with some metallic hydrosulphides, to form thio-arsenites (q.v. under ARSENIC, THIO-ACIDS OF). The following are the typical reactions:

 $\begin{array}{l} {\bf 1.} \ \ {\bf As_{a}S_{3}+K_{a}SAq} = {\bf 2(AsS.SKAq)}. \\ {\bf 2.} \ \ {\bf As_{a}S_{3}+6NH_{a}IISAq} = {\bf 2(As.(SNII_{a})_{a}Aq)} + {\bf 3H_{a}S.} \end{array}$

3. As S₃ + 2(NH₁) SAq = As S(SNH₁) Aq. 11I.—Arsenic Pentasulphide. As S sulphide of Arsenic). Mol. w. unknown.

Preparation.-1. By melting As with con-

siderable excess of S, a thin, transparent, liquid is obtained which solidifies to an elastic mass, and after some time becomes hard; if this hard solid is powdered and treated with NH,Aq a solution of As, S, is obtained from which the sulphide is thrown down on addition of HClAq (Gélis, A. Ch. [4] 30, 114).-2. A solution of Na S is digested with As S and enough S to form As.S., on evaporating and cooling largo erystals of 2Na.AsS., 1511.O are obtained (Rammelsberg, P. 52, 249; 90, 40); when HClAq is added to a solution of this salt, As S, is precipitated and H.S is evolved (Fuchs, Fr. 1, 189; Flückiger, Vierteljahrsschr. pr. Pharm. 12, 330; Eekert, ibid. 13, 357). The product of the action of ILS on HaAsO4Aq is not As.S5, as was once supposed, but is a mixture of As, Sa and S

(21I₃AsO₂Aq + 2H₂S = As₂O₃Aq + 5H₂O + S₃; As₂O₃Aq + 3H₂S = As₂S₃ + 3H₂O (200 Ludwig, Ar. Ph. [2] 97, 32; also H. Rose, P. 107, 186). Properties. - A yellow powder, easily fusible;

may be sublimed in a stream of a gas which does not act on it.

Reactions.-1. Heated in a stream of hydrogen, it is reduced to metallie As, and H2S. 2. Dissolves easily in ammonia, potash, and soda solutions, with production of thio-arsonate, and arsenate, of the alkali metal .- 3. Dissolves easily in solutions of alkali sulphides, forming thio arsenates. The sulphide As, S, behaves az a salt-forming compound, or as the anhydride of thio-arsenic acid; the salts which are generally formed directly from it are pyro-thio arsenates M.As.S.; these yield two other series of salts, viz. ortho-thio-arsenates M, Asil, and meta-thio arsenates MASS, (v. ARSENIC, THIO-ACIDS OF).

ARSENIC.

Arsenic, sulphe-acids of, v. ARSENIC THIO-

Arssnio, sulpho- (or thio-) bromide of. S.Br. (=AsSBr.SBr.). Mol. w. unknown. AsS,Br, (-AsSBr.SBr,). Dark red crystals deposited at -18° [-17°]. on addition of a small quantity of powdered As to a solution of S in Br in ratio S:Br2; decomposed by water into As,O,Aq, IIBrAq, and 8 (Hannay, C. J. 33, 291).

Arsenie, sulpho- (or thio-) iodide of. AsSI. Said to be formed by the mutual action of As S2

and I (Schneider, J. pr. [2] 23, 486).

Arsenic, sulphydrates (or hydrosulphides) of. Only one compound As, S, and H is definitely known, AsS(SH)3; v. Thio arsenic acids under ARSENIC, THIO-ACIDS OF (v. also the art. Hydro-SULP: QDES).

Arsenic, tellurides of, v. Arsenic, Combina-

tions, No. 6.

Arsenic, thio acids of. (In connection with these compounds v. the art. Hydrosulphides.) Arsenious sulphide, As.S., dissolves in alkalis or alkali sulphides to form salts, and from these other salts are obtained by double decomposition. The sulphide As, S, may be regarded as the anhydride of three thio-acids AsS.SH, As(S11), and As, S(SII), corresponding to the three hypothetical oxy-acids (v. Arsentous acro); none of theso acids is known, all attempts to prepare them liaving resulted only in the production of As.S. and H.S, but thio- or sulph- arsenites are known belonging to the three types, MAsS, MASS, and MAs.S. The more important of these salts are described below. Arsenic pentasulphide, As,S,, dissolves in alkalis and alkali sulphides to form salts from which other salts are obtained (v. infra). According to Nilson (J. pr. [2] 14, 1, 145) the pp. obtained by adding dilute HClAq to a solution of Na, AsS, (v. Arsknic PENTASULPHIER, Preparation of) has the composition of ortho-thio-arsenic acid HASS, [= AsS(SII), no other thio-arsenic acid is known, but the salts may be divided into three classes, analogous to the arsenates, viz.: pyro-thio-arsenates M4As,S, (hypothetical acid - H4As,S,) meta-thio-arsenates MAsS, (hypothetical acid RASS,), and ortho-thio-arsenates MaAsS, (acid (?) H,AsS.).

I. Thio-arsenites. As already stated, no thio-arsenious acid is known. The salts which have been examined belong for the most part to the type M4As.S5; they are produced either by the direct union of As, S, with metallic hydrosulphides, e.g. As. S., or by ordinary double decomposition of (NH,)As. S.Aq by solutions of motallio salts, e.g. Pb.As.S.. A few salts belonging to the forms MASS, and MaASS, are also known, e.g. KASS, and KaSS, they are formed by the action of alkali sulphides on AsS, (comp. reactions given for Ausenic Trisulphide, p. 315). The thio-arsenites of the alkali and alkaline cartle metals and of magnesium are soluble in water, but the solutions are decomposed on boiling, the others are insoluble in water. Most of these salts give off all their sulphur when strongly heated out of contact with air. These salts have been chiefly investigated by Berzelius (v. Gm. 4, 275).

Only those salts which have been fairly satisfactorily examined are mentioned in the

following brief account :-

Annonium thio-arsenites. (NH₄),As₂S₂ is obtained by dissoiving As₂S₂ in (NH₄)₂SAq and adding alcohol; if NH₄HSAq is added before precipitating by alcohol the salt obtained has the composition (NH,),AsS,

Barium thio-arsenites. Ba2As2S3 is obtained as a pasty brownish-red very soluble mass by digesting As, S, with BaS, H, Aq; from the solu-

tion alcohol throws down Ba₃(AsS₃)₂.

Calcium thio-arsenites. The salt Ca₃(AsS₃)₂. is obtained as crystals by digesting As, S, with milk of lime and allowing the solution to evaporate; from the brownish mother-liquor alcohol precipitates white Ca₃(AsS₃)₂.15H₂O.

Lithium thio-arsenites. Closely resemble

the potassium calts (q. v.).

Potassium thio-arsenites. The salt KAsS, may be obtained in solution by dissolving As, S, in K2SAq, but this solution decomposes on ovaporation; in the solid form by heating KAsS, or by fusing As₂S₃ with K₂CO₃. By adding alcohol to a solution of As₂S₃ in K₂SAq, a white ppoof K₃AsS₃ is obtained. All these salts readily undergo change in aqueous solutions. Berzelins describes several other more or less indefinite bodies as potassium thio-arsenites.

Sodium thio-arsenites. Closely analogous

to the potassium salts.

The following thie-arsenites seem also to exist; they are generally obtained from

CMB, As, S, Aq by double decomposition:—
(BiS),As,S, S, Ce,As,S, Cd,As,S,; Cd,As,S,; Cd,As,S,; Mg,As,S,; M Sn.As₂S₅, SnAs S,; Pt.As.S.; Ag, As, S,; (US), As, S,; Zn, As, S,. Thio-arsenites of chromium, molybdenum, and zirconium, seem also to exist.

II. THIO-ARSENATES. As already stated, it is probable that ortho-thio-arsenic acid HASS. has been prepared. The thio-arsenates may be divided into three classes, of which the three potassium salts are representatives: K, As, S,, K, AsS, and KAsS, Tho thio arsenates are obtained: 1. By digesting As. S. with solutions of the alkali sulphides, on cooling some As.S. is precipitated.—2. By dissolving As S, in solutions of alkali-polysulphides. - 3. By precipitating solutions of arsenates by H2S, or by (NH1)2SAq; in the latter case the liquids must be boiled to remove NH₃.—4. By fusing As₂S₃ with alkali curbonates.—5. By dissolving As₂S₃ in KOHAq or NaOH aq; arsenate is formed as well as thic arsenate. The thio-arsenates of the alkali metals are yellow or red, very soluble in water, crystallisable, fairly stable, compounds; their aqueons solutions are slowly decomposed by exposure to air. The other thio arsenates are more easily decomposed; those of the heavy metals are insoluble in water; they are best prepared by de-composing the solution of an alkali thio-arsenate by a solution of a salt of the metal. Soluble thio-arsenates are decomposed by HClAq with precipitation of As, S,. The salts obtained by the methods cnumerated are usually pyro-thioarsenates MAS2S7; the meta- and ortho- salts are produced from these, very frequently by the action of alcohol on their solutions; alcohol usually precipitates an ortho- salt and leaves a meta-salt in solution. The ortho-salts are froquently crystalline; most of the others are

amorphous. Heated in absence of air, most thio-arsenates yield thio-arsenites, and then As S, which sublimes, and a metallic sulphide which remains; some, however, are unchanged by heat alone, e.g. M, AsS, where M = Li, K, Na. Heated in air, the thio-arsenates, as a class, givs off As₂S₃ and As₄O₄, and leave a sulphate in tho cases of alkaline salts, or an oxide in the cases of salts of heavy metals. The thio arsonates have been chiefly investigated by Berzelius (v. Gm. 4, 275); also by Nilson (J. pr. [2] 12, 295; 13,1).
The following are the thie-arsenates which

have been fairly well investigated:

Ammonium thio-arsenates. The pyro- salt (NH,), As, S, has not been obtained as a solid: a solution of As,S, in (NII,) SAq probably contains this salt, it is decomposed on evaporation; alcohol precipitates the ortho- salt (NH,), AsS, in white prismatic crystals, while the meta-salt NH AsS, remains in solution.

Barium thio-arsenates. A solution of BaHAsO, is decomposed by H.S, but the puro pyro-thio-arsenate, Ba.As, S., has not been obtained; this solution is decomposed by alcohol into Ba₂(AsS₃)₂ which precipitates, and Ba(AsS₃)₂

which remains in solution.

Magnesium thio arsenates. The pyro- salt Mg. As. S, is a yellow solid, very soluble in water; by adding Mg(SII), Aq to this solution until II.S ceases to come off, and evaporating in vacuo, crystals of Mg3(AsS4)2 are obtained; ulcohol

lecomposes this salt, dissolving out Mg_AS_S.

Potassium thio-arsenates. The pyro-salt,
K_AS_S, is best obtained by treating K_HASO,Aq with H.S and evaporating in vacuo; it forms a yellow viscid mass which liqueties on exposure to the air and then crystallises in rhombic plates. By adding alcohol to a cone, solution of this salt an oily liquid is obtained which crystallises when warmed giving KaAsS, and KAsS, remains in solution. A salt containing both sulphur and oxygen, AsSO.OK.H.O is described by Bouquet and Clocz (A. Ch. [3] 13, 44), produced by the action of H.S on cold saturated KallAsO, Aq; it may perhaps be regarded as a double compound of the hypothetical oxysulphido As, S,O, with K,O, but the data ars

very meagre.

Sodium thio-arsenates. The ortho- sult 2Na, AsS, 15HO is obtained in large white, or yellowish, monoclinic prisms, by digesting Na2SAq with As2S2, or with As2S2 and sufficient S to form As2S2, and allowing to crystallise (Fresenius, Fr. 1, 192). The same salt is also obtained by decomposing Na_IIAsO,Aq by H.S, and adding alcohol to the solution; according to the conditions under which this liquid is allowed to crystallise, crystals of varying form and somewhat varying appearance are obtained (Berzelius). The crystals are not dehydrated in dry air, but when slowly heated the salt may be obtained without water of crystallisation. It is doubtful whether the meta- and pyro-thio-arsenates have been obtained; the solution from which the ortho- salt is thrown down by alcohol probably contains NaAsS3, and the solution before alcohol is added probably contains Na,As,S,. The double thio-arsenato Na_s(NH₄)_s(AsS₂), is also described by Berzelius

Bssides the above salts, the fellowing thicarssnates seem to have been obtained in fairly definito forms: (Ca,As,S.,, Ca,(AsS,),; Ce,As,S.,, Ce,(AsS,),, Ce,(AsS,),; Co,2As,S.,; Au,(As,S.),; Fe,(As,S.),; Fs,As,S.,; Pb,As,S., Pb,(AsS,),; Mn,As,S.,; Hg,As,S., Hg,As,S., Hg,As,S., Ag,As,S., S, Co,S., Thio-arsenates of Sb, Be, Bi, Cd, (US),As.S,. Cr. Li, Ni, Pt, Sr, Y, Zn, and Zr, probably exist.

Arsenic acid and Arssnates v. Ausenic, across

Arsenides. Binary compounds of arssnie with more positive elements, v. Arsenic, Com-

binations, No. 9.

Arsenious acids and Arsenites, v. Arsenio,

ARSENIC COMPOUNDS, ORGANIC. This article is devoted to compounds in whose molecules arsenic is supposed to be directly united to earbon. They are produced by distilling alkyl iodides with an alloy of arsenio with potassium or sodium (thus Mel gives As, Me, AsMe,, and AsMe,I—Cahonrs a. Riche, C. R. 39, 511), or by heating AsCl, with compounds of mercury with alkyls or aromatic radicles or by the action of sodium on a mixture of AsCl. and a haloid derivative. The methyl derivatives will be described first, followed by the methylethyl, ethyl, phenyl, and finally by the benzyl, derivatives. The nomenclature employed is somewhat different from that used for derivatives of nitrogen. Thus the radicles AsMe, AsMe, AsMe, and AsMe, are called methyl-arsino, dimethyl-arsine, tri-methyl-arsine, and tetramethyl arsonium respectively.

Msthyl-arsins dichloride AsMcCl.. (133°). At 40°-50° di-methyl arsino trichloride produces AsMeCl₂ thus: AsMc₂Cl₃ = McCl + AsMcCl₂. Liquid which does not fume. M. sol. water but not decomposed by it. It violently attacks the mucous membrane. At -10° absorbs Cl. to.m. ing AsMeCl, which at 0° splits up into MeCl and AsCl, (Bacyer, A. 107, 257).

Methyl-arsins di-iodids AsMcI, [c. 25°]. From the oxide, AsMeO, and III. From cacodyl and iodino (Cahours, C. R. 50, 1022). Yellow needles (from alcohol). Converted by H2S into AsMeS, and by HCl into AsMcCl.

Methyl-arsine sulphide AsMeS. From H2S and AsMeCl2. Plates (from alcohol). Insol. water. Pps. Ag, Cu, and Pb, as sulphides from their salts.

Methyl-arsins disulphide AsMeS... Formed by passing H.S into an aciditied solution of

methal-arsonic acid (G. Meyer, B. 16, 1440).

Methyl-arsins oxids AsMeO. [95°]. Formed by action of K₄CO₃ on the chloride AsMeCl₄.

Crystallises from CS₂ in irregular cubes, smells like Asa fortida. M. sol. cold, v. sol. hot, water; slightly volatile in vapour of water and alcohol; v. sol. aqueous acids forming neutral solutions.

Methans arsonic acid McAsO(OH)₂. From AsMcCl₂ and excess of moist Ag₂O. From ArMeO in aqueous solution by action of HgO. From aqueous sodium arsenito and MeI (M.). Large spear-shaped lamina composed of small

needles (from alcohol).

Salts. - BaA", 5H2O: ppd. as anhydrous rhombie crystals, by adding alcohol to aqueous solution; the crystals soon change to hydrated needles. Ag.A": nacroous crystals which sxplode above 100'.-- CaA" aq.

Tetra-methyl di-arsenide As, Me, Cacodyl. Alkarsin. Mcl. w. 210. [c. -6°]. (c. 170°).

V.D. 7-1 (air = 1).

Preparation.-By heating di-methyl-arsine chloride (cacodyl chloride) with zine at 100° in bnlbs filled with CO. (Hunsen, P. 40, 219; 42, 145; A. 87, 1; 42, 14; 46, 1).

Properties. — Stinking eil; heavier than

water. Takes fire in air or in chlorine. Reduces HgCl, to merenrous eliloride.

Reaction.—As, $Me_4 + 2MeI = AsMe_4I + AsMe_2I$

(Caliours, A. 122, 209).

Combinations.- When gradually mixed with air, chlorine, or bromine, it forms derivatives of cacodyl, behaving like a molecule of such a metal as potassium: $(AsMe_{\cdot})_2 + Cl_2 = 2(AsMe_{\cdot})Cl$; and $(AsMe_{\cdot})_2 + O = (AsMe_{\cdot})_2O$.

Tetra - methyl - di - arsine exide $(AsMe_{\cdot})_2O$.

Cacodyl oxide. Mel. w. 226. [o. -25°]. (120°). S.G. 12 1.462. V.D. 7.55 (calc. 7.83).

Formation .- Calet's fluid (Crell. N. Chem. Arch. I, 212), obtained by distilling KOAc with an equal weight of As_*O_{20} is cacodyl oxide mixed with some cacodyl. HgO converts both into cacodylic acid, whence a mixture of HgCl, and furning HCl forms cacodyl chloride. The latter is converted into cacedyl oxide by distilling with aqueeus potush in a current of CO2 (Baeyer, A. 107, 282): 2AsMe_Cl + 2KHO = $H_2O + 2KCl + (AsMe_2)O$.

Properties. Pungent, stinking oil. Slewly oxidises in air forming cacodylic acid. Acids

couvert it into salts of cacedyl.

Compounds. - Forms with HgCl2 a compound (AsMc2),O2HgCl2, crystallising in trimetric plates. S. 3-17 at 100°. Distilled with furning 11Cl this forms eacodyl chloride.— (AsMe₂)₂O2HgBr₂.—(AsMe₂)₂O 3AgNO₃: explodes at 100°, — (AsMe_)₂OPtCl₂aq: red-brown pp. converted by KBr into (AsMe₂)₄OPtBr₂aq, and by KI into (AsMe.), OPtI,

Di-methyl-arsine chleride AsMe, Cl. (c. 100°). V.D. 4.56 (calo. 4.85). Obtained from cacodylio acid as abeve; or by action of chlorine-water on cacodyl.- Heavy oil; attacks the mucous mombrane; combines with Cl. forming AsMe, Cla.

Zn, Sn, and Fe liberate As, Me,

Compounds. — AsMe Cl CuCl (Bunsen), -

(AsMe, Cl), PtCl,.

Di-metbyl-arsine bremide AsMe.Br: yelloweil Di-methyl-arsino iodide AsMe.I (160°); oil (Cahours a. Riche, A. 92, 364).

Di-methyl arsine cyanide AsMe, Cy. [33°]. (140°). V.D. 442. Prisms. Excessively poisonous.

Di-methyl-arsine sulphide (AsMey) 28. Combines with S to form (AsMe2)2S. [50°].

Di-methyl-arsine fluoride AsMegF. Di-methyl-arsine trichloride AsMe, Cla.

Cacodyl trichloride. From PCl, and cacodylic acid; er from cacodyl chloride and Cl.

Reactions .- 1. At 50 ' it splits up as follows: AsMe₂Cl₃ = MeCl + AsMcCl₂ -2. With water it forme oacodylic acid.

Di-methyl-arsinic acid AsMo₂O(OH). Cacodylic acid. Mol. w. 138. [200°].

Formation.—From cacodyl and HgO in

presence of water. Properties. - Large prisms (from alcohol), without odour, but peisonons. V. eol. water, m. sol. alcohol, insel. ether. Not acted on by HNO, HCl, aqua regia, KMnO, or CrO.

Reactions.-1. H.PO, reduces it to escodyl oxide.—2. Aqueons H.S forms cacodyl sulphide.—3. An alcoholic sclution gives with alcoholic HgCl₂ a pp. of (AeMe₂)₂O₂Hg₂Cl₂.— 4. Cacodylatee are converted by dry H.S into thio-eacodylates; e.g. (AsMo,S_d),Pb.— AsMe₃S₃Cn.—(AsMc₂S₃)₃Sb.—(AsMc₂S₂)₃Bi.—AsMe₂S₂Au.
Salts.—Solublo in water, but amorphous.

AgA': needles. $-AgH_zA'_a$: needles. $-AgA'\hat{A}gNO_{so}$

Compounds .- HCl forms a crystalline compound (AsMe_O.H HCl) decomposed by water. This compound distilled in a current of HCl splits up thus: AsMc_O_H IICl + 2HCl =

**MSMCL2+ McCl + 2H.O. — IIA'HF: prisms.

**Tri-methyl arsine AsMc., Mol. w. 120.
(c. 100°). **Formation.—1. 2AsCl3+3ZnMe2— 3ZnCl, + 2AsMe, (Hofmann).—2. From AsMe, I and solid potaeli (Cahours, C. R. 49, 87).

Properties.—1. Combines directly

Cl., Br., I., S, and O.

Iodido AsMe, I. Splits np on distillation into MeI and AsMe, I, cacodyl iodide.—Oxide. AsMe,O: deliquescent erystals. - Sulphide AsMeaS: prisms (frem alcohel). - Bromide AsMe₃Br₂.

Tetra-methyl-arsoninm iodido AsMe, I.

Formation .- 1. From sodium arsenide and McI at 180°, and treating the product (AsMc, IASI, with KOH (Calieure, C. R. 36, 1001; A. 122, 192) Properties .- Plates (Irom alcohol mixed with MeI).

Combinations. — AsMe,II, — (AsMc,I),ZnI,

- (AsMe₁1)₂CdI₂.—AsMe₁IAsI₃.

Reactions.—1. With ZnMe₂ gives AsMe₃(?)
(Cahours).—2. KOII no action.—3. Meist Ag₂O gives AsMe,OII, deliquescent alkaline crystals .-4. Ag₂SO₄ gives crystalline (AsMe₄)₂SO₄.— 5. AgNO₃ forms crystalline AsMe₄NO₃.

Penta-methyl-arsenide AsMe, Frem AsMe,I

and ZnMo.. With iedinc forms MeI and AsMe.I; with HCl forms CH, and AsMe, Cl (Cahours). Di-methyl-ethyl-arsine. -- AsMe, Et. From

AsMe I and ZnEt Liquid (Cahours).

Methyl-di-ethyl-arsinc AsMeEt... AsMeEt. From

AsMel, and ZnEt. (Cahours).

Di-methyl-di-cthyl arsonium ealts.

Iedidc .- As Me, Et, I. From encodyl and EtI, thus: As,Me, + 2EtI = AsMe,Et,I + AsMe,Cl (Caheurs a. Riche, C. R. 39, 541).

Hydroxide: very deliquescent.

Chloride AsMe, Et, Cl: deliquescent neodles. Platino chleride (AsMe, Et, Cl), PtCl, Bremide AsMe, Et, Br: deliquescent.

Todide AsNe, Pt. I.: priems.

Periodide AsNe, Pt. I.: priems.

Nitrate AsNe, Et. No.; doliqueeent grains.

Sulphate (AsMe, Et., No.; cotahedra.

Ethyl arsine iodide AsEtl.. From AsEt. I

and I₂ (Cahours, C. R. 50, 1022; A. 116, 367). With moist Ag₂O it ferms the acid AeEtO(OH)₂.

Ethyl-arsine chloride AsEtCl, (156°). From HgEt, and AeCl, (La Coste, A. 208, 33). Liquid, m. sol. water.

Ethane arsenic acid EtAsO(OH). From the preceding by the action of diluted HNO. Small crystale (from alcohol). -Ag2A": pearly ecales.

Tstra-ethyl-di-arsenide As, Et, Mol. w. 266. (185°-190°). From an alloy of areenic and sodium on EtI (Landolt, A. 89, 319). Heavy stinking oil, takes fire in air. Reduces calts of silver and mercnry. Unites directly with sulphur and halogens, Alcoholic HgCl, gives a crystalline precipitate AsEt, Cl, 2Hg, O(?).

Iodide AsEt,I. (c. 230°). Oil. Di-ethyl-arsinic acid AsEt,O(OH). [190°] From AsEt, and HgO under water (Landolt, A. 92, 865). Large plates, soluble in water. Not attacked by HNO, or aqua regia.

Salts.-BaA',HA' 2aq. Very sol. in water,

difficultly sol. in alcohol. Tri-ethyl-arsine AsEt,. Mol. (140°-170°). S.G. 11 1·151. V.D. 5·28 (calc.

Š·62).

Formation.—I. From Ascl, and ZuEt, (Hofmann a. Cahours, C. R. 41, 831).—2. Together with As, Et, by the action of EtI on an alloy of arsonic and sodium .- 3. By distilling AsEt,I with solid potash (Landolt, A. 89, 322).

Properties. Oil of disagreeable odour. Funes strongly in air. Combines directly with non-metals. Does not reduce aumoniacal silver

nitrate (difference from As,Et,).

Combinations.—AsEt,Br.: deliquescent.— AsEt,I.. [160°]. (190°). (Cahours a. Riche, A. 92, 365).—AsEt,S. [c. 100°]. Prisms (from ether); pps. sulphides from solutions of metallic (AsEt) PtCl ... (AsEt,) PtCl ... (AsEt,) PdCl2 (Cahours a. Gal, C. R. 71, 208).-(AsEt,OAsEt,Cl,Hg,CL(?). — AsEt,AuCl. AsEt,PEt,(C,H,Br)Br.

Tri-ethyl-arsine oxide AsEt, O. Formed by exposure of an ethereal solution of AsEt, to the air. An oil, insoluble in acids, except IINO3.

Tetra-ethyl arsenium iodide AsEt, I.

Formation.—1. From AsEt, (Landolt, A. 89, 331) .- 2. Arsenic with EtI at 180° gives red needles of AsEt,IAsI, which is then boiled with potash (Cahours a. Riche, C. R. 39, 516).-3. An alloy of arsenic with Zn or Cd heated with Etl gives (AsEt,1), ZnI, or (AsEt,I)₂CdI₂; these are boiled with potash (Cahours, A. 122, 200).

Properties.-Needles, v. sol. water and

alcohol, insol. ether.

Reactions .- 1. With moist Ag.O, gives an alkaline hydrate. -2. Combines with I forming brown needles of AsEt, I.

Tetra-ethyl arsenium salts (Landolt, A.

92, 371).

Chloride AsEt,Cl 4aq: deliquescent crystals, insol. ether. (AsEt,Cl), (BiCl,), (Jörgensen, J. pr. [2] 3, 346).

Platino chloride (AsEt, Cl), PtCl,: sl. sol.

cold water.

Bromide AsEt, Br: deliquescent mass. ~ (AsEt,Br)3(BiCla);

Sulphate AsEt, SO, II: grains, v. sol. water

and alcohol, sl. sol. ether.

Breme - tetra - ethyl - arseninm hromide (CH.Br.CH.)AsEt,Br. From ethylene bromide and AsEt, at 50° (Hofmann, Pr. II, 62). Rhombie dodccahedra (from alcohol). V. sol. water, sl. sol. alcohol. Aqueous AgNO_a pps. half its bromine as AgBr.

Reactions .- I. With moist Ag. O it gives vinyl triethyl arsonium hydroxide, C.H.Aslit, (OII). 2. With AsEt, it gives As, (C, H,)Et, Br, ... 3. With ammonia at 100° it gives a compound NAs(C₂H₄)Et₃H₂Br₂. This compound and the preceding are converted by Ag₂O into oxides and thence into platinochlorides (e.g., MAs(C,H,)Et,H,Cl,PtCl,) and other salts.

4. AuCl. gives crystals of AsEt, AnOi .- 5. PtOL gives crystals of As, Et, Pt (Hofmann, A. 103, 857).

Di-methyl-di-isoamyl-arsonium icdide

AsMe2(C,H11),I. From cacodyl and iso-amyl iodide at 180°, as follows: As.Me, +20,II, I = AsMe₂(C₅H₁₁)₂I + AsMe₂I (Cahours a. Riche).

Tri-propyl-arsine AsPr. At I80°, arsenic combines with PrI forming AsPr, IAsI,. Distilled with solid potash, this gives Asl'r, (Cahours, C. R. 76, 1383). Arsenie acts similarly on isobutyl iodide at I80° (Caliours, C. R. 77, 1406). Calcium butyrato distilled with As,O. gives a distillate resembling Cadet's liquid, probably containing the propyl homologues of eacodyl compounds (Wöhler, A. 68, 127). Potassium valerate distilled with As.O. appears similarly to give 'butyl-cacodyl' derivatives (Gibbs, Am. S. [2] 15, 118).

AROMATIC DERIVATIVES.

Literature.-Michaelis, A. 201, 184; 207, 195; 208, 1; 233, 60; B. 8, 1316; 9, 1566; 10, 622; II, 1883; I3, 2176; 14, 912; 15, 1952, 2876; 18, 42; La Coste, A. 181, 1; 208, I.

Di - phenyl - di - arscnide C. II. As: As.C.II. Arseno-benzene. [196°]. Prepared by reduction (best with phosphorous acid) of an alcoholic solution of phenyl-arsine oxido (Michaelis a, Schulte, B. 11, 912; 15, 1952). Yellowish needles. Sol. benzene, chloroform, and CS,; sl. sol. alcohol, insol. water and other. On heating it gives triphenyl-arsine and arsenie.

Reactions. - I. Hented with 1 mol. of sulphur phenyl-arsine sulphide is formed, with more sulphur, phenyl sulphide and As₂S₃. - 2. Alcoholic NH, HS reduces it on heating to benzeno, As, S, and As; III acts in a similar manner. -3. On oxidation it gives benzene-arsonic acid. -4. Com-

bines directly with halogens.

Di-iodide. - Ph.Asl.Asl.Ph. Yellow needles. Very unstable. Prepared by reduction of phenylarsine iodide (which is formed by dissolving phenyl-arsine oxide in 111).

Phenyl-arsine chloride PhAsCl. (c. 253°). Obtained in theoretical quantity by heating AsOl (800g.) with HgPh, (70 g.). Colourless liquid with unpleasant odonr; insol. water, sol. KOHAq.

Phenyl-arsine tetra-chleride PhasCl, [45° Formed by passing Cl into the preceding at 0°. Yellow needles, fuming in moist air; readily decomposed into Cl, and PhAsCl,; when heated at 150° it gives C. H. Cl and AsCl.

Phonyl-arsine hremide PhasBr.

S.G. 15 2 10. Colourless liquid formed by the action of conc. IIBr upon PhAsO. Gives with

bromine AsBr, and PhBr.

Phenyl-arsine oxide PbAsI₂. Oil. Phenyl-arsine oxide PbAsO. [120°]. Formed by treating PhAsCl, with Na, CO,. Crystals (from ulcohol); smells like anise; insol. water; sl. sol. cold, m. sol. hot, alcohol; slightly volatile with steam. Heated with HCl it forms PhAsCl. Above its melting-point it decomposes thus:

 $3PhAsO = AsPh_s + As_sO_s$

Phonyl-arsinc exy-chlorids PhAsOCl2. [100° Formed by decomposing the tetrachloride with the theoretical quantity of water; or by tho union of chlorine with the oxide. Crystalline; dissolved by water, being converted into benzens arsonic acid. At 120° it splits up thus:

PhAsOCl2 = PhCl + AsOCl.

Bensens-arsonic acid C.H.AsO(OH). Formed by dissolving PhAsOl, or PhAsOOl, in water. Long columns; begins to soften at 138°, ohanging to an amorphous anhydride, which is re-converted by water into the acid. M. sol. oold, v. sol. hot, water.

Reactions .- 1. Not affected by reducing or oxidising agonts. - 2. Potash fusion produces

Salts .-- NH .HA": needlos .-- KHA": amorphous.—Ball,A": needles, v. sol. water.—CaH₂A": needles.—CaA"2aq.—CuA": v. sl. sol. water.—PbA": insol. water.

Totra-phonyl-di-arsenide As (Calla), Phenyleacodyl. [135°]. Formed by reduction of tetraphenyl-di-arsino oxido with phosphorous acid. White crystals. Sl. sol. alcohol and acid. White crystals. Sl. sol. alcohol and ether. It quickly oxidises in the air, forming di-phenyl-arsinic anhydride (Ph.As.O.).

Di-phenyl-arsine ohloride Ph. AsCl. Phenyl-

cacodyl chloride. (333°). S.U. 14 1.42. Propared by heating HgPh, with a large excess of PhAsCl, at 320°. The product is then fractionally distilled. Yellow oil, insol. water, sol. alcohol, ether, and benzene, sl. sol. aqueous alkalis. Not affected by heating with Na.CO₃. Combines with hromine and chlorine. Conc. HNO₃ slowly converts it into di-phenyl-arsinie acid.

Di-phonyl-arsiao trichloride [174°]. From the preceding and oblorino. Colourloss tables (from benzene). At 200° it decomposes

thus: PhasCl, = PhasCl, + PhCl.

Di-phenyl-arsine chloro-hromido Ph. AsClBr., Formed by passing dry bromine-vapour into Ph, AsCl. Excess of Br produces di-broino-

Tetra phenyl-di-arsine oxide (Ph.As).O. [92°]. Formed by heating Ph. AsCl with alcoholic KOII. Di-pheayl-arsine bromide Ph.AsBr. (356°).

From the oxide and HBr.

Di-phoayl-arsino oxy-chlorido (Ph2AsCl2)2O. [117°]. From the oxide and chlorine.

Di-phonyl-arsinic acid Ph. AsO.OH. [174°]. S.G. 155. From the oxy-chlorido or the trichloride by the action of water. White needles; sol. water and alcohol, sl. sol. benzeno and ether. Not attacked by CrO, or boiling conc. HNO,

Salts.—NaA'.—NII,A': mustable feathery crystals.—BaA'₂.—CuA'₂.—HO.CuA'.—AgA'.—

PbΛ′2.

Tri-phenyl-arsine AsPh₃. [59°], (above 360°). S.G. 1 306. Prepared by heating phenyl-arsine oxido at 200°, thus: 3PhAsO = AsPh₃ + As₂O₃. More readily by the action of sodium (50 g.) on AsCl, (54 g.) and chloro-benzeno (101 g.), diluted with 4 vols. dry ether. Is also a by product in preparing Ph.AsCl from PhAsCl, and HgPh. Triclinic crystals isomorphous with SbPh, (Philips, B. 19, 1031). Insol. water and dilute acids, v. sol, hot alcohol, benzene, and ether. With HgCl, it forms leadlets of Asl'h, HgCl, whenco aqueous KOII forms AsPh₃(OH), [1082], thus: $\mathbf{AsPh_3HgCl_2 + 2KOH = AsPh_3(O11)_2 + 2KCl + Hg.}$

Tri-phenyl-arsiae chloride Ph, AsCl. [171°]. From AsPh, and chlorine. Tables; decomposed at 280° into PhaAsCl and PhCl.

Tri-phenyl-arsine sulphide Ph, AsS. [162°]. Prepared by digesting Ph, As with S dissolved in CB2; or by action of ammonium sulphido on Ph.AsCl. Silky needles, insol. water and ether. Tri-phenyl-arsine oxy-nitrate

(C.H.),As(OH)NO, [84°]. Formed by adding (C₄H₃)₃As(OH)₃C₅. Formed by saming HNO₂ to an aqueous solution of the hydroxide (C₄H₃)₃As(OH)₂ (Philips, B. 19, 1033). Long glistoning needles. V. sol. alcohol, sl. sol. water.

Tri-nitro-tri-phenyl-arsine oxide (C₈H₄.NO₂), AsO. [254°]. Formed by nitration of tri-phonyl-arsine-hydrate, (C, H,), As(OH), with HNO₃ and II₂SO₄. Nearly colourless large crystals. V. sol. acetic acid, insol. alcobol and ether.

Tri-amido-tri-phenyl-arsine (C,H,NH2),As. [o. 176°]. Formed by reduction of tri-nitro-triphenyl-arsino oxido (C₆H₅.NO₂), AsO with tin and HCl in acetio acid solution (P.). Colourless crystalline solid. V. eol. alcohol and dilute acids, insol, water.

Salts.-B"H3Cl3: crystallino solid, easily soluble in water and aloohol. — $(B'''H_3Cl_3)_2(PtCl_4)_3$:

yellow pp., insol. cold water.

Tri-acetyl derivative (C.H. NHAc), As. [o. 230°]. Very sparingly soluble in alcohol, more easily in acetic acid.

Tri-p-ethoxy-tri-phenyl-arsine

(EtO.C.H.) As. Tri - phenetyl - arsine. [890]. Formed by the action of sodium upon a mixture of p-bromo-phonetol and AsCl, (Michaelis a. Weitz, B. 20, 52).

tri-p-methoxy-tri-phenyl arsine (MeO.C., II,), As. Tri-anisyl-arsine. Obtained by the action of sodium upon a mixture of p-bromo-anisol and AsCl, containing some acctio ether. Transparent colourless erystals. V. sol. beuzene, sl. sol. alcohol and HI splits it up into di-anisyl-arsine iodide (C,H,OMe) AsI and anisol; hy longer and higher heating anisol and Asl, are formed. By heating with an excess of AsCl, it yields anisyl-arsino chloride C₈H₄(OMe).AsCl₂ (Michaelis a. Weitz, B. 20, 48).

Di-p-mothoxy-di-phenyl-arsiae chloride (CaH, OMe) AsCl [1:4]. Di-anisyl-arsine chloride. [80]. Formed by dissolving the oxide in 11Cl. Long thin needles. V. sol. ether, less in alcohol.

Di-mothoxy-di-phenyl-arsine oxido $\{(C_6H_1.OMe)_2As\}_2O$ [1:4]. Di-anisyl-arsine oxide. Tetra-anisyl-di-arsine oxide. [130°]. Crystalline. Formed by the action of alkalis on the iodide which is obtained by heating trianisyl-arsine with HI.

p-Methoxy-benzene-arsine chloride

C.H. (OMe). AsCl., [1:4]. p-Anisyl-arsine chloride. (230° at 117 mm.). Colourless liquid. Formed by heating tri-anisyl-arsine (C.H.OMo), As with an excess of AsCl, at 200°. Alkalis yield the oxide C_aH₄(OMe).AsO, a colourless crystal-line solid. It combines with Cl₂ to form C. II, (OMe) AsCl, which is a thick yellow liquid decomposed by water giving anisyl-arsinic acid C.H.(OMe).AsO(OH),

p-Methoxy-beazene-arsonic acid

(C,H,OMe).AsO(OH),. Anisyl-arsinic acid. [160°]. Formed by the action of water upon the chlorido C_aH₁(OMe).AsCl₄. Colourless crystalline solid. Sol. hot, sl. sol. cold, water; v. sol. alcohel. On heating it gives the anhydride C_aH₁(OMe).AsO₂.—Ag₂A": white pp.

Phenyl-di-methyl-arsine PhAsMo, From ZnMo2 and PhAsCl2. Mobile liquid, sol.

alcohol and henzene, insol. water.

Phenyl-tri-methyl-arsoninm iodide PhAsMe,I. [244°]. From the preceding and Mel. White needles; sol. water and alcohol, insol. ether.—(PhAsMe,Cl),PtCl, [219°]; v. sol. hot water.

Di-phenyl-methyl-arsine Ph.AeMe. From Ph2AsCl and ZnMe2 in benzene (Michaelis a. Link, A. 207, 199). Insol. water.

Di-phenyl-di-methyl-arsonium iodide

Ph_AsMe_I. [190°]. From the preceding and MeI. Needles; sl. sol. cold, v. sol. hot, water. Decomposed by heat into MeI and Ph.AsMe.—

(Ph_AsMc_Cl)_Ptcl [219].

Phonyl-di-ethyl arsins PhAsEt_. (210].

From PhAsCl, and ZnEt_. Colourless liquid.
Combines with Cl, forming PhAsEt_Cl..

Phenyl-tri-ethyl-arsonium iodids PhAsEt_L.

[II3°]. From the preceding and EtI at 100°. Prisms, turned yellow by sunlight; sol. water and alcohol, insol. ether. Decomposed when heated in an indifferent gas into EtI and PhAsEt, Gives with AgCl the chloride PhAsEt,Cl; whence (PhAsEt,Cl),PtCl, Gives with AgO the hydroxide PhAsEt,OII, an

alkaline syrup, absorbing CO, from the air.

Di-phsnyl-othyl arsine Ph.AsEt. (320'
From Ph.AsCl and ZnEt.. Colourless liquid.

Di-phonyl-othyl-arsine chloride Ph.AsEtCl. [137°]. From the preceding and Cl. Needles (from benzene); fumes in the air; decomposed by water.

Di - phenyl - di - ethyl - arsonium Ph_AsEt l. [181']. From Ph_AsEt and Ett.

Flat white needles.

Di - phenyl -methyl - ethyl - arsonium iodide Ph.AsMcEtt. [170°]. S. 11 at 15°; 844 at 100°. From Ph.AsMc and EtI or from Ph.AsEt and McI. Trunctric prisms; insol. ether. Split up by heat into Etl and Ph. AsMe.

Derivatives .- (Ph. AsMcEt), PtCl. Picrate Ph_AsMeEt,O.C. II. (NO2)3. [95°]. Sl. cold water.

Tolyl-arsins chloride C, II, AsCl.

Ortho (265°). Para [31°]. (267°).

From AsCl, and mercuric di-tolyl (o- or p-). Bromine converts them into di-bromo-toluenes. Tolyl-arsine tetrachloride C.H.AsCl..

Tolyl-arsins oxide C,H.AsO.

Ortho [145°], Para [156°]. From C.H.AsCl, and aqueous Na.CO, Combine with Cl., forming oxy-chlorides.

Tolnene arsonic acid Cll, C, H, AsO(OH)...

Ortho [160°]. The para compound decomposes

above 300° without previous fusion.

From the tetrachloride or the oxychloride, C.H.AsoCl₂, by treatment with water. The ortho acid forms a crystalline anhydride C.H.Aso...-Au, A". -- BaA". -- CaA" (La Coste a. Michaelis, A. 201, 255).

Di-p-tolyl-arsino ehlorido (C.H.) AsCl. (c. 343°). From C.H.AsCl. and Hg(C.H.). Liquid; not affected by aqueous Na CO3.

Chlorino gives (C.11.) AsCl.

Tetra-p-tolyl-di-arsine oxide ((C,II,),As),O. [98°]. Silky needles (from ether). Obtained by boiling the preceding with alcoholic KOH.

Di-p-tolyl-arsinie acid (C.H. AsO.OH. [167°]. Formed by boiling (C,II,) AsCl, with water. Oxidised to 'dibenzarsinic' acid.

Tri-p-tolyl-arsine (C.H,)2As. [145]. Obtained by heating C.Π.AsO.

Tri-p-tolyl-arsins dichloride (C,H,),AsCl2. [214°]. Not attacked by water.

Vol. I.

p-Carboxy-phenyl-arsine chloride

CO.H.C.H.As.Cl., [158°]. From the corresponding iodide and AgCl; or from the product (COCl.C.H.As.Cl., (?) of the action of PCl, upon CO.H.C.H.As.Cl.(OH), by treating with water. Needles (from benzene); decomposed by water.

p-Carboxy-phenyl-arsine iodide

CO₂H.C₆H₄As1₂ [153°]. From CO₂H.C₆H₄AsO(OH)₂ by III and P₄ Yellow needles (from chloroform).

p Carboxy-phenyl-arsino hydroxide

CO_H.C.H.As(OH)... Bencarrenious acid. From the preceding by heating with aqueous Na CO₄. Colourless needles (from water). At 145° 160° it gives off H_2O leaving the oxide $CO_2H_1C_6H_4AsO_1 \sim Ca(C_1H_4AsO_4)$ Aq: plates; changing at 200° into $Ca(C_1H_4AsO_3)_2$. changing at 200° AgO, II, AsO,

p-Carboxy-benzene arsonic acid CO.H.C.H.AsO(OH)... Benzarsinic acid. Formed by oxidising toluene arsonic acid with alkaline KMnO4. Transparent interlaced needles; m. sol. water, v. sl. sol. alcohol. At 190° it becomes CO.H.C. II. AsO.; at 230° it gives off benzoio acid. — Ag₃A". — CaIIA" aq. — KII. A". —

Mell.A".

 $p ext{-Di-earboxy-di-phonyl-arsino iodide}$ (CO.H.C.H.).AsI. [above 280°]. From (CO.H.C.H.).AsO.OH, conc. 111, and P. Converted by aqueous Na.CO, into the hydroxide (CO.11.C.11.) AsOH. (Dibenzarsenious acid).— CaA" 2aq.

p-Di-carboxy-di-phenyl-arsinic acid (CO,11.C,11.),AsO.O11. Dibenzarsinie Formed from (C,11,) Aso, Oll and alkaline KMnO_c at 60°. Leaflets, insol. water, 8l. sol. KMnO₄ at 60°. Leaflets, insol. water, sl. sol. alcohol.—Mc.HA" [above 280°].

p-Tri-earboxy-tri-phenyl-arsine (CO.H.C.11), As. Tribensarsenious acid. From the following acid and 111. Small colouriess needles.—Na₃A''' 2aq.—Ag₃A'''.

p-Tri-carboxy-tri-phenyl-arsine hydroxide (CO.H.C.H.), As(OH), Tribenzarsinic acid. From tri-tolyl-arsine and alkaline KMnO,... (CO.K.C.II.), AsO.

Benzyl-arsine chloride PhCH_AsCl. (175°) at 50mm. Formed by heating tri-benzyl-arsine with excess of AsCl₂. Easily oxidised by air: PhCH₂AsCl₂ + O = PhCH₂Cl + AsOCl.

Di-benzyl-arsinie acid (Ph.CII.)2AsO.OH

Preparation. - Sodium [50 g.) acting upon a solution of benzyl chloride (100 g.) and AsCl, (72 g.) in dry ether (500 g.) containing acetic ether (5g.) forms (PhCH₂),As, (PhCH₂) ArCl₃, and (PhCH₂),AsCl₃. Alcohol extracts the first, and converts the two latter into (PhCH2), AsCI(OH), and (PhCH_)_ASCI(OH) respectively, and they are then converted by agneous NaOH into (PhCH_), AsO.ONa and (PhCH₂)₃AsO (Michaelia a. Pactoff, A. 233, 60).

Properties. - Pearly white plates (from alcohol); attacks the nucous membrane; v. sol, hot alcohol, m. sol. hot water, sl. sol. ether. When strongly heated it gives benzoic aldehyde and dibenzyl.

Salts.—BaA', 8uq.—CaA', 6aq.—AgA'.

Reactions.-1. Conc. HCl forms AsCl, toluene, and benzyl chloride .-- 2. Boiling dilute HNO, has no effect; HNO, of S.G. 1.3 forms a. compound (PhCII₂)₂As(OH)₂NO₃ [129°]; cone

HNO, forms benzoic and arsenic acids.—8. Comnines with HCl forming (PhCH₂)₂AsCl(OH)₂ which crystallises from aqueous HCl in needles [128°]; this is decomposed by more water, giving (PhCH₂)₂AsO₂OH again.—4. HBr forms (PhCH₂)₂AsO₃OH)HBr.

Di-bonzyl-thio-arsinio acid (PhCH₂)₂AsO.SH. [199°]. From di-benzyl-arsinic acid and H₂S in

alkaline solution.

Tri.benzyl-arsine (PhCH₂)₃As. [104°]. The preparation is described under di-benzyl-arsinic acid (v. sup.). Monoclinic needles (from alcohol). Insol. water; v. sol. ether, benzene, and glacial acetio acid; sl. sol. cold alcohol. Like AsMe₃, but unlike AsPh₃, it combines with alkyl iodides. It is not affected by boiling cone. HCl. It combines with S and halogens. Boiling dilute IINO₃ forms benzoic and arsenic acids. An ethereal solution gives with an ethercal solution of HgCl₂ a pp. of (PhCH₂)₃AsHgCl₂ [159°].

Tri-bonzyl-arsine oxide (PhCH,),AsO. [220°]. From tri-benzyl-arsine chlorido or oxychlorido by treatment with alkalis; or together with dibenzyl-arsinic acid by the action of wet ether upon the product of the action of sodium upon benzyl chloride and AsCl,. Prisms (from dilute alcohol); v. e. sol. alcohol, sl. sol. water and

ether.

Tri-benzyl-arsino oxy-chloride

(PhCH_),AsČl(OH). [163°]. Iformed by union of HCl with the preceding. V. o. sol. aloohol; insol. dilute HCl.

Tri-benzyl-arsine oxy-bromide

(PhCH_)₃AsBr(OII). [129°]. Tables (from alcohol). Tri-benzyl-araine iodide (PhCH_)₃AsL_. [o. 95°]. Formed in impure condition by action of aquoons 1H on the oxide. Converted by alcohol into the oxy-iodide, (PhCH_)₃Asl(OH) aq [78°].

Tri-benzyl-arsiac-oxy-nitrate (PhCH₂)₃As(NO₃)(OII). [170°]. Slender needles

(from alcohol).

Tri-benzyl-arsine sulphide (PhCH.), AsS.
[214°]. Prisms (from glacial HOAc). Insol.

alcohol and ether. Tri-benzyl-methyl-arsonium iodide

(PhCH₂)₂AsMoI. [143°]. From (PhCH₂)₂As and MeI at 100°. Slender needles (from water). Gives with moist Ag₂O the alkaline hydroxide, (PhCH₂)₂AsMo(OH).

Tri-benzyl-methyl-arsonium chloride (PhOH₂)₂AsMeCl.[201°]. — Platinochloride ((PhOH₂)₃AsMe)₂PtCl₂. [173°].

Tri-benzyl-othyl-arsonium iodide (PhCH₂), AsEtl. (148°). White plates (from water).

Tri-benzyl-propyl-arsonium iodide (PhCH₂)₂AsPrI. [146°]. Tho isomoride, (PhCH₂)₂AsPrI melts at [143°].

Tri-benzyl-isoamyl-arsonium iodide (PhCH₂)₃As(C_3H_{11})I. [146°].

Tetra-benzyl arsonium chloride (PhCH₁),AsG I. [160°]. From (PhCH₂)₄As and PhCl1,Cl at 170°. Triclinio orystals containing aq (from water); insol. diluto IICl. Converted by aqueous KBr into the bromide (PhCH₂),AsBr. [173°], and by aqueous KI into the iodide, (PhCH₂),AsI, [160°]. Moist Ag₂O forms an alkaline hydroxide, split up by heat thus: (PhCH₂),AsOI = PhCH₂+ (PhCH₂),AsO.

Platinochloride ((PhCII2),As),PtCl.

Di-naphthyl di-arsenide O₁₂H₁.As:As.O₁₀H₂. Arseno-naphthalene. [221°]. Prepared by heating an alcoholic solution of naphthyl-arsine oxide with phesphorons acid (Michaelis a. Schulte, B. 15, 1954). Slender yellow needles; sl. sol. alcohol, benzene, CS₂ and chloroform; insol. water and ether. Converted by Cl. into C₁₀H₂.AsCl₂. With sulphur it gives C₁₀H₂.AsS. It is oxidised by HNO₃ to naphthalone arsenic acid.

Naphthyl-arsine chloride C₁₀H,AsCl₂. [63°]. From mercury ai-naphthyl and AsCl₂. Crystalline powder; insol. water, v. sol. alcohol.

Naphthyl-arsine oxide CloH,AsO. [245°]. From the preceding by treatment with aqueons Na₂CO₂. Powder; sl. sol. alcohol, ether, and water. On dry distillation it gives C, As, and naphtbalene.

Naphthalene arsonio acid C₁₀H,AsO(OH)₂. [197^o]. Needles. Formed by action of water on C₁₀H,AsCl, which is obtained by treating C₁₀H,AsCl, with chlorine (W. Kelbe, B. 11, 1503).

ASAFŒTIDA. A gum-resin obtained by drying the juice contained in the root of Ferula asafætida, a Persian plant. Potash fusion gives resorein and protocatechuic acid. Asafætida contains ferulic acid (q. v.), but its odour is dae to 3 p.o. of an essential oil (135°-140°) which appears to be a mixture of C₁₂II₂S and C₁₂II₂S. Its alcoholic solution is ppd. by HgCl₂ (Pelletior, Bull. Pharm. 3, 556; Johnston, P. M. Dec. 1838; Hllasiwetz, A. 71, 23).

ASARITE.—Impuro asarono.

ASARONE (1,H₁₆O₃, [59°]). (296°). S.G. 18
1·165. Contained in the root of Asarum europeum. Needles or plates; v. sol. alcohol, ether, and glacial HOAo, sl. sol. hot water (Blanchet a. Sell, A. 6, 296; C. Schmidt, A. 53, 156; Butlerow a. Rizza, B. 17, 1159; Bl. [2] 43, 114; Poleck, B. 17, 1415).

ASCLEPIONE C₂₀H₃₁O₃. [104°]. Extracted by ether from the coagulum got by heating the milky juice of Asclepias syriaca. Radiating crystals; insol. water and alcohol. Not attacked by boiling KOHAq (List, A. 69, 125; Gram, C. C. 1886, 735).

ASEBÓTOXIN. C. 60·5 p.c.; H. 7·4 p.c.; O. 32·1 p.c. [120°]. A glucoside extraoted by water from the leaves of Andromeda japonica. Brittle mass. The addition of cono. HCl to its alcoholic solution gives a blue colour (Bijkman, R. 1, 224; Ph. [3] 13, 365). It is accompanied by a glucoside, aschoin $C_{u_1}H_{u_2}O_{u_2}$, crystallising in yellow needles [147·5°] and also by aseboquerectin $C_{u_1}H_{u_2}O_{u_1}$ and aschofnsoin $C_{u_2}H_{u_3}O_{u_4}$ (Eijkman, J. 1883, 1410; R. 2, 99, 200).

ASH OF ORGANIC BODIES.

The inerganic constituents contained in vegetable and animal products are usually determined by incineration of the substance, and determination of the weight and composition of the ash. The first question to be considered is —Does this ash accurately represent the inorganic constituents of the substance?

The sulphuric acid originally present is undoubtedly but imperfectly represented. The tendency to the reduction of sulphates to sulphides during ignition with carbonaceous, and especially with nitrogenous, matter, is generally overborne by the oxidation of the sulphur contained in the albuminoids. The sulphuric acid found in the ash is thus greater.

than that originally present; it entirely fails, however, to represent the sulphnr present in the original substance; this must be determined by a special experiment.

The carbonic acid originally present in the substance is generally quite undiscoverable by an analysis of the ash. Carbonic acid may be lost by the decomposition of calcium and magnesium carbonates during ignition; or by the decomposition of carbonates by the action of silica, or of phosphates containing less than three equivalents of base. On the other hand carbonates are produced when tribasic alkali phosphates are ignited with carbon; they are also formed in large quantity during the incineration of organic substances containing nitrates, or salts of organic acids. Treatment of an ash

Phosphoric soid may be lost if acid phosphates are heated to a high temperature with carbonaceous matter. The alkali metals are also liable under some oircumstances to suffer loss by volatilisation.

The ash constituents are obtained with the smallest loss when the ignition is conducted at a low temperature, preferably in a mufile. In some cases an excess of line or baryta must be added to prevent losses of phosphorio acid and chlorine; this treatment also prevents the fusion of the ash (Strecker, A. 73, 366).

1. Ash of Annals.—The proportion of ash in some of the principal parts and products of the animal body, and its percentage composition, are shown in the following table. The figures are taken from Wolff's Aschen Analyses.

ASH OF ANIMAL PARTS AND PRODUCTS.

210	II OF AL	1771777	1 2213	110 2	IND	1 16().	DOOTE				
	Number	Pure ash in									
	of Analyses	100 dry sub- stance	К,0	Na _z O	CaO	MgO	Fe ₃ O ₄	P.O.	SO.	SiO.	Œ
Blood, luman	4	_	26.6		0.9		8.2	8.8	7.1		30.7
,, ox	7	3.77		45.0			9.4	5.3	3.1	0.8	34.4
" calf	2		11.2				8.3	7.8	1.3		34.7
"sheep	2 3	_	7.1	45.0			9.6	5.5	1.9	l —	35.8
" pig		4.29		29.4			8.9	12.2	1.0	l —	28.5
" horso	1			21.2			9.5	8.4	6.3		28.6
" dog	8			39.9		1.1	9.0	12.6	3.3		32.0
Flesh of mammalia .	8	4.32		10.1	2.1	3.2	0.4	41.2	1.0	0.7	4.7
" fowls	2	- 1		18.7		4.2		36.4			8.1
" marino fish .	1			14.9	15.2	3.9		31.5			11.4
Meat extract	11	20.89	43.9			3.1	0.4	29.8	2.2	0.9	10.0
Meat flour	2	1.80	5.4	3.0	22.1		13.3	43.5	1.2	0.9	1.8
Bone of ox carcase	1				53.5		0.2	40.3		0.5	 -
Wool, unwashed	3	8.33	79.4			0.6	0.7	1.0	4.7	2.9	4.5
" washod	1	1.11	19.1		24.7	6.0	18·2	3.2	-	25.3	0.8
Colostrum, cow	1	1.18*			34.9		0.5	41.4	0.2		11.8
Milk, human	4	0.49*			16.7		0.2	22.7	1.0	<u> </u>	18.4
,, cow	9	0.72			23.2		0.4	28.0	1.3		13.5
" cwo	2	0.73*	21.3		29.3		1.0	35.8	1.6	2.0	7.5
" mare	1	0.37*	25.1	3.4	30.1	3.0	0.4	31.9		l —	7.5
,, SOW	1	1.05*			39.2	1.8	0.9	37.2	1.3	<u> </u>	9.3
"bitch	2	0.73*			33.7	1.6	0.1	37.2		-	13.1
Whey, cow	1 2 3 3	0.51*		13.8			0.6	17.1	2.7		15.8
Hen's egg, without sholl.	3	3.48		22.9			0.4	37.6	0.3	0.3	9.0
" whito	3	4.61		31.6			0.6	4.4	2.1	1.1	28.8
,, yolk	3	2.91	9.3	5.9	13.0	2.1	1.7	65.5	- 1	0.9	1.9
			r					1	1	1	,

^{*} These numbers represent per cent, of pure ash in fresh substance.

with ammonium carbonate, and re-ignition, is sometimes employed with the view of converting caustic lime and magnesia into carbonates. Such treatment converts stllphides and cyanides into carbonates; sulphate of calcium is also partially converted into carbonate if only a small proportion of alkali carbonate is present. Treatment with carbonic acid water is more free from objection, but its action is slow.

The chlorides found in an ash may be below the truth from volatilisation of alkali chlorides, if too high a temperature has been employed; or from loss of hydrochloric acid due to the action of organic acids produced during the charring of the organic matter; or from a similar action of silica, or dibasic phosphates, at a high temperature.

(1880), with the exception of the analysis of bone ash, which is quoted from G. J. 24, 80, The bone-ash represents the mean composition of all the carcase bones of the 'fat ox' analysed at Rothan sted. In this analysis, alkalis, and sulphurie and carbonic acids, were not determined; the whole amount of these constituents was, however, but 4.5 p.c. The 'pure ash' in Wollf's Tables is exclusive of sand, charcoal, and carbonic acid.

The amount of ash yielded by the entire bodies of the principal animals reared on the farm, and its composition, have been determined by Lawes and Gilbert (T. 1883, 865); they also separately analysed the ash of the carcase and offal parts. The percentage composition of the ash of the entire bodies of oalf, ox, lamb, sheep,

Reactions .- 1. Gives on oxidation benzoio acid and tri-phenyl-carbinol.—2. Heating with alcoholio KOH produces tri-phenyl-methane and benzoio acid (Zagumenny, Bl. [2] 31, 330). 3. Reduced by HI to s-tetra-phenyl-ethane (?).

BENZPINACONE C.,H.,O, i.e.,.C(OII).C(OH).Ph., Tetra-phenyl-ethylene Ph. C(OH).C(OH).Ph. Tetra-phenyl-ethylene glycol. [168°]. S. (benzene) 3.8 at 80°; S. (HOAc) 8.7 at 118°; S. (15 p.e. alcohol) 2.5 at

Formation .- From benzophenone by roduoing the alcoholic solution with Zn and H2SO4 (Linnsmann, A. 133, 26) or a solution in acstio acid (10 pts.) diluted with water (2 pts.) with zine (Zagnmenny, J. R. 12, 426).

Properties. - Minute prisms, sl. sol. boiling alcohol, v. sol. ether. On fusion it splits up into benzhydrol and benzophenono (Thorner a.

Zincke, B. 10, 1473).

Reactions.—1. Chromic acid oxidises it to benzophenone.—2. Sodium analyam reduces it to di-phenyl-earbinol .- 3. Readily converted into (a) or (B) benzpinacoline by dehydration; this ie effected by BzCl, AcCl, dilute acids, or even by recrystallisation from alcohol (Za.) .- 4. Ac.O gives benzhydrol and benzophenene. - 5. HI and P at 1703 give tetra-phenyl-ethane (Graebe, B. 8, 1054).

BENZ URAMIDOXIM C.H.N. i.e. C.H., C(NOH)(NILCO.NH.). [115]. Formed by the action of potassium cyanate upon benzamidoxim hydrochloride in conc. aqueous solution (Falck, B. 19, 1186). Long thin white needles. V. sol. alcohol, ether, benzene, and ligroïn, sl. sol. water.

BENZ - URANILIDOX1M $C_{14}H_{12}N_{4}O_{2}$ i.e. C.H., C(NOH). NPh.CO.NH. ("). Benz. phenyl-uramidoxim. [167°]. Formed by the action of potassium cyanats upon benzanilidexim hydrochlorido in concentrated aqueous solution (Müllor, B. 19, 1671). Yellowish needles. Sol. alcohol, ether, benzene, and chlorolorm, insol. water.

BENZYL. The radiels phenyl-methyl, C.H. CH. It is isomeric with methyl-phonyl or tolyl CH, C,H.

DIBENZYL v. s. DI-PHENYL-ETHANE.

BENZYL-ACETAMIDE v. Acetyl - Bunzyl-

BENZYL ACETATE C.H. CH. O.CO.C11, (206°). S.G. 163 1 057. From benzyl alcohol (2 vole.), acetic acid (4 vols.) and H.,SO, (1 vol.), or by boiling benzyl chloride with alcoholic KOAo (Cannizzaro, A. 88, 130). Formed also by boiling a mixture of benzoic aldehyde and glacial aestic acid with zinc dus (Tiemann, B. 19, 355). Oil, smelling of pears. Sodiumacting upon benzyl acetate does not form benzyl aceto-acetate but the chicl product is benzyl 8. phenyl propionate: 4CH, CO, C, H, + Na₂ = 2CH, CO, Na + 2C, H, CH₂, CO₂C, H, + H₁, and by a secondary reaction, sodio phenyl propionate, sodic phenyl-acrylate, and tolueno;

CH,CH,CC (Conrad a. Hodgkinson, A. 193, 300).

BRHZYL.ACETIO ACID v. B. PHENYL-PROPI-DHIC ACID,

Di-beasyl-afette H. CH.), CH.CO.H. Di-phonyl-isobutyric soid 185°]. Obtained by saponifying the ether, by heating di-benzyl-malonio ether with alcoholie KOH (Lellmann a. Schleich, B. 20, 439), or by heating di-benzyl-malonic scid (Bischoff s. Siebert, A. 239, 101).

Properties .- Prisms (from ligroin), al. sol. cold water, v. sol. alcohol. Heated with soda-

lime it gives di-benzyl-methane.

Salts .- AgA': trimetric prisms, sol. boiling water (Michael a. Palmer, An. 7, 70). - BaA',

Ethyl ether EtA'. (above 300). Formed, together with B-phenyl-propionio ether by heating acetio ether with benzyl chloride and sodium (Lydia Sesemann, B. 6, 1086; Merz a. Weith, B. 10, 759).

BENZYL-ACETO-ACETIO ETHER v. p. 21.

BENZYL-ACETONE C, II, O i.e. C, II, CII, CH, CO, CII, Methyl phenylethyl ketone. (236°). S.G. 233 989.

Formation .- 1. By the dry distillation of a mixture of calcium hydrocimamate and calcium acetate; the yield is 33 p.c. (Jackson, B. 14, 890). 2. From benzyl-aceto-acetic ether by boiling with alcoholic KOII (Ehrlich, A. 187, 15).

Properties .- Oil. Combines with NallSO, forming C₁₀H₁₇ONnHSO₃nq. Oxidised by CrO₂

to nectio and benzoic acids.

BENZYL ACETONE Y-CARBOXYLIC ACID

V. ACKTYL-PHENYL-PROPIONIO ACID.

Benzyl-acetone o-carboxylic-acid C, II, O. i.e. CO.H.C.H.,CH.,CH.,CO.CH., [114°]. Obtained by boiling o carboxy benzyl aceto acetio ether with baryta-water (Bidow, A. 236, 192). Slender needles (from water).

BENZYL-ACETOXIM v. ACKTOXIM, p. 38. BENZYL ACETYL SUCCINIC ETHER v. ACETYL-BENZYL-SUCCINIC ETHER, p. 39.

P. BENZYL DI ACETYL PYRROL v. 151 NZYL-

PYRKYLENE-DI-METHYL-KRYONE.

BENZYL ALCOHOL C 11,0 i.e. C, 11 .C11, OF Mol. w. 108. (206 5°). S.G. 2 10129 (Bruhl S. 4 at 17°, μ_{σ} 15518. R_L 53 16. H.F. 38,73 (Stohmann, J. pr. [2] 36, 4).

Occurrence. - Bulsam of Peru contains bonzy benzonte, benzyl cinnamate and small quantitie of benzyl alcohol (Kraut, A, 152, 129). Liquic storax contains benzyl cinnamato (Laubenheimer A. 167, 285). Balsam of tolu contains benzyl cinnamate and some benzyl benzoate (Busse, B. 9, 830). In small quantity, together with benzoic aldehyde, prussic and, and a resin in the volatile oil of cherry-haurel (Tilden, Ph.

[3] 5, 761).

Formation. -1. Together with KOBz by the action of alcoholic KOU on benzoic aldebyde (Cannizzaro, A. 88, 129). 2. From benzyl chloride by converting it into benzyl acetate by alcoholic KOAc, and boiling the product with alcoholic KOH (Cannizzaro, A. 96, 216).—3. From benzyl chloride by heating with an aqueous solution of K,CO, (Mennier, Bl. [2]) 38, 159); with water (10 pts.) and freshly ppd. Ph(OH)₂ (3 pts.) (Lauth a. Grimanx, A. 143, 81)₃ or merely with water (30 pts.) (Niederist, A. 196, 353).—4. From balsam of Peru by bolling with aqueous KOH (Kachler, J. pr. 107, 807) .-5. By the action of sodium amalgam upon

benzoic aldehyde (Friedel, J. 1862, 263), benzoic acid, hippuric acid (Hermann, A. 182, 76; 188, 835), benzoyl chloride in presence of HCl (Lippmann, Bl. [2] 4, 249), or benzamide (Guarsschi, G. 4, 465).

Preparation.—10 pts. of beazaldehyde are shaken in a stoppered oylinder with e solution of 9 pts. of KOH in 6 pts. of weter, and left to stand over night. Sullicient water is then added to dissolve the potessium benzoate which has seperated, and the solution is extracted with ether; after evaporeting the sther the residue is distilled; the yisld is 92 p.c. of the theoretical. Benzyl alcohol cannot be dried with CaCl, as it combines with it (Meyer, B. 14, 2394).

Properties .- Liquid with little odour, al. sol.

water, sol. elcohol and ether.

Reactions .- I. Oxidised by dilute HNO, or air end platinum black to benzoic aldehyde, and by CrO, to benzoic acid. 2. III and P at 140° reduce it to toluene (Grache, B. 8, 1051). -3. Alcoholic KOH forms toluene and benzoie acid (Camuzzaro, A. 90, 253). 4. Conc. II,SO, acm (Galmizzaro, A. 30, 250); A. Cone, I. 37, 1920, and ZnCl₂ form a resin (Galmizzaro, A. 92, 113), ... 5. B.O. at 110? forms di benzyl oxide (C.H., CH.), O. -6. Solid cyanogen chloride forms bonzyl carbamate and di-benzyl-uren (Cannizzaro, (I. 1, 83; B. 3, 517). 7. Urea nitrate at 120° forms di-benzyl area and benzoic aldeliyde; at 140° it forms benzyl carbamate (Campisi a. Amato, G. I. 39). 8. BCl, forces s-di-phenyl-ethane and benzyl chloride (Councler, B. 10, 1655).

Methylether C.H.,CH.,OMc. (168°). From benzyl cldoride, KOII, and McOII (Sintenis, A. 161, 334). Also from benzyl sulphide, methyl alcohol, and Mcl (Calours, J. Ch. [5] 10, 23).

Ethyl ether C.H.CH.OEt. (1859). Gives anthracene when located with P.O. Chlorine in the cold forms HCl, ethyl clderide, and C.H.CHO; at a higher temperature it gives EtCl and berzyl chloride. Chlorine in the cold in presence of I forms oldoro-beazoic aldehydes and Etl (Sintenis, A. 161, 331). Br forms in the cold HBr, EtBr, kenzyl bromisle, benzoic aldeliyde, and BzBr (Paterno, B. 5, 288).

Isobutyl other Call Oll O.Call (e. 210)

(Claus a. Trainer, B. 19, 3006).

Phenyl other Pho.Cll.Ph. [391]. (2870). From phenot potassium, benzyl elderide and a little alcohol at 100° with inverted condenser for 3 hours (Staclel, A. 217, 41; Lauth a. Grimaux, A. 143, 41; Sintenis, A. 161, 337). Chitering white plates which feel greasy (from alcohol). Cone. HC at 100 splits it up into phenol and benzyl chloride. Cldorine in presence of HgO forms the chloro-phenyl ether. C.H., CH., O.C. H.Cl [71°]; bromine forms similarly C.H., CH., O.C.H. Br (60 '. o-Tolyl ether C.H., CH., O.C.H., CH., [1:2

Beneyl o crosyl oxide. (285° 290°) (Staedel, B.

 Tolyi ether C_aH_aCH_aO.C_aH_aCH_a [1:3].
 (300°-305°). Satiny tablets.
 p.Tolyi ether C_aH_aCH_aO.C_aH_aCH_a [1:4].
 From potassium p-cresol, a little alcohol, and benzyl ohloride (Staedel, A. 217, 44). The yield is 86 p.c. White silky scales or transparent straight them. aix-sided columns (from alcohol). Feels greasy.

(a) Naphthyl ether. An oil, decomposed : by distillation.

(β)-Naphthylether C, H.O.CH.Ph. [999]. From (β)-naphthol (70g.), KOH (37g.), a little water and alcohol, and benzyl ohloride (70g.). White plates (from alcohol). No smell. Not volatils with steam (Staedol, A. 217, 47).

Other benzyl ethers are described under the hydroxylated compounds from which they are derived.

BENZYL-O-AMIDO-ACETOPHENONE

CeH4(NHC,H,).CO.CH2. [81°]. Formed by heating o-amido-acetophsnone with benzyl chlorids (Bacycr, B. 17, 971). Large prisms. elcohol, ether, benzene, chloroform and CS., sl. sol. ligroin. Weak basc.

Nitrosamine C.H.(N(C,H.).NO)CO.CH, [55°]; long colourless needles. By heating with H.SO, it gives a mixture of indigo end benzyl-

indigo.

BENZYL-o-AMIDO-BENZOIC ACID

C,H,NH.C,H,CO.H. [176° uncor.]. Formsd together with its formyl derivative by oxidation of benzyl-quinoline with alkaline KMnO, Long needles or thick prisms. Salts C₁,H₁₃NO,1lCl: [105° uncor.]; large tables.—(C₁,H₁₃NO,),H.Cl,l'tCl,: [158° uncor.];

orenge yellow tables.

Formul derivative C.H.N(CHO).C.H., CO.H. [196°]; largo colourless tables (Claus a. Glyckherr, B. 16, 1283).

BENZYL-AMIDO IRI-PHENYL-METHANE Ph₂C.NH.CH Ph. [H0°]. The hydrochloride B'HCl [249°] is formed by the action of benzyl chloride on w-amido-tri-phenyl-methane (Elbs. B. 17, 703).

Di-benzyl-amide di-phenyl-methane

Ph.CH., C. II, N(CH., Ph). From aniliue hydrochloride and benzyl chloride at 120°. Also from acetanilide and benzyl chloride at 120°. White moorphous powder (Meldola, C. J. 41, 200). Soluble in benzene. Solutions heve a blue fluorescence.

BENZYLAMINE C.II,N i.e. C.H., CH., NII,

Mol. w. 107. (181°). S.G. 14 99.

Formation.-1. Together with di- and trioenzylamine by heating benzyl chloride with alcoholic NII, (Camcizzaro, A. 131, 128; Limpricht, A. 14I, 304). - 2. Together with di- and tri-benzylamine by the action of Zn and HCl upon benzonitrile (Mendius, A. 121, 144; Spica, G. 10, 515). -3. By reducing thiobenzamide C_sH_s.CSNH₂ with Zn and HCl (Holmann, B. I, 102) .- 4. From benzyl eyanate and KOH (Strakosch, B. 5, 692). - 5. By seponification of its acetyl derivative, obtained by the action of sectamide on benzyl chloride (kudolph, B. 12, 1297). - 6. By the ection of bromins in alkaliae solution on phenyl-acet-amide C 11 .C11 .. CONH .: the yield is 60 p.e. of the theoretical quantity (Holmann B. 18, 2738; Hoogsworff a. Van Porp, R. T. C. 5, 252).—7. Together with toluene, by energetic reduction of hydrobenzamide dissolved in absolute alcohol by means of sodium or sodium-amalgam; very good yield (O. Fischer, B. 19, 748).—8. By reduction of an alcoholic solution of benzaldehyds-phenyl-hydrazide by means of sodium-amalgam and acetic acid (Tefel, B. 19, 1928).—9. By reduction of benzaldoxim (5 pts.), dissolved in alcohol (15 pts.), at 50⁵-60⁶ with sodium-amalgam (160 pts. of 2½ p.c. Na), keeping noid by gradual addition of acetic acid: good yisld (Goldachmidt, tion in 1,000 parts of the fasted live weight! Graminacec.

The animal body is also given. The latter of the animal body is also given. will afford data for calculating the loss which a farm suffers by sale of stock. The fasted liveweight is inclusive of contents of stomach and intestines, but the constituents of these contents are not reekoned among the animal con-stituents. The 'puro ash' is inclusive of earbonic acid.

2. Asu or Plants. The composition of the ash of plants of agricultural importance will be given under the headings of the different crops, some general considerations will, however, be best made in the present place.

tion of the plant are (1) partly employed in the formation of new tissuo; (2) partly deposited as nerusting matter on the older tissues; while (3) bluble salts that are of no advantage to the biblible salts that are of no advantage to the Pant first accumulate in the sap, and then are grainhally removed from the plant by the action of right, and possibly by diffusion into the soil through the roots. The ash constituents most largery consumed in the formation of tissue are potasty consumed in the formation of tissue are potasty and phosphoric acid; in all the actively growing parts of a plant potash and phosphoric acid igneratly prependerate. Magnesia, lime,

Percentage Composition of Ash from captire Bodies of Animals.

	 K,0	Na ₂ 0	CaO	MgO	4 (c ₂ 1) ₃	$\mathbf{P}_{\mathbf{z}} O_{\mathbf{z}}$	SO,	SiO ₂	CI	CO,
Fat calf Half-fat ox Fat ox Fat lamb Store sheep Half-fat old sheep Fat sheep Very fat sheep Store pig Pat pig	5·4 4·4 4·5 5·7 5·6 5·3 5·5 5·5 7·1 8·6	3·8 8·1 8·0 8·6 3·9 8·4 8·6 4·5 4·2 4·4	44·0 45·3 46·6 41·6 43·1 44·4 41·6 43·3 40·6 38·5	2·2 2·0 1·5 1·8 1·8 1·7 1·8 1·9 2·0 2·0	0.5 0.1 0.5 1.2 1.4 1.0 0.9 0.8	40·4 40·2 39·8 39·0 39·0 39·2 38·7 40·1 40·1	1·1 0·9 0·8 1·2 1·8 1·1 1·0 1·0 2·3 2·2	0·1 0·2 0·1 0·3 0·7 0·6 0·9 0·6 0·2 0·1	1·6 1·2 1·5 1·9 2·3 1·6 1·6 2·3 2·2 2·8	1·3 2·0 2·1 1·5 1·1 1·8 1·7 1·7 0·6 1·2

Composition of Animal Bodies, per 1,000 parts fasted live weight.

						s jasica	acco acc	rigito.		
	Fat Calf.	Half-fat Ox.	Fat Ox.	Fat Lamb.	Stora Shcep.	ilaif-fat old Sheep,	Fat Sheep.	Very fat Sbeep.	Store Pig.	Fat Pig
Contents of stomach and intestines, moist Water	32·0 630·0 118·0 152·0 38·0 37·8	515·0 191·0 166·0 46·6	145.0	85·0 478·0 285·0 123·0 29·4 28·9	60·0 573·0 187·0 148·0 31·6 30·6	91·0 502·0 235·0 140·0 31·7 30·6	60·0 434·0 356·0 122·0 28·1 26·8	52·0 352·0 458·0 109·0 29·0 28·6	52·0 551·0 233·0 137·0 26·7 26·5	40·0 413·0 422·0 109·0 16·5 16·3
K ₂ O	2·06 1·48 16·46 0·79 0·21 15·35 0·41 0·05 0·63 0·47	1·46 21·11 0·85 0·41 18·39 0·38 0·13 0·59	1·26 17·92	1.66 1.03 12.81 0.52 0.26 11.26 0.39 0.12 0.53 0.43	1·74 1·20 13·21 0·56 0·37 11·88 0·52 0·21 0·72 0·37	1.68 1.04 13.50 0.52 0.42 11.99 0.35 0.20 0.51 0.53	1·48 0·97 11·84 0·48 0·34 10·40 0·31 0·26 0·44 0·41	1·29 12·40 0·55 0·30 11·08 0·29 0·16	1.96 1.10 10.79 0.53 0.22 10.66 0.53 0.05 0.57	1·38 0·73 6·36 0·32 0·13 6·54 0·29 0·03 0·43

The ash constituents of a plant are obtained from the soil by the roots. All matters in the soil which are soluble and diffusible will enter the plant by the root, the abundant evaporation of water from the surface of a growing plant maintaining a rise of liquid in the eapillary vessels. The substances entering the plant are not, however, limited to those existing in solunot, lowever, limited to those existing in solution in the soil, as the roots of plants exercise a solvont or digestive action on constituents of the soil not otherwise soluble in water. The wide differences in the assimilating powers of the roots of different plants are well illustrated by the special assimilation of alumina by the Lyco-

oxide of iron, and sulphuric acid, must also be reckoned as essential for plant growth. The incrusting ash constituents are calcium salts and silica; these are shiefly precipitated in the leaves, where evaporation is most active. The soluble salts remaining unused in the sap generally contain a large proportion of chlorides, and of sodium salts.

potash or lime according as one or the other preponderates in the soil. On the other hand plants olearly exercise a selective power, potash being stored up in large quantity, though soda rather than potash may be abundant in tho soil. This selective power is apparently not a property of the roots, but simply results from the fact that potash is removed from the sap to form tissue, while soda is not; potash salts can thus continue to enter the roots by diffusion or otherwise, while sodium salts having accumulated in the sap the tendency of dilfusion is new for them to pass through the roots into the moist soil (see Dehérain, Cours de Chimic agricole [1873] 77). The Rothamsted experiments show that potash greatly preponderates in hay, and in barley straw, when the seil supplies a sufficient quantity; but when potash fails, soda is retained by the plant to a considerable extent.

The variations in the composition of the ash of any plant do not extend to the seed; the ash of this is of very definite composition whatever the nature of the soil. The ash of a seed consists chiefly of potassium phosphate; soda is practically absent.

While the seed is forming, a migration of phosphorio acid and potash, and of mtregenous matter and carbohydrates, sets in from all parts of the plant, the roots included; a great part of these important constituents is finally stored in the seed. The extent to which the exhaustion of the plant, and the enrichment of the seed, proceeds, depends on the climate during the ripening period.

ASPARAGINE

C₁H₁N₂O₂ i.e. CO₂H.CH₂.CH(NH₂).CO.NH₂ or CONH₂.CH(NH₂).CO.H. Amido-succinamic acid. M. w. 150 (containing aq) S.G. 14 1.52. S. o. 1.8 at 10.5°; 53 at 100°. Occurs in juice of most plants, especially in growing buds and germinating seeds (e.g. asparagus, marsh-mallow, comfrey, potatoes, deadly nightshade, chestnuts, liquorico root, lettuce, convolvulus root, dahlia tubers, young shoots of vetch, peas, beans, and other leguminous plants). Lupine seeds that have not begun to grow contain no asparagino; after 15 days' germination more than 20 p.c. of asparagine may be extracted by water (Schulze a. Barbieri, J. pr. [2] 27, 339). When twigs full of young leaf-buds of the plane, birch, or horsechestnut are cut off and allowed to open by placing the cut end in water, the leaves are found to contain asparagino (S. a. B., J. pr. 133, 145). Asparagino may be formed by adding conc. animonia to mono-ethyl aspartate (Schaal, A. 157, 24).

Properties .- Trimetric prisms (containing aq) exhibiting left-handed hemiliedry. Sol. water, acids, and alkalis; insol. alcohol and ether. Its solution in water or alkalis is lavorotatory, in acids it is dextrorotatory. In HCl solution [a]p = about + 36°; in aqueeus solution about -6°; in ammoniacal solution about -11°.

Reactions .- 1. Boiling with lime, or barytawater, or with dilute H.SO, rapidly converts it into aspartic seid. - 2. Nitrous acid forms malio acid. -3. Impure asparagine is liable to undergo fermentation, changing to ammonio succinate. Am. 6, 419; Griess, B. 12, 2117).

Salts. — HA'HCl. — $(HA)_2HCl.$ — CuA'_2 . — CaA'_2 . — EAA'_3 . — EAA'_4 . yellow prisms (Smolka, M. 6, 915).

Estimation.—Heat the extract containing it with dilute HCl for some hours and determine the amount of NII, Cl formed. This corresponds to half the nitrogen in asparagine (Sachsse, J. pr. [2] 6, 118). Glutamino also splits off half its amidogen as ammonia when treated with HCl. Or the extract may be treated with bromine and NaOH (measuring evolved N2) both before and after heating with 11Cl (Sachsse). But asparagine gives off too much No when so treated (Morgen, Fr. 20, 37). It even gives off a little Nawhen treated with NaBrO beforeheating with liCl. These two errors nearly balance one another (E. Schulze, J. pr. [2] 31, 235). Solutions of sodic aspartate give off no N2 with NaBrO, but if NH3 be present more N2 is evolved than corresponds to the NII3. The increase may be 6 p.c. Leucine behaves in the same way as asparagine, but tyrosino behaves in exactly the opposito manner. Urea has the same in-thuence as NH₃. It is therefore hetter to determine the free NII, by distilling with MgO, CaO or even NaOII (comp. Berthelot a. André, C. R. 103, 1051). The presence of peptones will, of course, invalidate the determination, these are often absent from vegetable solutions; if present they must be removed: albuminoids may be ppd. by lead salts, peptones by tannin or phosphotungstic acid (E. Schultze, I.c.). Dextro-asparagine

 $C_2H_2(NH_2)(CO_2H).CO.NH_2$. Dextro-hemihedral erystals. Dextroretatory $[\alpha]_0 = +5^{\circ}41'$. Very sweet taste (ordinary asparagine is tasteless). Rather more solublo in water than ordinary asparagine. Occurs in the mother-liquors obtained in recrystallising the erudo asparagino prepared from the shoots of the vetch; 20 kilos of crude asparagine, obtained from 6500 kilos, of vetch gave 100 grms. of the pure dextro-asparagine.

The compounds prepared from dextroasparagino exhibit the same properties as those prepared from the lave-asparagine except that their rotatory power is reversed. By heating with 2 mols. of aqueous IICl at 170°-180° both asparagines give the same inactive aspartic acid (Pintti, C. R. 103, 134; B. 19, 1691).

Additional References.—Vauquelin a. Robiquet, A. Ch. 57, 88; Dessaignes, A. 82, 237; Piria, A. Ch. [3] 22, 160; Pasteur, A. Ch. [3] 31,70; Mercadante, G.5, 187; Portes, B.9, 1934; Dubrunfaut, J. pr. 53, 508; Corup-Besanez, A. 125, 291; Champion a. Pellet, B. 9, 724; Becker, B. 14, 1031; Do Luca a. Ubaldini, C. R. 59, 527; Buelner, Z. 1862, 117; Campani, Z. [2] 6, 87; E. Schulze, B. 15, 2855; J. pr. [2] 20, 397; 27, 339.

ASPARTIC ACID C.H.NO. i.e. CO.H.CII, CH(NII,).CO,H. Amido-succinic acid. Mol. w. 133, S.G. 124 1.66. S. 45 at 20°; 5.4 1009

Formation.-1. By boiling asparagine with lime, baryta, PbO, KOII, or HCl dissolved in water .- 2. By boiling albumen or casein with dilute H.SO, (Kreussler, J. pr. 107, 239; Ritt-4. MeI and KOH produce an amide of fumaric | hausen, J. pr. 107, 218). -3. By treating proacid, CO.H.CH:CH:CONH, (Michael a. Wing, teids with bromine (Illasiwetz a. Habermann, 4m, 6, 419; Griess, B. 12, 2117).

A. 159, 325).—4. From casein by treatment with SnCl, and HCl (H. a. H., A. 169, 162).-5. From diazo-succinic ether by reduction with zine dust and acetic acid (Curtius a. Koch, B.

19, 2460).

Preparation: 100 grms. of asparagine are boiled for 2 or 3 hours with an inverted condenser with 408 c.c. of pure aqueous hydric shloride (containing:11925 g. HCl per c.c.). To the cooled solution is then added 204 c.e. of aqueous NH3 (corresponding to the acid volume for volume). On standing for several hours the aspartie acid separates in colourless crystals. The yield is 90 p.o. of the theoretical (Schiff, B. 17, 2929).

Properties. - Small trimetrio rectangular plates. Sl. sol. water, insol. alcohol. Its solutions in alkalis are lavorotatory; its solution in HClAq is dextrorotatory, $[a]_D = +28^{\circ}$. The rotation is affected by the nature of the solution (Becker, B. 14, 1035). Aspartic acid (1 mol.) provents the ppn. of Cu(OII), (I mol.) by KOII.

Reactions .- 1. Nitrous acid converts it into malic acid,-2. McI and KOII form fumaric acid (Korner a. Menozzi, R. Istit. Lombard, 13, 352) .- 3. Not affected by boiling water or by magnesia .- 4. Heating in a current of HCl at 130°-200° produces two anhydrides: (a) insoluble in water $(C_{10}H_{11}N_{10}O_{11})$, (b) slightly soluble in water $(C_{10}H_{11}N_{10}O_{0})$. Both are consoluble in water $(C_{10}H_{11}N_{10}O_{0})$. verted by boiling baryta into aspartic acid; but when the former is heated for 2 hours at 125° with half its weight of urea it produces a gummy mass, soluble in water forming a solution that has all the characters of a proteïd. It is ppd. by acids, by NaCl, MgSO,, tannin, and HgCl, forming gelatinous pps. CuSO, and KOH give a violet solution (Grimaux, C. R. 93, 771).

Salts. II.A' HCl : deliquescent crystals .-H2A"HSO4 -- NaHA" aq: trimetric prisms; S. 89 at 12°, —BaILA", 4aq. —BaA"3aq.—GaA"4aq. — HgA", — PbILA", ... — PbA", —AgHA", —AgA", —CuA" 4 aq. S. 035 at 15°; 43 at 10°; v. sol. dilute HOAc. The insolubility of this salt may be used to detect and to isolate aspartic acid (Hoffmeister, Sitz. B. 75, 469). - CuA" 3nq (Curtius a. Koch, B. 19, 2460).

Mono-ethyl ether A"EtH. Its hydrochloride (A"HEt) HCl forms large colourless needles, [199°].

Di-ethyl ether A"Et.. Its hydrochloride A"Et, HCl forms excessively hygroscopic concentrio needles.

Di-methyl ether A"Meg. Its hydrochloride A"Me HCl forms very hygroscopic glistening prisms (Curtius a. Koch, B. 18, 1293).

Amide v. ASPARAOINE.

Di-phenyl - amide CO.H.C.H.NH. CONPh. [230°]. Formed, together with phthalimide, by the action of NH, on the diphenylamide of phtimlyl-amido-succinic acid CO.H.C.H.N(C.O.C.H.).CONPh. (Piutti, G. 16, 14).

Inactive aspartic acid C.H.NO. S. 42 at 7°. Formation.—1. By the action of boiling HCl on the product obtained by heating the acid ammonium salts of malic, maleic or fumaric acid.-2. By heating an aqueous solution of the hydrochloride of activo aspartio aoid for several hours at 170° (Michael a. Wing, B. 17, 2984; Am. 7, 278).

Properties. - Monoclinic needles. Converted by nitrous acid into inactive malic acid.

Salts.—PbA".—Ag,A".—H,A"HCl. Leve-aspartic acid O,H,NO,. Obtained from dextro-asparagine by treatment with HCl (Piutti, B. 19, 1693). Laworotatory. Its properties are the same as those of the dextrorotatory acid. Combines with dextro-acid to form an inactive modification.

Additional References.-Plisson, A. Ch. 40. 303; 45, 315; Boutron-Chautard a. Pelouze, A. Ch. 52, 90; Liebig, P. 31, 232; A. 26, 125, 161; Wolff, A. 75, 293; Piria, A. Ch. [3] 22, 160; Dessaignes, C. R. 30, 324; 31, 432; A. 83, 83; J. Ph. [3] 32, 49; Pasteur, A. Ch. [3] 34, 30; A. 82, 324; Pott, J. pr. [2] 6, 31; Radziszewski a. Salkowski, B. 7, 1050; Ritthausen a. Kreussler, J. pr. [2] 3, 314; Scheibler, J. Ph. [4] 4, 152; B. 2, 296; Kreussler, Z. [2] 6, 93.

ASPHALT. A natural product of the decomposition of vegetable substances. It is found on the shores of the Dead Sea, also in a molten state in Trinidad, and as a mineral deposit at Seyssel. It frequently impregnates other rocks. When distilled with water, petrolene ConH32

(280°), S.G. 2¹ 89, V.D. 9°5, passes over (Boussingault, A. Ch. [2] 64, 111; Voclekel, A. 87, 139).

ASPIDOSAMINE C.,H., N.O., (c. 100°). In quebracho bark (Hesse, A. 211, 263). Turns yellow in air. V. e. sol. ether, chloroform, benzeno or alcohol, v. sl. sol. light petroleum, insol. water. Its alcoholio solution turns litmus blue, neutralises HCl and tastes bitter.

Reactions. - 1. Solution of hydrochloride gives with Fe,Cl, a brownish-red colour .-- 2. Conc. II_uSO₄ gives a bluish solution.—3. Conc. H_uSO₄ and MoO₂ gives a blue liquid.—4. Conc. H_uSO₄ and K.Cr.O. gives a dark blue colour. -5. Boiling aqueous HClO, gives a magenta colour.

Platinochloride. B' H PtCl 3aq.

ASPIDOSPERMATINE C₂₀H₂₈N₁O₂. [162°]. In quebracho bark (Hesse, A. 211, 259). Crystalline. V. sol. chloroform, alcohol, or ether. In alcohol (97 p.c.) it turns litmus blue, has a bitter taste, and is lavorotatory $[a]_0 = -72.3^{\circ}$ at 15° in a 2 p.c. solution.

Reactions. - 1. HClO4 gives a magenta colour. 2. Conc. H2SO, and K2Cr2O, give no colour.-3. Fe₂Cl₆ gives no colour.

Salts.—Dilute HCl is neutralised by aspido. spermatine. NaOH or NH, give in the solution a thecoleut pp. (u. sol. pure water) which soon becomes crystalline. Salts are amorphous.-(B'HCl) PtCl, 1ag.

ASPIDOSPERMINE C., H., N., O., $[a]_{\rm b}$ (alcohol) - 100.2° ; (chloroform) -83.60 (dilute 11Cl) -62° (in all cases 2 p.c. solution at 15°). S. (alcohol) 2 at 14°; (ether) 71 at 14° (Wulfsberg, Ph. [3] 11, 269). An alkaloid present (with others) in bark called in the Argentine Republic quebraelio blanco or quebracho, colorado (Fraude, B. 11, 2189; Ilesse, A. 211, 251; Arata, C. J. 40, 622). Needles or pointed prisms (from alcohol or light petroloum). M. sol. alcohol, sl. sol. ether or light petroleum, v. sol. benzeue or chloroform. Levorotatory.

Reactions .- 1. HCl and PtCl, give a blue .- 2. HClO, gives a magenta colour.-3. Conc. H₂SO₄ no colour.—4. Cono. H₂SO₄ and MoO₂ no colour.—5. Conc. H₂SO₄ and K₂Cr₂O₇, a brownish-red turning dark green.—6. Salts give with NH, NaOH, Na,CO, or NaHCO, a

flocculent pp. becoming crystalline.
Salts.—Very unstable; even ether or CHCl, can partly decompose them .- B'2H2PtCl, 4aq.

ASSAMAR. A name given by Reichenbach (A. 49, 3) to a bitter, deliquoscent, transparent yellow solid which may be extracted by alcohol from toasted bread. It is insol. other. The same namo was given by Völckel (A. 85, 74) to a thick yellow neutral syrup obtained from the aquoous portion of the product of the distillation of cane-sugar. It is sol, other. Both substances reduce aqueous AgNO₃.

ASYMMETRIC CARBON. A name applied to an atom of carbon that is united to four different atoms or radicles. All compounds that in the liquid state or in solution rotate light contain asymmetric earbon (Van 't 11off, La chimie dans l'espace; Le Bel, Bl. [2] 22, 337).

ATHAMANTA, OIL OF. C₍₀H_{1c}. (163°). S.G. 84. An essential oil obtained from the leaves of Athamanta orcosclinum. It forms a liquid compound with HCl (190°) (Schnedermann a. Winckler, A, 51, 336).

ATHAMANTIN C. H. O., [79°]. In the root and seeds of Athamanta orcosclinum. Fibrous, silky crystals, or sometimes rectangular prisms; insol. water, v. sol. alcohol and ether. It gives valerie acid on dry distillation. Aqueous acids and alkalis split it up into valeric acid and oreosclone CaHcOr. Chloro-, and tri-nitro-, athamantin are amorphous (Schnedermann, A. 51, 315; Geyger, A. 110, 359).

ATHEROSPERMINE [128]. An alkaloid in the bark of Atherosperma moschatum. A greyish-white powder with bitter taste. V. sl. sol, water, m. sol. alcohol, sl. sol. ether. The solution of its hydrochloride gives pps, with phosphomolybdic acid, picric acid, taunin, and PtCl. It liberates iodine from iodic acid (Zeyer, J. 1861, 769).

ATMOSPHERE. The word Atmosphere (ἀτμός, vapour; σφαίρα, a globe) in its most extended sense signifies the gaseous envelope which surrounds any liquid or solid body: more commonly, however, it is taken to mean the invisible elastic fluid which surrounds the earth. A variety of phenomena, e.g. solar and terrestrial radiation. animal and vegetable life, weather, the dis-integration of rocks and the formation of soils, the propagation of sound, &c., are dependent on the existence of a terrestrial atmosphere. The earth is not the only planetary body which possesses an atmosphere. The Sun, body which possesses an atmosphere. Jupiter, Mars, Saturn, have doubtless very dense atmospheres, but as yet we have no exact knowledge of their physical and chemical natures.

The phenomena of solar eclipses, and the facts that a single star scems to disappear instantly when it is occulted opposite the smooth part of the moon's limb, and that there is no change of colour or other effect such as a refractive atmosphere would occasion, make it ocrtain that the moon's atmosphere, if it exists at all, must be of extreme tenuity. This conclusion is strengthened by evidence afforded by the spectroscope. It has been observed that the spectrum of the moon's light is identical with the solar spectrum and there is no trace of any absorptive action; moreover, it is found that on the assumption that the air contains 0001

the spectrum of a star during its occultation disappears as suddenly as the star itself.

Wollaston's arguments as to the finite extent of the terrestrial atmosphere were deemed inconclusive even by his contemporaries. There is indeed direct evidence for the belief that air is present in a state of sensible density at much greater heights than 40 or 45 miles which was the limit Wollaston assigned. Linis, from observations on the phenomena of smilight at Rio Janeiro, arrived at a superior limit of 200 miles; and Secchi, from observations on luminous meteors, calculated that air exists of appreciable density even at a height of 200 kilometres above the earth's surface. It is in fact probable that no actual limit exists. Up to the present it has heen impossible to arrive at direct results otherwise than by astronomical observations, as the law of the diminution of temperature which in great measure governs the extent of the repulsion among gaseous particles is unknown for the upper strata of the atmosphere. No arguments can be based on the finite expansibility of gases. Faraday's experiments on the limits of vaporisation of mercury have been controverted by Merget. It is obvious that thorelative distribution of the mass of the air will be modified by the increase of attraction at the pides as compared with that at the equator; by the increase of temperature as we approach tho torrid zone; and by the earth's motion.

The ponderability of air although suspected before the time of Arislotle was first conclusively demonstrated by Galileo, who found that a copper ball containing condensed air weighed more than when filled with air of ordinary tension. The weight of I litre of air, freed from aqueous vapour, carbonic acid, and ammonia, at 0 C., and under a pressure of 0.76 m. of mercury, at Paris (lat. 48° 50'), and at a height of 60 m. above the sea level, was found by Regnault to be 1-293187 grams. According to Regnault I litre of oxygen at the normal temperature and pressure weighs I 429802 grams; 1 litre of nitrogen under the same conditions weighs 1 256167 grams. If x be the volume of exygen contained in I litre of air, and I-x that of the nitrogen, then 1.129802x + (1-x) 1.256167 = 1.293187, whence x = 0.2132 or in per cents. 21.32, which is considerably higher than that found by oudiometric analysis. According to Magnus I litre of pure air at 0° and 76 m, weighs at Berlin (lat. 52° 36') 1.29306 grams. Ph. v. Jolly found that at Munich (lat. 48° 8', 515 m. above the sea's level) 1 litro of oxyger at 0' and 76 m. weighed 1:429094 grains; and 1 litre of nitrogen under the same conditions weighed I 257614 grams. Reducing these numbers to the lat. of Paris and to a height of 60 m. above the sea's level, they become:

Jolly 1 429388 Oxygen 1.429802 1.257873 1.256167 Nitrogen

The Bureau Internat. des Poids et Mesures adopts for the weight of I litro of dry air under a normal barometric height of 1 mm. and at the normal temperature τ

 $\mathbf{P}\tau \mathbf{i} = \frac{1 \cdot 293052}{1 + 0 \cdot 00367} + \frac{1}{760}$

parts of carbonic acid, and that '00367 is the coefficient of expansion of air at constant pressure for a normal degree.

This expression is obviously only true for a particular ratio of oxygen and nitrogen. The composition of the air varies sufficiently to affect its value at different times (Ph. v. Jolly,

W. 6, 520).

The pressnro exerted by the atmosphere upon the oarth's surface, at the sea's level or upon any substance at that level, may be expressed by saying that it is equivalent to a barometric column about 76 centimetres (29.92 inches) high. Now at ordinary temperatures 1 c.c. of mercury weighs 13:58 grams. If we suppose that the base of the mercurial column is 1 sq. centimetro it follows that the weight of the counterbalancing atmospheric column is $76 \times 13.58 = 1032$ grams. This is equivalent to 1473 lbs. upon a sq. inch. It can be readily calculated that the total weight of the atmosphere of this average pressure is about 113 trillions of pounds, or 51 trillion kilos. Allowing for the space occupied by the land above the soa's level, the mass of the atmosphere may be taken as Izuovo part of that of the earth (Herschel).

The heights of the counterbalancing columns of air and increury will of course he in the same ratio as the weights of equal volumes if it be assumed that the air is of uniform tension throughout. The height of this homogeneous atmosphere is between five and six miles: it was first calculated by Rohert Boyle to disprove the conjectures of Kepler and others that the air could not extend beyond a couple of miles or so from the earth's surface.

As the air is an elastic fluid it follows from Boyle's law that its pressure must duminish as we ascend; hence the mercurial column stands lower on a mountain top than in the valley below. The fact that the barometric column is less on the top of an elevation than at the bottom was first noticed in 1643 by Claudic Bereguardi from observations on the tower of Pisa—that is, five years before Perrier made his famous experiments on the Puy-dc-Dôme. The relation between the pressure and density of the air at different altitudes may be seen from the following table:—

Metres above sea level	Buik of sic	Density	Barometer lum.		
0	1 och. metro	1.	760		
5,520	2 ,,	0.5	380		
11,040	4 ,,	0.25	190		
16,560	8 ,,	0.125	95		
22,080	16 ,,	0.0625	47.5		
27,600	32	0.0312	23.8		

A pressure equivalent to the average pressure of the atmosphere at the level of the sea is frequently adopted by engineers and others as a unit of pressure and is styled an atmosphere. In this country an atmosphere is the pressure equal to 29 905 inches of mercury at 32° P. at Loudon, and is about 14 73 lbs. on the sq. inch. In the metric system it is the pressure of 760 mm. (29 922 inches) of mercury at 0°C at Paris, and is equal to 1 933 kilos on a

sq. centimetre. Hence the English 'atmosphere' is 0.99968 that of the metric system.

That the mercury in the Torricellian tube, or barometer as it was tormed by Boyle, is constantly varying in height even at the same place, and that these variations are due to the fluctuating pressure of the atmosphere, appears to have been first clearly recognised by Descartes and by Boyle in 1658. It is, however, only within the last few years that we have acquired any very definite information respecting the distribution of the mass of the atmosphere over the earth. The pressure of the air at any given spot depends upon its relative position on the earth's surface: at this spot it varies also with the season of the year and the hour of the day. According to Buchan, whose isobaric charts are really the foundation of our exact knowledge of the subject, there are two broad belts of high pressure passing completely round the globe, one to the north and the other to the south of the equator. The southern belt of high pressure is nearly parallel to the equator; but the northern belt is more irregular in outline in consequence of the anequal distribution of land and water in the northern hemisphere. Between them is the low pressure of the tropical regions, through the centre of which is a narrow belt of still lower pressure towards which the north and south trades blow. A region of low pressure exists also round each pole; that round the north pole having two distinct centres, one in the north Atlantic, the other in the Pacific: at each of these the diminution of pressure is much below the average of the north polar depression. As regards the seasons, it is found that in January the highest pressures are over the continents of the northern hemisphere, and the lowest pressures are over the northern portions of the Atlantic and Pacitic, S. America and S. Africa, and the Antarctic Ocean. The maximum mean pressure at this time is found in Central Asia where it is 30 4 inches, the minimum is in the N. Atlantic and round 1celand, where it is only 29:34 inches. The area of high pressure passes westwards through central and southern Europe. over the N. Atlantic between the parallels of 5° and 45°, across N. America (except to the North and North West), and over some portion of the Pacific. In July the mean pressure of Central Asia is only 29:468 inches or one inch less than in January. The lowest pressures of the western hemisphere are now to be found over the continents, whilst the highest are over the ocean between 50° N. lat. and 50° S. lat. Pressures are also higher at this time over S. Africa and Australia.

Speaking generally, atmospheric pressure is more regular throughout the year over the ocean than over the land. To the westward of each continent there is at all seasons an area of higher pressure over the ocean than over the land, in amount varying from 0·1 to 0·3 inch. These regions of high pressure extend over about 30° of longitude and attain their maxima during winter. The prevailing winds and the general circulation of the atmosphere are intimately associated with these areas of high and low pressure. Winds, in fact, are caused by the flowing away of air from regions of high pressure to those of low pressure, in accordance

with Buys-Ballot's law, which has been thus expressed by Buchan. 'The wind neithst blows round the space af lowest pressure in oircle returning on themselves, nor does it blow directly towards that space, but it takes a direction intermediate, approaching, howover, more nearly to the direction and acurse of circular curves than of radii to a contre. More exactly the angle ie not a right angle, but from 45° to 80°.'

The mest important of the influences affecting atmospherio pressure during the months are tomperature, and, as a secondary effect of temperature, humidity. By comparing the average pressure during the two months which exhibit the greatest divergence of temperature, viz. January and July, Buchan finds the following general result:—The January pressure exceeds that in July over tho whole of Asia except in tho north east, the highest pressures being near the middle of the continent; over Europe to the eauth and east of a line drawn from the north of Russia to the south of Norway, thence to the north coast of Germany, across France to Bordeaux, along the north of Spain, and passing out into the Atlantic at Corunna; over N. America except in the N. East and N. West. The July pressure exceeds that in January over the whole of the sonthern hemisphere, over the northern portion of the N. Atlantic, and over the northern part of the Pacific. The pressure which is thus removed from Asia, Europe, and America in the northern hemisphere in July is transferred partly ta the southern hemisphere, and partly to the more northerly portions of the Atlantic and Pacific Oceans.1

At all places on the earth's surface where the alternation of day and night exists, the pressure of the atmosphere exhibits a remarkable diurnal variation. Generally speaking, the pressure is highest at about 9 A.M. and 9 P.M., and lewest at about 3 A.M. and 3 P.M., but the exact times vary somewhat with the locality and with the season of the year. The regularity of this variation within the tropics is so great that, as Humboldt remarked, the hour of the day may be approximately ascertained from the height of the mercurial column. This oscillation in atmoepheric pressure is not confined to the sca's level: it takes place with equal regularity at heights of 13,000 feet. Within the tropies the oecillation amounts to about 2.2 nm., but as we approach the poles it decreases, until at 70° N. lat. it is only 0.3 mm. In our latitudes theso horary variations are much less strongly marked than in the tropics, and are usually masked by olimatic disturbances; but by comparing the results of a large number of observations, the fluctuation, which in these islands amounts to about 0.5 inm. on the mean of the year, can be olearly made out. In Paris eleven years' observation shows that the mean baremetric oscillation amounts from 9 a.m. to 3 p.m. to 0.756 mm., and from 3 P.M. to 9 P.M. to 0.373 mm. The amount of the diurnal variation differs during the seasons of the year, being greater in summer

than in wintsr. This peouliar phenomenon has given rise to much discussian, but as yet tho oause cannot be said to be satisfactorily determined. Unlike the oceanio tide, it cannat be ascribed to the influence of the meon, since Bouvard has shown that the portion of the horary oscillation of the pressure of the atmasphere which depends on the attraction af the moon cannot raise the mercury in the barometer at Paris more than 0.018 mm., whilst the total variation deduced from the 11 years' observation amounts to 1:129 mm. The fact that the two maxima of pressure occur when the temperature is about canal to the daily mean, and the two minima when the temperature is at its highest and lewest, has led to the supposition that the fluctuations in pressure are connected with the daily march of temperature, and also with the humidity of the air. Dove, Sabine, and Hopkins have effered explanations based on such connections, but they are insufficient to account for the facts. Lamont and Brown have sought to refer the phenomenon to the magnete-electria influence of the sun, or in other words to connect it with the cause of the diurnal changes in ter-restrial magnetism. There is every reason for supposing that the cause of the dinrnal variation in atmospheric pressuro is in some way dependent on, or originates with, the sun, but that its effects are greatly modified by a variety of local er accidental circumstances, as for example the prevailing winds, the amount of moisture in the air, and the relative distribution of land and

The atmosphere appears to receive its heat (1) from the direct rays of the sun, (2) by tha reverberation of these rays from the surface af the earth, (3) by contact with the ground, and (4) through the influence of aqueous vapour.

Although the air ie nat absolutely diathermanous, the heat received by the air from the direct rays of the sun is the least important of the sources enumerated. We know very little at present as to whether the diathermancy of air varies with its density: that is, we have little evidence to determine whether the absorption of the sun's rays increases as they pass further into an atmosphere compressed by its own weight.

The greater portion of the heat which finds its way into the atmosphere is due to radiation from the carth's surface and to the air being in contact with the ground. The amount of heat thus sent into the air depends to a great extent on the nature of the soil which receives the solar radiations and on its capacity for retaining heat. Hence places in the same lattindes and not very far distant from each other, and in the same condition as regards protection, may have very different mean temperatures on account of the different enpacities of varione soils for absorbing and retaining heat.

Aqueous vapour is one of the most important agents in modifying the temperature of the atmosphere. A relatively large amount of heat is rendered latent in the process of evaporation from the surface of the earth, and becomes sensible on the condensation of the vapour in the upper regions of the air. Aqueous vapour also acts even when in the conditian af a perfect gas

¹ For further details see Buchan, ^c The mean pressure of the Atmosphere and the prevailing Winds over the Globe for the Months of the Year ^c (T.E. 25); also Julius Hann's *Erdlunde; and R. H. Scott's *Elementary Meteorolory.

by retarding the transmission of the sun's rays through the air. As the quantity of aqueous vapour decreases as we ascend through the atmosphere, it follows that the amount of this absorption increases as the sun's rays penotrate

further into the atmosphere.

The temperature of the atmosphere varies with a multitude of oauses, such as the latitude, the season of the year, the hour of the day, the degree of humidity, &c. Among the causes which tend to raiso the temperature of the air may be enumerated: the proximity of a western coast in the temperate zone; the divided configuration of a continent into peninsulas with deeply indented bays and inland seas; the aspect or position of a portion of the land with reference either to a sea of ice spreading far into the polar circle, or to a mass of continental land of considorable extent lying in the same meridian, either under the equator or at least within a portion of the tropical zone; the prevalence of sontherly or westerly winds on the western shore of a continent in the temperate northern zone; chains of mountains acting as protecting walls against winds coming from colder regions: the infrequency of swamps which in the spring and beginning of summer long remain covered with ico; and the absence of woods in a dry sandy soil; finally the constant screnity of the sky in the summer months; and the vicinity of an occunic current bringing water which is of a higher temperature than that of the surrounding

On the other hand, the following causes lower the temperature of the air of a place: elevation above the level of the sea, when not forming part of an extended plain; the vicinity of an eastern coast in high and middle latitudes; the compact configuration of a continent having no littoral curvatures or bays; the extension of land towards the poles into the regions of perpetual ice without the intervention of a sea remaining open in the winter; a geographical position in which the equatorial and tropical regions are occupied by the sea, and consequently the absence under the same meridian of a continental tropical land having a strong capacity for the absorption and radiation of heat: mountain chains whose form and direction impede the access of warm winds; tho vicinity of isolated peaks occasioning the descent of cold currents of air down their declivities; extensive woods which hinder the insolation of the soil by the gital activity of their foliage. which produces great evaporation owing to tho large surface it exposes, and increase the surface that is cooled by radiation, acting consequently in a three-fold manner-by shade, evaporation, and radiation; the frequency of swamps or marshes which in the north form a kind of subterranean glacier in the plains lasting till the middle of summer; a cloudy summer sky which weakens the action of the solar rays; and finally a very clear winter sky favouring the radiation of heat (Humboldt: Recherches sur les Causes des Inflexions des Lignes Isothermes. See also Molin's Grundzilge der Meteorologie).

The temperature of the air varies in different strata of the mass, decreasing generally after a sertain clevation in proportion as the distance

from the earth's surface increases, but it is not possible to connect the diminntion in temperature with the elevation in accordance with any definite law. It is nsually assumed that the temperature falls about 1°C. for every 300 feet of perfectly dry air. As, however, the air invariably contains moisture, which is condensed by cooling and so produces heat, the decromen, may be taken practically at about 1°C. for every 500 feet. This estimate can only be taken as an extremely rough upproximation, for it is obvious that the rate of cooling must be affected by a great variety of causes. Indeed the extensive series of aeronantical observations made at the instance of the British Association showed such great irregularities in the rate of diminution that Mr. Glaisher concluded that no law exists.

The atmosphere always contains free electricity, which is generally positive, that is, of an opposite kind to that of the earth. Atmospherio electricity increases rapidly after sunrise, and reaches its first maximum for the day at about 8 A.M. In general the variation in potential follows the diurnal range of atmospheric pressure. In summer the hours of maxima appear to be 8 A.M. and 10 P.M. and the minima 4 A.M. and 4 P.M. In winter the hours of maximum intensity are 9 A.M. and 10 P.M. und the minima 4 A.M. and 4 P.M. This diurnal variation seems to depend mainly on the degree of humidity of the air, the humid months manifesting the greatest potential. The potential seems to increase from July to January, and then to decrease. According to Everett, the maxima occur in February and October, and the minima in June and November.

In clear weather the air is usually positively electrified; it is only during rain, or more properly speaking when rain begins, that the electricity is negative. On the approach of a storm the air is almost invariably negatively electrified, even when the storm-clouds are at a considerable distance from the place of observation. When rain begins, the drops show negative electricity like the uir. In light rain the potential is moderate, but heavy rain is almost invariably accompanied by a high potential. Dellmann's observations have shown, however, that the air may have a very high potential, extending over many days, without any other ovidences of an approaching storm.

• The sources from which the electricity of the atmosphero is derived are not clearly recognised. De la Rive attributed it mainly to chemical action at work on the earth; Pouillet to the evaporation of water; Volta and Saussure to the inequalities of atmospheric temperature. In all probability atmospheric electricity is not wholly due to any one of these causes: they may all be regarded as contributing to the

amount.

The sun's light in its passage to the earth is partially absorbed and reflected by the atmosphere. Clausius has calculated that of the direct sunlight entering the atmosphere on a clear day 6.4 p.c. is absorbed, 18.6 is reflected and diffusod, leaving therefore 75 p.c. to reach the earth. This light is, of course, refracted in its passage in amount depending upon the density of the air. Each ray entering the atmosphere otherwise than perpondicularly may be

supposed to describe a curve in coming to the earth, and as objects are seen in the tangent of the curve on entering the eye, all celestial bodies not in the zenith appear further removed from

the horizon than they actually are.

The refractive power of dry air free from carbonic acid is the mean of the refractive powers of the oxygen and nitrogen under the pressure which each gas exerts in the mixture. This fact furnishes a proof of the physical nature of the atmosphere; since, as Dulong has shown, the refractive power of a compound gas is not equal to the refractive powers of its components, but is sometimes greater and sometimes less. Moist air is rather less refractive thau dry air: precipitated vapour, as mist or fog, slightly increases the refractive power.

Although many of the more striking physical properties of the air were recognised even in the earliest ages, it is only within comparatively recent time that anything very definite has been known concerning its chemical nature.

It had long been observed that many metals on exposure to fire lost their peculiar lustre, and it was also known that by the prolonged action of heat they were ultimately converted into calces or carthy powders often possessing characteristic colours. The fact that the calx weighs more than the metal from which it was derived was known to Geber, and was well understood by the alchemists of the 16th century. Cardan (1506-1576) in noticing the increase of weight which accompanies the calcination of lead, says that it is due to a gas (flatus) which feeds flamo and rekindles a body presenting an ignited point; and Cesalpinus in the De Metallicis (published at Nuremberg in 1602) ulso states that the 'crasse' which forms ou the surface of lead exposed to heated air contains au acrial substance which increases the weight of the metal. Rey of Perigord seems to have first clearly recognised that the augmentation in weight was due to the action of the air. 'Je responds et soutiens glorieusement que cc surcroît de poids vient do l'air qui dans le vase a été épaissi.' Hooko in 1665 asserted that air contains a principle analogous to if not identical with that contained in nitre, and he seems to have believed that a certain portion only of the air is required to support combustion and respiration. The conception of the complex nature of the air was greatly strengthened by the observatious of Mayow on respiration : his experiments are so precise and his facts so incontestable that, to quote Chevreul, one is surprised that the truth was not fully recognised until a century after his researches. Boylo also considerably extended our knowledge of the physical and chemical constitution of the air in the various treatises which he published between 1672 and 1692.

Two years after the segacity of Rutherford had demonstrated the existence of nitrogen, Priestley obtained oxygen gas by heating the ealx of mercury or red precipitate. The significance of this discovery in its relation to the constitution of the air and true nature of calcination was first clearly and irrefragably demonstrated by Lavoisier. By heating increury in contact with a measured volume of air, he showed that about one-fifth of the volume of

the air is absorbed by the metal with the formation of 'red precipitate,' from which the gas oan be recovered by heating to a still higher temperature, and that the remaining four-fifths had all the properties of the 'mephitic air,' or nitrogen, of Rutherford. This experiment not only demonstrated the compound nature of the air and the character of its constituents, but it also showed approximately the relative quantities in which these constituents were present. It was of course quickly recognised that the active properties of air depended upon oxygen, and it was reasonable to assume that the relative amount of this gas determined the quality of air; hence aroso the art of eudiometry. Priestley, who discovered nitrie oxide in 1772, had observed that this gas became red in contact with the air and that the ruddy gas, unlike nitric oxide, was readily soluble in water. When it was subsequently ascertained that the formation of the soluble red gas was due to the action of oxygen on the nitric oxide, the idea of basing a eudiometric method upon this reaction was suggested by Priestley. Careful experimenters were, however, unable to distinguish air which was reputed to be unhealthy from that which experience had proved to be beneficial and salubrious. Thus, in Priestley's hands, air from the country seemed no better than that obtained from the worst-ventilated workshops of Birmingham. Cavendish, after a critical examination of the method, made numerous analyses of air, 'During the last half of the year 1781,' he says, 'I tried the air of near sixty different days in order to find whether it was sensibly more phlogisticated at one time than another, but found no difference that I could be sure of, though the wind and weather on these days were very various, some of them being very fine and clear, others very wet, and others very foggy . . . On the whole, there is great reason to think that the air was in reality not sensibly more dephlogisticated on any one of the sixty days on which I tried it than the rest.' Cavendish devised a scale of graduation applicable to all nitric oxido eudiometers, by means of which the late Dr. Wilson calculated that the mean of his results furnished the following numbers, expressing the centesimal composition of the air by volume:

> Oxygen . . . 20-833 Nitrogen . . 79-167

100:000

Cavendish concludes his account of these observations by pointing out the character of the information furnished by the endiometer. Etymologically the name was without significance. In so far as the instrument takes eegmisance of the impurity of the atmosphere, it betrays no difference between one specimen of air and another; so that, apparently, there are no degrees of goodness to be measured.... Thus it may be inferred that our sense of smelling can, in many eases, perceive infinitely smaller alterations in the purity of the air than can be perceived by the nitrous test. (Cavendish, 'Account of a New Eudiometer,' T. 1783).

These conclusions were confirmed by Humboldt and Gay-Lussac in their celebrated memoir on the composition of the air, published in 1804. They employed the eudiometrio method of Volta, i.e. explosion with hydrogen, and from an extensive series of analyses made on air collected in the most variable weather they concluded that 100 vols. of air contained 21 of express and 70 of nitrogen

oxygen and 79 of nitrogen. The constant proportion of the two principal constituents of the atmosphere appeared now to be se well established that many chemists, after the recognition of the atomio theory, were inclined to think that air was a definito compound of oxygen and nitrogen. The two main constituents of the air are, however, not present in the simple ratio demanded by the law of Gay-Lussac. There is no evidence of chemical combination on mixing oxygen and nitrogen in the proportion in which these gases are present in air: the properties of the mixture are identical with those of air and are such as might be predicated to result from such a mixture. Moreover, oxygen and nitrogen can be isolated from air by mechanical means, or by taking advantage of the different intestinal movements of the gases. Graham separated the gases by atmolysis, and Bunsen demonstrated that the two gases were absorbed by solvents on which they exerted no chemical action in exact accordance with the law of partial pressures. Lastly the more accurate eudiometric processes of the last forty years have shown that the proportion of oxygen to nitrogen oven in so-called normal air is not absolutely constant. This fact was first clearly demonstrated by Bunsen: in a series of analyses made during January and February 1846, he found that the percentage amount of oxygen varied from 20.97 to 20.84 by volume, i.e. a difference of 0.13 p.c., whereas the error of experiment never exceeded 03 p.c. Even wider variations were found by Regnault in the course of a long series of analyses made on air collected in different parts of the world. In more than 100 analyses of air taken at various times of the year in and about Paris the lowest quantity of oxygen found was 20 913 and the highest 20.999; an extreme difference of 0.086; the experimental error being 0.02 p.c. Air collected from different parts of Europe, from valleys and from the tops of mountains and during different seasons of the year, showed

Angus Smith found similar differences in London air in the course of numerous analyses made during 1869; the percentage amount of oxygen varied between 20.857 and 20.95. That these variations are due to local or accidental causes in the case of a town is established by the circumstance that the air in the streets is almost invariably poorer in oxygen than the air of the parks and open spaces. As types of normal air, Angus Smith found the following means of numerous analyses of air in Scotland (1863-5):

variations in the amount of oxygen from 20.903

to 21.0 p.c.

Scashore and the heath 20.999
Tops of hills 20.98
Not mountainous 20.978
Forests 20.97

In marshy places the oxygen sank as low as 20.922.

In Glasgow, in a series of 30 analyses the oxygen varied from 20.889 in the closer parts to 20 929 in the more open places. A. R. Leeds found that the air of New York showed variations from 20.821 to 21.029 p.o.; and lastly Jolly found that air in Munich freed from carbonic acid and aqueous vapour varied in weight as much as 9 mgm. per litre, this variation depending upon the direction of the wind. By cudiometric measurement he obtained variations from 20.53 to 21.01 p.c. Southerly winds as a rule showed a relatively low percentage of oxygen. According to E. W. Morley these deficiencies in the relative amount of oxygen arc to be attributed to the down-rush of air poorer in oxygen from the higher regions of the atmosphere. It was conjectured by Dalton and Babinet that air in the upper strata of the atmosphere contained relatively less oxygen than that immediately above the earth.

From Regnault's observations it would seem that sea-air contains slightly less oxygen than land air. The mean of 17 samples collected in the Arctic seas was 20 91, the extremes being 20.94 and 20.85. The mean of all the samples collected at sea was 20.84; in a series of twonty only five showed amounts of 20 96 and upwards. On the other hand, the observations of Lewy indicate that sea-air differs but little in composition from land-air, but that in the tropics it exporiences close to the sea a diurnal variation in the amount of oxygen and carbonic acid, due to the action of the sun's heat in disengaging these gases from the water. Subsequent experiments on the composition of air over the sca have not confirmed these observations so far as the carbonic acid is concerned (vide infra).

We have comparatively little information in regard to the relative quantities of the constituents of the air at great heights. Such experimental evidence as exists seems to indicate that air contains relatively less oxygen in the higher strata than near the surface of the earth.

Very little is known respecting the proportion of ozone in the atmosphere, or of the circumstances which influence its production. The ozonometric methods hitherto devised are incapable of affording accurate quantitative estimations. Air over marshes or in places infested by malaria contains little or no ozone. No ozone can be detected in towns or in inhabited houses.

Houzeau (A. Ch. [4] 27, 5) determines the relative amount of ozone in the air by exposing strips of red-litmus paper dipped to half their length in a 1 p.c. solution of potassium iodide. The paper in contact with ozone acquires a blue colour from the action of the liberated potash upon the red litmus. The iodised litmus paper is preferable to iodised starch paper (Schönbein's test-paper) which exhibits a blue coloration with any reagent which liberates iodine, e.y. nitrous acid, chlorine, &c. From observations made with iodised litmus paper Houzeau concludes that ozone exists in the air normally, but the intensity with which it acts at any given point of the atmosphere is very variable. Country air contains at most relocated of its weight or relocated of its volume of ozone

The frequency of the ozone manifestations varies with the seasons, being greatest in spring, strong in summer, weaker in autumn, and weakest in winter. The maximum of ozone is found in May and June, and the minimum in December and January. In general ozone is more frequently observed on rainy days than in fine weather. Strong atmospheric disturbances, as thunder storms, gales, and hurricanes, are frequently accompanied by great manifestations of ozone. According to Houzeau atmospheric electricity appears to be the most active cause of the formation of atmospheric ozone.

The existence of hydrogen peroxide in air was first established by Meissner in 1863, but we have no knowledge of the proportion in which it is present. All information as to its relative distribution is obtained from determinations of its amount in rain water and snow. The proportion seems to vary, like that of ozone, with the seasons of the year and with the temperature of the air. It is not improbable that the amount of hydrogen peroxide in air is greater than that of ozone, and it is possible that many so-called ozone manifestations are in reality due to peroxide of hydrogen (v. Houzeau, C. R. 76, 491; Schönbein, J. pr. 106, 270; Meissner, J. 68, 181; Schöne, B. 12, 316; 13, 1503).

The amount of aqueous rapour in the air is subject to great variations. It depends principally upon the temperature, on the distance from the equator, and on the level of the sea; on the form in which the aqueous vapour is precipitated; on the connexion between such precipitations and the change of temperature; and on the direction and succession of winds. The air is rarely saturated with aqueous vapour. In our moist climate saturation is sometimes very nearly attained, but in some parts of Central Asia, Russia, and Africa, extraordinary degrees of dryness have been noticed. In these islands the most humid month is January, and the dries is May.

The existence of carbonic acid in the atmosphere was first inferred by Dr. Macbride of Dublin, in 1764, from the observation that quicklime after exposure to the air effervesced on treatment with an acid. From the case with which determinations of its amount may be effected our knowledge of the distribution of atmospheric carbonic acid and of the causes which affect its proportion is probably more precise than in the case of any other constituent of the air. In fresh country air the amount is remarkably constant, and may be stated as about 034 p.c. In large towns and cities it is usually greater; thus Angus Smith, from numerous analyses made in London during November 1869, found as a mean 044 p.c.; in upwards of 70 analyses the proportion fell below ·04 p.c. on only 5 occasions. In Glasgow, Smith found on an average 05 p.e. The amount will of course be affected by any circumstances which interfere with the rapid diffusion of the oarbonic acid produced by respiration and the combinstion of fuel: hence during fogs the proportion is very greatly increased, an amount as high as 0.1 p.c. having been occasionally noticed. Smith gives the following summary of results obtained in Manchester (Air and Rain, p. 52).

In Manchester streets in usual weather .0408 . 0679 During fogs . Where the fields begin . . .0369 In country air the amount of carbonic acid is invariably greater at night than during the day. This remarkable diurnal variation was first pointed out by Saussure (P. 19, 391), and has heen fully confirmed by subsequent observers. Thus, as the mean of numerous analyses made at Clermont-Ferrand, Truchot (C. R. 77, 675) obtained during the day 0353, and during the night 0403 (v. also G. F. Armstrong, Pr. 1880. 343; and Muntz a. Aubin, C. R. 92, 1299). These differences are mainly due to the exhalation of carbonic acid from plants at night. In the air of towns, and in the absence of vegetation, no such diurnal variations can be detected.

The amount of carbonic acid in the cir is not sensibly altered by rain: this indeed would follow from the law of partial pressure. Over the sea the amount of carbonic acid is about '03 p.c., and, contrary to the statement of Lewy, no diurnal variation in the amount can be perceived (Thorpe, C. J. [2] 5, 189). Schulzo (Landw. Versuchs.-St. 14, 366) obtained similar results with sea air at Rostock: the mean of a large number of observations made from 1868 to 1871 was '0292 p.c. No definite change in the amount was observed at different exasons of the year or at different times of the day. Fog and also a fall of snow were often associated with an increase of carbon dioxide (o. also Fittbogen a. Hasselbarth, C. C. 1874. 694).

Very little is known concerning the distribution of carbonic acid in the higher strata of the atnosphere. According to Saussner and Schlagentweit the amount of carbonic acid on the mountains is greater than on the plains: Truchot, however, found only 0203 on the top of the Puy-de-Dôme (1416 m.), and 0172 on the Peak of Sancy (1884 m.), as against 0313 at Clermont-Ferrand. Additional observations are required.

The existence of nitric acid in the air was first inferred by Priestley. The amount, however is so small that it can only be detected in rain -water. Nitroxy-in compounds are occasionally produced during thunder-storms, and it is said that the rain collected during a storm often contains notable quantities of nitrous and nitric acids. Boussingault found that a million parts of rain water contained 0-83 parts of nitric acid. Reichardt found in hailstones collected during a thunderstorm 0.526 parts per million.

According to Barral each hectare at Paris receives annually from the rain about 63.6 kilos of combined nitric acid. Bineau found that 1 litre of rain-water at Lyons contained in winter 0.3 mgm. nitric acid; in spring 1.0; in summer 2.6 mgm.; and in autumn 1.0 mgm. Bobierro found that a cubic metre of rain-water collected at Nantes in 1663 contained on an average 7.36 grams in the upper part, and 5.682 grams in the lower part, of the town (C.R. 1861, 755). Angus Smith (Air and Rain, p. 287) obtained the following results from a large number of observations on rain-water.

					Nitrie	acid
				(pi	s. per n	allion)
Scotland:	inland	countr	y plac	68		0.305
Ireland-	Valentia		•			0.370
England-	sea coa	st cour	try pl	aces		0.371
Scotland	- 11	oour	itry pl	aces,	west.	0.372
99	"		"		east.	0.476
			91	81	verago	0.424
Liverpool			•			0.582
England:		country	place	8		0.749
London:			•			0.840
England:						0.863
Manohosto		n o f 1 8	69 and	1 187	ο.	1.032
Scotland:						1.164
Glasgow		• •				2.436
Glasgow						5.436

The amount of nitrio acid in the rain-water of towns is uniformly greater than in rainwater collected in the country, from which we infer that much of the nitrio acid in the air is due to the exidation of ammonia derived from the docomposition of nitrogenous organic matter.

The ammonia in the air exists partly as carbonate, partly as nitrate and nitrite; ammonia itself being converted into nitrous and nitrie soids and water by ozone. Solicolo observed that a bottle containing hydrochloric acid became coated near the stopper with a film of salamineniao on exposuro to the air. A piece of pipcolay heated to redness and oxposed to the air for a few days yields a perceptible amount of ammonia when reheated: this is not the case if the clay is kept in a stoppered bottle.

The quantity of ammonia contained in the air is extremely variable: the results on record differ from 135 to 0.1 of ammonia (calculated as carbonate) in 1,000,000 parts of air. Frescnius found that a million parts by weight of air contained during the day 0.098 parts of ammonia, and during the night 0.169 parts. According to H. T. Brown the amount ordinarily present is much larger than this: a million parts of country air at a height of 2 metres from the ground contained from 5:1 to 6:08 parts; the same amount of town air contained from 4:06 to 8:73 parts of ammonium carbonate (Pr. 18, 286). Direction of wind appears to have no influence on the amount. The quantity decreases after heavy rain but is restored to the normal amount (about 6 pts. in 1,000,000) in a few hours. Truchot (C. R. 77, 1159) found from 0.93 to 2.79 mgm. per cubic metre in the air of Auvergne, the highest results being obtained on misty days and the lowest on clear days. From observations made on the Puy-de-Dôme, Truchot concludes that the quantity increases with the elevation and is greater in cloudy than in elear air. On the other hand, Muntz and Aubin (C. R. 95, 788), from observations made on rainwater, find that the upper strata of the air contain much less ammonia than air near the surface of the earth. Nitric acid also was entirely absent from rain water collected at an elevation of 2877 m. Lewy (C. R. 91, 94) finds that the air in summer contains invariably larger quantities of aumonia (2.3 mgm. NH, in 1000 o.m.) than in winter (1.7 mgm. in 1000 o.m.).

The proportion of animouia contained in rain water is as might be expected subject to equally wide variations. Lawes and Gilbert

in the country contained from 0-927 to 1-142 pts. of ammonia. Water collected in towns always contains much larger amounts than that oollected in the country. Barral found that 1,000,000 parts of Paris rain-water contained 3-49 pts. of ammonia. Angus Smith obtained 1-07 pts. of ammonia in the rain-water of inland country places in England, whereas the water collected in the inland country places and more sparsely populated districts of Scotland contained only 0.53 pts. of ammonia per million. The rain water of London contained 3.45; that of Liverpool 5:38; that of Manchestor 6:47; and that of Glasgow 9.10 parts per million. The larger proportion in the cities is due to the influence of animal life and to the constant presence of azotised organio matter in the air of thickly populated districts. Dews and fogs and snow always contain larger quantities of ammonia than rain-water. (For references, see Angus Smith, Air and Rain.)

In addition to those substances oxygen, nitrogen, carbon dioxide, ozone, water-vapour, ammonia, and nitrous and nitrie acids-which are the essential and necessary constituents of atmospherio air, it frequently contains a variety of accidental substances such as common salt, alkalino sulphates, and organic matter dead and living, derived from the proximity of the sea and of marshy districts, or to the influences of towns. Moscati nearly 80 years ago observed that the dew condensed on bottles filled with ice and suspended over the rice-fields of Tuscany, whon collected quickly became putrescent and deposited flakes of a body containing nitrogen; and similar appearances were noticed by Rigaud de Lisle in 1812 in the dew collected in the marshes of Languedoc. The water deposited flakes of nitrogenised organic matter and gave with silver nitrate a precipitate which became immediately purple. (Compare A. H. Smco, Pr. 20, 442.) Vogel also observed that the moisture condensed on cold surfaces in inhabited rooms quickly became putrid owing to the presence of organic matter resembling albumin. Angus Smith found that the moisture condensed from breath after standing for some time formed a thick glutinous mass, which was seen under the microscope to be a closely-matted confervoid growth. Between the stalks of the conferve a number of greenish globules were to be seen in a state of constant movement; also various species of volvox accompanied by monads many times smaller. As far back as 1722, Loewenhoeck (Opera omnia, vol. i. 1722) showed that rain-water, even when recently collected, contained infusoria derived apparently from the air. Similar observations wero mado by Ehrenberg and Gaultier de Claubry (C. R. 41, 645). The first attempt to throw light upon the question of the relative distribution of the organisms present in air was made by Pasteur, by subjecting certain putrescible solu-tions to the action of the air obtained from various localities.

Tyndall (Les Microbes, Paris, 1882) has shown that the micro-organisms contained in air are rapidly deposited in the absence of any strong aerial currents. Upon this fact Hesse (Mitheilungen aus dem kaiserlichen Gesund-heitsamte: Borlin, 1884) has basod a method found that 1,000,000 pts. of rain-water collected for quantitatively estimating the relative proportion of micro-organisms in air. The method consists in aspirating air through wide-mouthed glass tubes, coated internally with gelatine-peptone, prepared according to Koch's method, which is afterwards kept at a temperature of 20°-26° for a few days, when the organisms which have been arrested—consisting of the various monads, bacilli, and micrococci, capable of development and growth in the nutrient gelatine-are recognised by the colonies to which they give riso. By means of this method Dr. Percy F. Frankland has made a number of estimations of the micro-organisms contained in the air of towns, and in the country, and in inhabited buildings. By simultaneously oxposing small circular glass dishes partially filled with the nutrient gelatine to the action of the air, a rough estimate was obtained not only of the number of mioro-organisms in a givon volume of the air, but also of the number which fell during a given time on a definite horizontal area. As the mean of a series of observations made ou tho roof of the South Kensington Museum between Jan. and June 1886, it was found that there were 35 organisms in 10 litres of air, whilst 279 was the average number which fell in I sq. ft. in 1 minute. Similar experiments made near Reigate and in the vicinity of Norwich showed an average of 14 organisms in 10 litres of air, whilst 79 fell per sq. ft. per minute. Experiments made in Kensington Gardens, Hyde Park, and on Primroso Hill, gave an average of 24 organisms in 10 litres, and a deposition of 85 per sq. ft. per minute. At St. Paul's Cathedral, 56 organisms were found at the base, 29 in the Stone Gallery, and 11 in the Golden Gallery, in 10 litres of air. At Norwich Cathedral 18 at the base, 9 at a height of 180 ft., and 7 at 300 ft. In inhabited buildings great variations were observed; as a rulo the number of micro-organisms was less than that found in the open air when the air of the room was undisturbed, but roso rapidly when the air was set in motion by draughts or by the presence of many people (P. F. Frankland, Pr.

Angus Smith has sought to base a system of chemical climatology on the examination of rainwater collected under different conditions and at various places. Rain falling through the air over the sca always contains common salt and sulphates, the latter in larger proportion to the chlorides than is found in sea-water. The sulphates increase inland: they seem to be a measure of the products of decomposition, the sulphuretted hydrogen which is evolved in the putrefaction of ecrtain organic compounds being oxidised in tho atmosphere. In largo towns, the amount of the sulphates is greatly increased owing to the combustion of coal containing ironpyrites. Indeed the rain water of large towns is frequently acid from the presence of free sulphuric acid. When rain contains 40 parts per million of free acid, vegetation is rapidly affected. The following analyses by Angus Smith will servo to show the general character of the rainwater (and therefore of the impurity of the atmosphere) in various parts of Great Britain. The results, which are the mean of many experiments, are expressed in parts per million of rain water.

	I.	II.	III.	I₹.	₹.	VI.	VII.
Sea Air:				{	1	1	
Scotland	12.28	8.61		0.74		424	
England	not det.	88.8	0	1.90	not det	. 371	not dot.
1nland	ł		ł	1		1	
cntry. air :			i	1	i	1	
Scotland	3.37	2.06	0	0.53	•04	.305	*264
England	3.99	5.22	0	1.07	.109	.749	*466
Towns:			i o			1	
London	1.25	20.49	3.87	3.45	*205	*840	not det.
Livernt.	10.16	39.59	11.56	5.38	159	582	
Muchstr.	5.83	44.82	10.17	6:47	251	1.032	*4401
Glasgow	8.97	70.19	15.13	9.10	.300	2.438	10.040

I. Amount of hydrochloric acid (chlorides),

II. , sulphuric acid (sulphutes),

III. , acidity (free sulphuric acid),

III. , amounts.

V. , abbunionid ammonia; decomposition

of organio matter,

nitric acid.

veight of oxygen required to oxidise

organio matter (measure of organio

matter and nitrites).

Although the atmosphere is subject to continual change from a multitude of causes, such as the respiration of animals and plants, the combustion of organic matter, various processes in the arts &c., still from its immense mass and minterrupted motion such changes have only the very slightest effect on its composition. Let us very briefly consider the chief circumstances which tend to influence the proportion of its components.

Nitrogen is undoubtedly a primitive substance: no other body occurs in such large quantities as an element. This gas is probably the source of all nitrogenous bodies, in the formation of which it is continually abstracted from the air. A portion only of the nitrogen so abstracted finds its way back to the air as such: the most considerable compensating influence known to us is the nitrogen evolved by volcanocs,

By the respiration of animals and the oxidution of the spent portions of thoir tissuo, by the respiration of plants at night-time, and by the combistion of fuel, large quantities of carbonic acid are being continually added to the atmosphero. Enormous quantities also are evolved from volcanoes and other subterranean sources. Poggendorff has indeed calculated that the amount so added is at least ten times as much as is derived from all other sources put together. Taking the weight of carhonio acid in the air as 06 per cent., it can be calculated from the area of the terrestrial oblate spheroid that the weight of the carbonic acid in the atmosphero is about 3,223,000 x 10 kilos (Le Conte, P.M. [5] 15, 46; v. also E. H. Cook, P. M. [5] 14, 387). At least 50,000 million kilos of carbonic acid are daily added to the air. The main compensating influence is of course the action of growing plants in sunshine; earbon dioxido is also removed directly and indirectly by zoophytes and by certain chemical actions such as the conversion of felspar into kaolin, &c. Sterry Hunt ('Chemical and Geological Relations of the Atmosphoro,' Am. S. 1880) has calculated that a weight of carbonic acid equal to more than twenty-one times that of our present atmosphero would be absorbed in the production from orthoclase of a layer of kaolin extending over the earth's surface with a thickness of 500 metres, an amount ropresenting

but a small proportion of the results of felspathic decay in the sedimentary strata of the

Dumas and Boussingault, in their woll-known memoir on the Composition of the Air (A. Ch. [3] 3) made some interesting calculations on the duration of the supply of atmespheric oxygen. They found that, taking all the knewn sources of diminution, and assuming that tho oxygen disengaged by plants compensates only for the causes of diminution at present unknown, then even in this exaggerated easo three times the amount of exygen thus abstracted would only amount in 100 years to about \$\frac{1}{8000}\$ of the total quantity, an amount barely appreciable by our most exact eudiometric melhods, -T. E. T.

ATOMIC AND MOLECULAR WEIGHTS .-Two theories regarding the ultimate constitution of matter have opposed each other from tho beginnings of philosophy; one asserts that any mass of apparently homogeneous matter is really homogeneous; the other affirms that every pertion of matter of sensible size is built np of a vast number of small particles which are not themselves capable of further sub-division. The carliest exponent of the second theory of whom we possess any definite record was the Greek philosopher Democritus, who donrished about 450 B.o. The doctrines of Democritus were developed by Epicurus, and the teachings of the Epicurean philosophy are preserved in the Latin poem of Lucretius (s.c. 99-55). These early atomists tried to assign to the atoms, of which they said matter is composed, only such properties as should suffice for their presentation in time and space. They taught that nothing exists save atoms and empty space; that the atoms or 'firstbeginnings,' are of many different forms, and different weights, and the number of atoms of each form is infinite; that all change is only combination or separation of alons; and that the atoms are in constant motion. To meet the objection that if a mass of matter is at rest the parts of it cannot be in motion, Lucretius uses the illustration of a flock of grazing sheep with skipping lambs; to one looking from a distance the flock appears as a white motionless patch on the green hillside, but a closer view shows that the parts of the flock are continually changing their positions. Every atom, Lacretius asserts, is indestructible, and its motion is indestructible likewise; if this were not so how could we account for the preservation of fixed types in nature? Rieses always bear roses, and each animal reproduces its like, because the first bsginnings' (or atoms) of which each is composed are the same and are never destroyed or worn out. 'First-beginnings are of solid singleness, and in no other way can they have been preserved through ages during infinite time past in order to reproduce things." Here we see how clearly the early atomists recognised that every event in nuture occurs in accordance with strict laws. Nothing happens by chance, was a fundamental doctrine of these philosophers. 'I . . . teach . . . by what law all things are made, what necessity there is then for them to continue in that law, and how impetent they are to annul

the binding statutos of time.' The way to gain a knowledge of the laws of nature, Lucretius teaches, is to examine natural events. (See for instance the analysis of the effects of the thunderbolt in Book vi. 323-398.) The differences between a hard body such as iron, and a soft body such as air, depend, according to Lucretius, on the motions of the atoms of the two bodies; in the hard body the atoms move to and fro within very small distances, in the soft body they move freely and rebound from each other only at comparatively leng intervals. 'Bodies are partly first-beginnings of things, partly those which are formed by a union of first heginnings.': The latter are produced by the atoms grouping themselves in concilio; this term seems to mean something very like our expression in combination. The properties of the body fermed by the grouping tegether of atems need not resemble the properties of the atoms themselves (see, for instance, Book I. 915-920). Not only must the atoms enter into concilium with each other in order that any kind of matter may be produced, but the properties of the matter thus formed depend on the mutual relations of the atoms; 'it matters much with what others' and in what positions the same first-beginnings of things are held in union, and what motions they do mutually impart and receive.'3

Although this theory was so nearly complete, yet, as taught by Lucretius, it had few of what we now regard as the essential features of a good scientific theory; it was not stated in terms which permitted of numerical applications to actual phenomena. Few or no exact applicatiens of the theory could be made to natural phenomena. It was searcely able to predict events in nature except in a wide and loose way. It savoured too much of a dogma. It was rather a speculation as to what might be the cause of natural occurrences, than an attempt to determine what these causes really were.

The teachings of the Epicurean philosophers were opposed by those of the school of Aristotle. The Aristotelians magnified the names of things and made them as real or even more real than the things themselves; they identified 'modes of predication with modes of existence' (Lange). Matter occupied a foremost position in the Epicurcan scheme of the universe, but by the followers of Aristotle it was regarded only as the 'potentiality of becoming anything or everything. Aristotelianism prevailed in the middle ages and atomism fell more and more into disrepute.

But in 1592, Gassendi, Canon and Provost at Digno in Provence, revived the atomic theory of the Greek philosophers, and attempted to found on it an interpretation of natural events. The influence of Gassendi was continued through Newton and Boyle; the former of whom, as we knew, demonstrated that not only de masses of matter attract each other, but that every particle of each mass attracts overy particle of the other mass with a force varying directly as the masses of the particles and inversely as the square of the distance between the particles. As Newton accepted the atemic conception of the structure of matter, his demonstration of the action of the force of gravitation gave a new

Lucretius, De Rerum Natura, I. 548-550 (Munro's translation).

¹ Id. V. 55-58. * Id. I. 483-4. * Id. II. 1007-9.

point of departure for the theory of atoms. From this time a science of atomic physics became possible. But the difficulty was, and still is, to form a olsar mental pietnrs of the mechanism of the action of the force of gravitation in terms of the atomic conception of matter. Newton gave the mathematical construction, and clearly ssparated this from the physical explanation which belonged to the region of conjecture.

Not much was done, after Newton, to advance the application of the atomic theory until tho early years of the present century, when Dalton mads a serious attempt to determine the cenditions under which the atems of elementary bodies unite to ferm the atoms of compound bodies.

The great advance made by Dalton consisted in his asserting the possibility of finding the relative weights of the atoms of all kinds of matter, and in his demonstration of the method whereby these relative weights could be determined.

Many analyses of chemical compounds had been made before the time of Dalton; the results were usually stated in percentages, and they seemed to have but few mutual relations. Ric ter (1791-1802) had shewn that a definite mass of each acid combines with a specified mass of a given base; he had arranged several acids and bases in order of neutralisation. Fischer, in 1803, published a table of chemical equivalents which expressed the quantities of bases which were of equal value as regarded power of neutralising a constant quantity of a specified acid. Laveisier, Cavendish, and others, had to some extent grasped the conception of the slements combining in definite proportions. They had never doubted that every chemical substance was of definite composition, and that it would be possible by careful analyses of many compounds to find the laws of elementary combinations. Preust had analysed several pairs of oxidss of the same metal; from some of his numbers the law of combining weights might have been deduced, had he stated his results so as to thow the quantities of oxygen in combina-tion with a fixed quantity of metal.

Dalton analysed two compounds of carbon and hydrogen, and found that in one there was twice as much hydrogen as in the other, combined with the same quantity of carbon. He found similar regularities in the quantities of caygen which combined with a specified quantity of carbon, in the quantities of oxygen which combined with a specified quantity of nitrogen, &c. Meanwhile he had been thinking much regarding the nltimats particles of bedies; he had pictured to himself a quantity of gaseous matter as resembling a heap of small shot, as built up of little definits parts or atoms. He saw how the facts of chemical combination he had been studying would halp him to find the relative weights of these small particles. Dalton's genius recognissd the unity which bound together so many diverse physical and chemical facts. He at once stated clearly the quantitative laws of chemical combination and referred these laws to one underlying conception, the conception namely of 'In all chemical investigations it the atom. to accertain the teletive weights of the simples which constitute a compound. But unfortnately the inquiry has terminated here; whereas

from the relative weights in the mass, the relative weights of the ultimate particles or atoms of the bodies might have been inferred, from which their number and weights in various other componnds would appear, in order to assist and to guide inture investigation, and to correct their results. Now it is one great object of this work to show the importance and advantage of ascertaining the relative weights of the ultimate particles both of simple and compound bodies, the number of simple elementary particles which constitute one compound particle, and the number of less compound particles which enter into the formation of one more compound particle.' 1

That he might determine the relative weight of the 'ultimate particle' of an element it was necessary for Dalton to have some means of fixing the number of particles of that element in one 'ultimate particle' of several of its compounds. Thus, masses of hydrogen and oxygen combine in the ratio of 1 to 8; now, if we assume that the ultimate partiele, er atom, of water is 9 times heavier than the atom of hydrogen, ths mest probable conclusion is that one atom of water is formed by the union of one atom of hydrogen, the mass of which is taken as unity. with one atom of oxygen, the mass of which is 8 thues that of the hydrogen atem; but if we cheese to assume that the atom of water is 16 times heavier than that of hydrogen, then the experimental results-1 of hydregen combines with 8 of oxygen, by weight-are most readily interpreted by saying that one atom of water is formed by the union of 2 atoms of hydrogen, weighing 2, with one atom of exygen, weighing 16. We cannot then determine how many times the atem of oxygen is heavier than that of hydrogen unless we have previously determined how many times the atom of the compound formed by the union of hydrogen and oxygen, that is the atom of water, is heavier than the atem of hydrogen.

Dalton framed certain empirical rules rogarding the composition of the atoms of compennds formed by the union of two elements. His principal rules were these: 'If there are two bedies, A and B, which are disposed to combine, the following is the order in which combination may take place, beginning with the most simple. namely:

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1 atom of A+1 atom of B=1 atom of C, binary;
1 atom of A+2 atoms of B=1 ,
2 atoms of A+1 atom of B=1 ,
1 atom of A+3 atoms of B=1 ,
3 atoms of A+1 atom of B=1 ,
6c. &c.
                                                                                                  D, ternary;
                                                                                                F, quaternary;
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'1st. When only one combination of two bodies [elements] can be obtained, it must be presumed to be a binary one, unless some cause appears to the contrary

'2nd. When two combinations are observed they must be presumed to be a binary and a

'3rd. When three combinations are obtained, we may expect one to be a binary, and the other two ternary.

'4th. When four combinations are observed, we should expect one binary, two ternary, and one quaternary, &c. &c.'
'From the application of these rules to the

Dalton, A New System of Chemical Philosophy (1808)

chemical facts already well ascertained, we deduce the following conclusions: 1st. That water is a binary compound of hydrogen and oxygen, and the relative weights of the two elementary atoms are as 1:7 nearly [more correctly 2nd. That ammonia is a binary compound of hydrogen and azote, and that the relative weights of the two atoms are as 1:5 nearly [more correctly 1:4.66]. . . . In all these cases the weights are expressed in atoms of hydrogen, each of which is denoted by unity.'1 But oven if these rules were admitted, it was not always possible to fix the relative weight of an elementary atom; thus, two compounds of carbon and oxygen were known to Dalton, containing, according to his analyses, 2.7 parts hy weight of carbon combined with (i) 7 and (ii) 3.5 parts by weight of oxygen; henco, by rule 2, the first of these is a compound of one atom carbon with one atom oxygen, and hence the atomic weight of carbon is 2.7, and the second is a compound of 2 atoms carbon (=54) with 1 atom oxygen $(=3.5 \times 2)$. But the results of analyses might also be stated thus: (i) 5.4 carbon + 14 oxygen, (ii) 5.4 carbon + 7 oxygen; and the conclusion might be drawn that the first is a compound of 1 atom carbon (5.4) with 2 atoms exygen (7×2) , and the second is a compound of I atom carbon (5.4) with one atom oxygen (7). Both ways of stating the results of experiments would be in keeping with Dalton's rules, but the first would lead to the number 2.7, and the second to the number 5.4, as representing the relative weight of the atom of carbon. Another objection to the Daltonian rules of atomic syntheses was that, although to-day we may know of but one coinpound of two specified elements, to-morrow we may know of several compounds of these elements.

Berzelius centinued the work which Dalton had begun; his aim was to discover the laws of atomic combinations. Why doss a specified element by combining with exygen produce only two or perhaps three different exides? Why do not the elementary atoms centine in a great many different ratios? What are the limiting forms of the compound atoms produced by the union of any specified elementary atoms? Borzelius busied himself with such questions as these. And that he might find some solutions to such questions, Berzelius was obliged to frame empirical rules, as Dalton had dono before him.

The following may be taken as an example of the Berzeliam rules. If an element forms two oxides with twice as much oxygen by weight in one as in the other, relatively to a fixed mass of the element, the atom of that compound which contains the smaller mass of oxygen is to be regarded as composed of one atom of oxygen and one atom of the specified element, and the atom of the other compound is to be regarded as composed of two atoms of oxygen and one atom of the specified element; but if the masses of oxygen in the two oxides are in the ratio 2:3 relatively to a specified mass of the other element, then the atom of the compound with less oxygen is to be regarded, as before, as composed of one atom of oxygen and one atom of the specified element, but the

atom of the compound with more oxygen is to be regarded as composed of three atoms of oxygen and two atoms of the other element.

But such rules were only empirical, and, however satisfactory might be the particular results obtained by their application, it was impossible to rest contented until somo general principle had been attained which should admit of universal application. In the course of his inquiries regarding the syntheses of atoms, Berzelius porformed a vast number of very careful analyses, the results of which firmly established the quantitative laws of chemical combination. These laws (v. Combination, Chemical, Laws of) assert:—(1) that the masses of the constituents of every homogeneous kind of matter stand in an unalterable proportion to one another, and also to the mass of the compound they produce—the mass of the compound being always equal to the sum of the masses of the constituents; (2) that when two elements combine to form more than one compound, the masses of one of the elements which combine with a constant mass of the other element bear a simple relation to each other; and (3) that the masses of different elements which combine with one and the same mass of another element are also the masses of these different elements which combine with each other, or they stand in a simple relation to those masses. laws may all be expressed in the statement that the elements combine only in the ratios of their combining weights, or, in simple multiples of these ratios. By the combining weight of an element is here meant the smallest mass of that element which is found to combine with one part by weight of hydrogen or with 8 parts by weight of oxygen.

As Berzelius was pursuing his investigations into the gravimetric composition of compounds, Gay-Lussae was making experiments on the volumetric composition of gaseous compounds. In 1809 this naturalist was able to prove (1) that the volumes of the gaseous elements which combine to form a gaseous compound stand in an unalterable proportion to each other; (2) that when two gaseous elements combine to form more than one gaseous compound, the volumes of one of the elements which combine with a constant volume of the other element bear a simple relation to each other; and (3) that the volumes of different gaseous clemeots which combine with one and the same volume of another gaseous element are also the volumes of these different elements which combine with each other, or they stand in a simple relation to those volumes. These laws may all be expressed by saying that the gaseous elements combine only in the ratios of their combining volumes, or in simple multiples of these ratios. By the combining volume of a gaseous element is here nicant the smallest volume of that element which is found to combine with one unit volume of hydrogen, and a unit volume of hydrogen is defined to be the volume, at normal temperature and pressure, occupied by one unit mass of this eloment.

Gay-Lussac argued that the ratios of the masses of the combining volumes of gaseous elements are also the ratios of the masses of the atoms of these elements; and the conclusion

was drawn that equal volumes of gaseous bodies. measured at the same temperature and pressure, contain equal numbers of atoms. This conclusion, if admitted, seems to put into our hands a means for finding the relative masses of the atoms of many compounds and hence of many elements. But the application leads to unlookedfor results. Consider the case of hydrogen and oxygen: experiment shows that two volumes of hydrogen-weighing two-combine with one volume of oxygen-weighing 16-and produce two volumes of water-gas-weighing 18; hence, if equal volumes of gases contain equal numbers of atoms, two atoms of hydrogonweighing two-combine with one atom of oxygen-weighing 16-and the product is two atoms of water-gas-each weighing 9. But each of these atoms of water-gas contains atoms of hydrogen and oxygen; now, the atom of oxygen has been shown to weigh 16 times as much as the atom of hydrogen; hence the atom of water-gas contains half an atom of oxygen. Again, consider the case of hydrogen and chlorine: oxperiment shows that one volume of hydrogen-weighing one - combines with one volume of chlorine -weighing 35.5-and that the product is two volumes of hydrochloric acid weighing 36.5; hence, if equal volumes of gases contain equal numbers of atoms, one atom of hydrogen has combined with one atom of ehlorine to produce two atoms of hydrochloric acid. But as each atom of hydrochloric acid is composed of both hydrogen and chlorine, it follows that each atom of hydrochlorio acid is formed by the union of half an atom of hydrogen and half an atom of chlorine. But these conclusions are at variance with the fundamontal definition of the atom, which states that the atom is the smallest mass of a body that can exhibit the properties of that body.

The discovery that gascous olements combine in fixed quantities by volume had done something to advance the study of atomic synthesis, but it had not removed the fundamental difficulty, the difficulty, namely, of finding some generally applicable principle by means of which the relative weights of the ultimato particles, or atoms, of compounds might be determined. This difficulty was overcome by Avogadro. In 1811 this Italian naturalist introduced into chemistry the conception of two orders of small particles-the molecule, and the atom. The molecule of an element or a compound, said Avogadro, is the smallest mass of it which exhibits the properties of that element or compound; the molecule of an element or a compound is formed by the union of smaller particles of matter which we shall call atoms; in the case of the molecule of an element the atoms are all of one kind, in the case of the molecule of a compound the atoms are of two. or more than two, different kinds. As the properties of the molecule of a compound are which compose it, so it is probable that the properties of the atoms which compose it, so it is probable that the properties of the molecule of an element are different from the properties of the atoms by the union of which the molecule is produced. A chemical action between two gases was conceived by Avogadro as being separable, in thought if not in actuality, into two stagos; in the first stage the molecules of the reacting gases are shattered, and in the second stage the parts of these molecules, that is the atoms, are rearranged to form the molecules of the new bodies.

Avogadro modified the generalisation made by Gay-Lussao, and re-stated it thus :- ' Equal volumes of gases, temperature and pressure being the same, contain equal numbers of molecules. The reactions between hydrogen and oxygen, and hydrogen and chlorine, which could not be oxplained by the generalisation of Gay-Lussac, are perfectly consistent with the generalisation of Avogadro. Two volumes of hydrogen combine with one volume of oxygen, and the product is two volumes of water-gas; that is, in terms of Avogadro's statement, 2p molecules of hydrogen, each composed of x atoms, combine with p molecules of oxygen, each composed of x' atoms (x may or may not equal x'), and the product is 2p molecules of water-gas. One volume of hydrogen combines with one volume of chlorine to form two volumes of hydrochloric acid; that is, in terms of Avogadro's statement, p molecules of hydrogen, containing x atoms, combine with p molecules of ohlorine, containing x' atoms (x may or may not equal x'), to form 2n molecules of hydrochlorie acid.

Not only are these, and other, reactions, between gases explicable in terms of the generalisation of the Italian naturalist, but this statement gives us a means of determining the relative masses of the molecules of all gaseous bodies, and also of determining the minimum number of atoms in each of these molecules. That is to say, the generalisation of Avogadro gives us what we could not obtain from the rules of Dalton or Berzelius, or from the generalisation of Gay-Lussac. For it is evident that, if the number of molecules in equal volumes of two gases is the same, the masses of the two kinds of molecules must be in the same ratio as the densitier of the two gases; and hence, if tho density of one of the gases bo taken as unity, the density of the other, in terms of this one, expresses the relative mass of a molecule of this other gas. Let tho two gasos be hydrogen and oxygen; experiment shows that a given volume of oxygen is sixteen times heavier than the same volume of bydrogen; hence, if equal volumes contain equal numbers of molecules, a molecule of oxygen is sixteen times heavier than a molecule of hydrogen. Let us call the mass of a molecule of hydrogen one, then, in order to find how many times greater than the mass of this molecule is the mass of the molecule of any gas, we have only to determine the density of the specified gas in terms of hydrogen as unity; the number expressing the density of the gas expresses also the relative mass of the molecule of the gas. But, further, the generalisation of Avogadro puts into our hands a means whereby the minimum number of atoms in a gaseous molecule may be determined, and hence a means whereby the maximum relative values to be assigned to the masses of atoms may be determined. Consider the mutual action of hydrogen and chlorine, hydrogon and bromine, nitrogen and hydrogen, and oxygen and hydrogen. Having regard only to the volumes of the re-soting gaseoos elements and the volumes of the

gaseous compounds produced, the actions in question may be stated thus :-

(i.) One volume of hydrogen combines with one volume of chlorine to produce two volumes of hydrochloric acid; (ii.) One volume of hydrogen combinee with one volume of bromine-gas to produce two volumee of hydrobromic acid; (iii.) one volume of nitrogen combines with three volumes of hydrogen to produce two volumes of ammonia; (iv.) one volume of oxygen combinee with two volumes of hydrogen to produce two volumes of water-gas.

Now, as equal volumes contain equal numbere of molecules, these statements may be put as follows :-

(i.) p moleculee of hydrogen combine with p molecules of chlorine, and the product is 2p molecules of hydrochloric acid; (ii.) p molecules of hydrogen combine with p molecules of bromine gae, and the product is 2p molecules of hydrobromic acid; (iii.) p molecules of nitrogen combine with 3p molecules of hydrogen, and the product is 2p molecules of ammonia; (iv.) p molecules of oxygen combine with 2p molecules of hydrogen, and the product is 2p molecules of water-gas.

Therefore in (i.) one molecule of hydrogen has produced, by union with chlorine, two molecules of hydrochloric acid, both of which are composed of hydrogen and chlorine; in (ii.) one molecule of hydrogen has produced, by union with bromine, two molecules of hydrobromic acid, both of which are composed of hydrogen and bromine; in (iii.) one molecule of nitrogen hae produced, by nnion with hydrogen, two molecules of ammonia, both of which are composed of nitrogen and hydrogen; and in (iv.) one molecule of oxygen has produced, by mion with hydrogen, two molecules of water-gas, both of which are composed of oxygen and hydrogen. In other words, in reactions (i.) and (ii.) every molecule of hydrogen has separated into at least two parts; in reaction (iii.) every molecule of nitrogen has separated into at least two parts; and in reaction (iv.) every molecule of oxygen has separated into at least two parts.

These parts of molecules are called atoms. If we assume the truth of Avogadro's goneralisation, then the foregoing reactions are moet simply interpreted by saying that the molecules of hydrogen, nitrogen, and oxygen, are each built up or composed of two atoms. As hydrogen is the standard element to which the atomic and molecular weights of all other bodies are referred, we say that the atomic weight of hydrogen is one, and, because of such reactions as those just stated, that the molecular weight of hydrogen is two. But if the molecular weight of hydrogen is two, the molecular weight of oxygen must be 32, the molecular weight of nitrogen must be 28, the molecular weight of hydrochloric acid must be 36.5, the molecular weight of hydrobromic acid must be 81, the molecular weight of ammonia must be 34, and the molecular weight of water-gas must be 18; because oxygen is 16 times heavier than an equal volume of hydrogen, nitrogen ie 14 times, hydrochlorie acid ie 18.25 times, hydrobromic acid is 40.5 times, ammonia is 17 times, and water-gas is 9 times, heavier than an equal volume of hydrogen. By such reactions and euch modee of reason-

specified element or compound which occupies (under the same conditions of temperature and pressure) the same volume as is occupied by these two unit masses of hydrogen. Detorminations of the sp. gravs. of gasee are

ing as these, we arrive at the following practica: definition of the molecular weight of a gaseous

element or compound :- The molecular weight

of a gaseous element or compound is a number which expresses how many times greater than

two unit masses of hydrogen is the mass of the

subject to several sources of error. But the values to be assigned to the combining weights of the elements, that is, the masses of the elements which combine with one part by weight of hydrogen or 8 parts by weight of oxygen, can be determined with great accuracy. Now, it is evident that the molecular weight of an element must be equal to the combining weight of this element or to a simple multiple of thie number, and that the molecular weight of a compound must be equal to the sum, or to a multiple of the sum, of the combining weights of its constituent elements; hence the data which are required for an accurate determination of the molecular weight of an element are (i.) an exact determination of the combining weight of the element, and (ii.) a measurement of the relative density of the element in the state of gas; similarly the data which are required for an accurate determination of the molecular weight of a compound are (i.) exact determinations of the combining weights of the constituent elements. and (ii.) a measurement of the relative density of the compound in the state of gas. Thus, 35.37 parts by weight of chlorine combine with 1 part by weight of hydrogen, therefore the molecular weight of chlorine is n35.37; but a given volume of chlorine is 35.5 times heavier than an equal volume of hydrogen, therefore the molecular weight of chlorine is approximately $35.5 \times 2 = 71$; now, $2 \times 35.37 = 70.74$; hence the molecular weight of gaseous chlorine is 70.74. Again, phosphorus hydride is composed of masses of phosphorus and hydrogen united in the ratio 10:32 to 1, therefore the molecular weight of this compound is n11.32; but this compound in the state of gae is 17 times heavier than hydrogen, therefore its molecular weight is approximately equal to $17 \times 2 = 34$; now, $3 \times 11^{\circ}32 = 33^{\circ}96$; hence the molecular weight of gaseous phosphorus hydride is 33 96.

Having thus arrived, by the helpof Avogadro's generalisation, at a definition of molecular weight, and having determined that the molecules of hydrogen, nitrogen, and oxygen, and of come other elements, are very probably composed cach of two parts or atoms, we proceed to find an exact meaning for the term atom. If the atom is assumed to be the ultimate portion of any homogeneous kind of matter of which cognisance is to be taken in chemistry, then it is evident that a molecule of a compound gas, formed by the union of (say) three elements, A, B, and C, must be formed by the union of at least one atom of the element A, one atom of the element B, and one atom of the element C. In general terme, no molecule of a compound gas can be formed by the combination of lees than a single atom of each of the elemente by the union of which the compound in question is

atom of an element is the smallest mass of that element which combines with other atoms to produce a molecule.

We cannot as yet determine the absolute mass of the atom of any element, but we have agreed to call the mass of an atom of hydrogen unity, and to represent the masses of the atoms of other elements in terms of the atom of hydrogen; hence we arrive at the practical definition of the maximum atomic weight of an element as follows:---

The maximum atomic weight of an element is a number which expresses how many times greater is the smallest mass of that element which combines with other elements to form a compound gaseous molecule, than the smallest mass of hydrogen which combines with other elements to

produced. This is equivalent to saying, the 14:485 times heavier than the same volume of hydrogen at the same temperature and pressure; therefore the relative density of any gas referred to air as unity multiplied by $14.435 \times 2 (-28.87)$ gives the relative density of that gas referred to hydrogen as twice unity, that is, gives (approximately) the unolecular weight of the gas. Let it now be required to determine the atomic weight of oxygen; the definition of atomic weight tells that the molecular weights of several gaseous compounds containing oxygen must be determined, that these compounds must be analysed and the results in each case stated in parts by weight of each olemont per molecule of the compound, and that the smallest mass of oxygen thus found in any molecule is to be taken as the atomic weight of oxygen. Here are some of the data which have been thus accumulated :-

Data for determining the atomic weight of Oxygen.

Gaseous compound	Sp. Gr. air = 1	Sp. Gr. × 28.87; i.e. approximate molecular weight	Molecular weight	Analysis, stated in parts by wt. per molecule
Carbon dioxide .	1·53	44·2	43:89	31-92 oxygen + 11-97 earbon
Sulphur dioxide .	2·25	64·9	63:90	31-92 , + 31-98 sulphur
Sulphur trioxide .	2·9	83·7	79 :86	47-88 , + 31-98 ,

produce a compound gaseous molecule, such smallest mass of hydrogen being taken as unity.

The term, and the conception underlying the term, molecule, are applied to compounds and elements alike; the term, and the conception show that this conclusion is incorrect:-

Were these the only known gaseous compounds containing oxygen we should conclude that the atomic weight of oxygen is 31.92, that of hydrogen being unity. But the following numbers

Data for determining the atomic weight of Oxygen.

Gaseous compound	Sp. Gr. alr=1	Sp. Gr. × 28.87; f.c. approximate molecular weight	Molecular weight	Analysis, stated in parts by wt, per molecule
Carbon monoxide	·97	27:97	28·93	15:96 oxygen + 11:97 carbon
Water	·63	18:2	17·96	15:96 ,, + 2 hydrogen
Nitric oxide	1·04	30:0	29·97	15:96 ,, + 14:01 nitrogen

underlying the term, atom, are applied in strictness to elements only.

The foregoing definitions of atomic weight and molecular weight are practical, because they indicate the nature of the data which must be obtained before the atomic or molecular weight of a gaseous body can be found. Suppose it is required to find the molecular weight of oxygon; tho mass of this element which combines with unit mass of hydrogen must be accurately measured; and the relative density of oxygen gas must be determined, the standard of reference being hydrogen taken as twice unity. Now, the relative densities of gases are determined by experiments in terms of air taken as unity; but a specified volume of air is the atomic weight of aluminium:-

These numbers show that at least three compounds exist the gaseous molecule of each of which contains 15.96 parts by weight of oxygen; hence, as no molecule is known containing less than this mass of oxygen, 15.96 is taken as the atomic weight of oxygen. Before, then, the atomic weight of an element can be determined with a fair degree of probability a number of gaseous compounds of the element must be analysed; if only a few gaseous compounds of a specified element are known it is probable that the value deduced, from analyses of these compounds, for the atomic weight of the clement, is too largo; it certainly cannot be too small. Thus, let us consider the data for finding

Data for determining the atomic weight of Aluminium.

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Gaseons compound	Sp. Gr. sir=1	Sp. Gr. × 28·87; f.e. approximate molecular weight	Molecular weight	Analysis, stated in parts by wt. per molecule
Aluminium ohloride ,, bromide ,, iodide		270·0 537·5 780 ·0	266·26 532·54 813·22	51-04 aluminium + 212-22 chlorine 54-04 , + 478-5 bromine 54-04 , + 759-18 iodine

Specific Heats of the Solid Elements.*

Mement	Spec. beat	Temp.	Atomic weight		Ob- cerve	Liement	Spec. heat	Temp.	Atomio weight	Sp. bt. ×at. wt.	Ob-
Lithlam	.941		7.01	8.8	Rg.	Seignion	_		·		
Bervillum	-62	450° to 500°	9-1	5.6	Hs.	crystalline	-0641		76.8	8.7	B.W.
* Boron	7.6	about 1000°	10.9	5.8	Wb.	Bromino	***		,,,,	1 67	₽.₩.
* Carbon	463	9600	11.97	5.5	Wb.	solid	-0643	-78° to -20°	79-76	87	P
Sodium	.293	-34° to+7°	23	8.7	Rg.	" Zirconium	-0688	-10.10-20.	900	8.0	Rg.
Magnsslum	.246	** ** 1	24	5-9	Kp.	11 Molybde-	1 0000] 800	ייס	M.D.
-	.25			6.0	Rg.	num	.0722		95-6	۰ ۰ ۱	١
Aluminium	202	1	27.02	5.6	Kp.	Rhodium	0723			6.9	Rg.
	-214								104	8.0	Rg.
10	225		,,	5.8	ltg.	Ruthenium			104.6	6.4	Bn.
* Billoon	203		26	6.1	Mt.	Paliadium	-0593		106.2	6.6	Rg,
	203	232°	26	5.7	Wb.	Siiver	-056		107-66	6.0	Kp.
Phospho-			1			,,	.0559		l " I	6.0	Bu.
rus cryst.	174	-76° to +10°	30.96	5.4	Rg.	,,	-057		1 . 1	6.1	Rg.
29 11	.169		,,	2.8	Rg.	Cadmium	*0542		112	6-0	Kp.
""	202		,, 1	6.5	Kp.	i I	-0546		i " I	8-1	Bn.
" red	170			5-3	Rg.	,,	0567		",	6.3	Rg.
Sulphur	188		31.98	6.0	D.P.	Indiam	+057		118.4	6.5	Bn.
" rhombie	•163	ļ i	"	5.2	Kp.	Tlu	*0548		117.8	6.5	Kia.
9 9	•171		",	5.5	Bn.	H I	0569		,	6.6	
	176		' <u>"</u> '	5.7	Rg.	"	-0562		"		Bn.
Potassium	166	-76° to +10°	39.04	6.5	Rg.	"	0514		"	6.6	Rg.
Calolum	170	-10 00 +10	39-9	6.6	Bn.	Autlmony	0523		120.0	8.0	D.P.
Titanium	1465	0° to 39)°	46	7:1		Antimony			120.0	6.2	Kp,
Chromlum	.10	0. 00 397	524		N.P.	, ,,	0495		,,		Bu.
Manganese	122			5.9	Kp.	1 " 1	·0506	i	"		Rg.
Iron	112	1	55	6.7	Rg.	!» !	0507	1	,, 1	6-0	D.P.
	114		55.9	6.3	Kp.	Teilurium	*0475		125		Kp.
"			"	6.4	Rg.	۱,,,	.0474	- 1	.,		Rg.
Nickel	110		56.6	6.1	D.P.	Iodine	.0541	- 1	126.53		Rg.
	106			6.3	Rg.	Lantha-	- 1				~~6.
Cobalt	107		59	6.3	Rg.	num	.0449		136-5	6.2	Hd.
Copper	-093	ſ	03.4		Kp.	Cerinm	.0446	ŀ	141		Hd.
,, !	-095		" í	6-1	Rg.	Didymium	.0456	- 1	114		Ha.
,,	·095		I	6.1	Ď.P.	Tungsten	.0334	1	183-6		na. Rg.
Zino	-0932	1	64.9		Kp.	Osminu	0311		193		
,,	-0935	1	,,		Bn.	Irldlum	0326		194		ltg.
"	-0955	1	"	6.2	Rg.	Platinum	0325		195		Rg.
	-093 I	1	"	8.0	D.P.		0324	1	199		Kp.
Gallium	-079	12º to 23º	cë l	5.4	Bt.	n	0314	1	"	6.3	Rg.
Germaninm	-077	0° to 200°	72.3		N.P.	" Gold		i i	197		D.P.
Arseulo	۷ ا	0.00.200	12.3	9.04			0324		197	6.4	Rg.
amorphous	-0762	ł	74.9	5.7		16 Mercury		1		1	
erystalline	083		14.0		B.W.	solid	.0319 -	-78° to -40°	199.6		Rg,
	1614	1	74.9		B.W.	14 Thallium	0335		203.6	6.8	Rg.
"	0822		74.9	6.1	Rg.	Lead	0307		206.4		Rg.
Selenion	V072	ŀ	12	6.3	N.	,,	0315	ľ	,,		Kp.
	.05.40	0504 . 00	1		i	l I	0314				Rg.
amorphous	0746	-97° to +6°	76.6	5.9	Rg.	Bismuth	0305		208		Kp.
crystalline	.0745	-18° to +7°	,,	5.9	Rg.		.0308				Rg.
11	*0782 *0681		,,	6.0	Rg.	Thorium	.0276		232.4		Nn.
				6.6	N.	Uraninm	.028				

_		_			38, 129; 46, 257; 63, 6; 67, 427,
Kp.	79	Kopp	**	1)	A. 126, 362; and do. Suppl. 3, 1 and 289.
N.		NEUMANN	**	17	to 100 140
Bn.	29		**	29	P. 126, 123,
.aa	39	Bunsen	39	22	P. 141, 1.
Wb. D.P.		WEBER	•	-	P. 154, 367 (translation in P. M. [4] 42.
n n			n	**	
D.4.		DULONG AND PETIT	39	-	A. Ch. 10, 395. [181 and 276).
Bt.		BERTHELOT		-	C. R. 86, 786.
Hd.		HILLEBRAND	**	n	
T1 117	**		29		P. 163,71 (transistion in P. M. [5] 3, 109).
B.W.	**	BETTENDORF AND WULLNER	**	**	P. 133, 293,
M.D.	99	MIXTER AND DANA	"		4. 169. 388.
Nn.	**		79	19	
Nu.		NILSON	*	20	B. 15, 2519.
N.P.	-	NILSON AND PETTERSSON	7		Z, P. C. 1, 27,
Mt.	-	MALLET		19	20, 21, 01, 12, 22,
	10				C. N. 46, 178,
Zn.		ZIMMERMANN	-		B. 16, 849.
He.	-	HUMPIDGE	~	10	Pr. 30. 1.
	*	WANT MAY	-	-	FT. 82. 1.

As no other gaseons compounds of aluminium, except these three, have been prepared in a state of purity and analysed, we conclude that the atomic weight of this metal is not greater than 54.04; but as only three gaeeous compounde of aluminium are known, it is not unlikely that the trne value to be assigned to the atomic weight of thie element is $\frac{54^{104}}{2}$ or $\frac{54^{104}}{3}$ or $\frac{54^{104}}{4}$, &o. The greater the number of compounde of a given element which have been gasified and analysed, the greater is the probability that the value thence obtained for the atomic weight of the olement represente the true value of this con-

Avogadro'e generalisation-equal volumes of gases coutain equal number of molecules—places in the hande of chemists an instrument whereby they may determine the relative weights of the molecules of all gaseous or gasifiable compounde and elements, and the maximum valuee to be aseigned to the atomic weights of all elements which form gaseous or gasifiable compounds. But at present the densities of only 14 elemente have been determined in the gaseous etate, and gaecoue compounds of only 42 different elemente have been prepared and analysed. Hence the application of the method introduced by Avogadro is limited. There are two other methods of general applicability for determining the values to be assigned to the atomic weighte of elements; let us consider these methods briefly.

In 1819 a paper was published by two French naturaliets, Dulong and Petit, on the specific heats of 13 solid elements, viz., copper, gold, iron, lead, nickel, platinum, sulphur, tin, zinc, bismuth, cobalt, silver, and tellurium (A. Ch.

10, 395).

The products obtained by multiplying tho generally accepted atomic weights of the nine elements from copper to zinc in this list by the specific heats of these elements, and cub-multiples of the generally accepted atomic weights of the remaining four elements by the specific heats of these elements, had all nearly the same value. Generalising from these results, Dulong and Petit concluded that 'the atoms of all the simple bodies have exactly the same capacity for heat.' Thie generalisation has, on the whole, been borne out by subsequent regearch.

The table on p. 342 contains most of the well-established data regarding the epecific heats of solid elements in so far as direct determina-

tions are concerned.

The values to be assigned to the specific heate of beryllium, boron, earbon, and silicon, have been the subject of many experiments and of much discussion: Nilson and Pettersson (B. 13, 1451; v. aleo C. N. 42, 297) made a series of determinations with a specimen of metallic beryllium containing about 5 per. cent. of beryllium and iron oxides. The following were the most important results :-

Specific Heat of Beryllium (Nilson & Pettereson). Temp. interval Spec. ht. Spec. ht. x 9-1 Spec. ht. x 13-65 0-46 3973 3-6 5-4 0 -100 ·4246 3.86 5.8 0 -214 475 4.26 6.4 6.9 .5055 4.6 0 -300

Nileon and Pettereeon concluded from theee numbers that the atomic weight of beryllium is three elements (P. M. [4] 49, 161 and 278); his

13.65; bnt L. Meyer (B. 13, 1780) showed that the true values for the spec. heat of this metal at varioue temperatures, as calculated from the data cummarised in the proceding table, are as follows :--

Specific Heat of Beryllium (Meyer).

Temp.	Spec. ht.	Increase in spec, ht, for 1°	Sp. ht. x 9·1	Sp. ht. x 13.65
20.20	·3973		3.62	5.43
		.00101		
73.2	•4481		4.08	6.12
•		.00085		
157	•5193	00000	4.73	7.10
201	0100	.00063		, 10
256.8	.5819	00000	5.29	8.94
2000	0010		0 40	0 9 2

These numbers show that the specific heat of beryllium increases as temperature increases, but that the rate of this increase is considerably less for the interval 157° to 256° than for that of 20° to 157°. Humpidge (Pr. 39, 1), working with a specimen of beryllium prepared with great care and containing 99.2 per cent. of the metal and 7 per cent. of beryllium oxide, obtained the following results:-

Specific Heat of Beryllium (Humpidge).

Temp.	Spec. heat	Spec. ht. x 9 1
100°	4702	4.28
200	·542 0	4.93
400	.6172	5.61
500	.6206	5.65

The value approximates to a constant between 450° and 500°. There can now be little doubt that the specific heat of beryllium is considerably larger at high than at low temperatures, that this value is nearly constant at about 500° and upwards, and that at those temperaturee beryllium ie not an exception to the law of Dulong and Petit. (For more details v. Beryllium.)

Very varying values had been obtained for the specific heats of the three elements, boron. carbon, and eilicon, before the researchee of Weber. The following table summarises the chief resulte:-

Specific Heats of Boron, Carbon, and Silicon (Weber's numbers not included).

(Temp. about 35°-55°)
Spec. heat. Sp. ht. x at. wt. Observer, Date.

Boron	•	•			-
amor phous	254		2.8	Kρ.	1864
crystalline	.530		2.6	ılo.	do.
" "	.252		2.8	M.D.	1873
"	.257	•	2.8	itg.	1869
graphitic	.235		2.6	do.	do.
CARBON					
diamond	.143		1.7	B. W.	1868
1)	147		1.8	lig.	1841
"		Temp. 20°-1000°	4.4	Dewar	
graphite	.174		2.1	Kp.	1864
"	.188		2.3	B. W.	1868
"	-198		2.4	Rg.	1866
gas-carbon	.165		2.0	Kp.	1864
"	•186		2.2	B. W.	1868
"	197		2.4	Rg.	1841
"	-32	Temp. 200-1000°	3-8	Dewar	
Surcox					
fused	.138		8-9	Kp.	1864
-	.166		4.6	Rg.	1881
erystalline	165		4.6	Kp.	1864
	-171		4.8	М. D.	1872
**	173		4.8	Rg.	1861
11	-,,			81	

Weber, about 1872, made a careful series of determinatione of the specific heate of these

more important results are presented in the | For this interval of temperature only oan any following table :-

Specific Heats of Boron, Carbon, and Silicon

		Temp. S	ec. heat.	sp. nt.
BORON	crystattine1	-400	1915	2·11
19	"	+77°	2737	3.01
**	"	177°	3378	3.72
"	"	233°	.3663	4.03
	n diamond	-50°	0635	0.76
D	**	+ 10°	.1128	1.35
99	24	85°	·1765	2.12
"	,,	250°	3026	3.63
97	,,	6 06°	1108	5.29
**	1)	985°	4589	5.51
,, 9	raphit e	-50°	·1138	1.37
11	"	+10°	.1601	1.93
,,	**	61°	1990	2.39
91	,,	2 01°	$\cdot 2966$	3.56
**	,,	250°	$\cdot 325$	3.88
91	**	641°	1454	5.35
. ,,	11	978°	·167	5.50
Porous	wood carbon	0°-23°	-1653	1.95
11	**	$0_{\circ} - 99_{\circ}$	$\cdot 1935$	2.07
. ,,	"	0°-223°	·2385	2.84
Buicon	i crystaltised	40°	136	3.81
**	11	+ 57°	1833	5.13
,,	,,	128°	·196	5.50
**	1)	184°	$\cdot 2011$	6.63
**	11	232°	$\cdot 2029$	5.68

These numbers show that the specific heats of horen, earben, and silicon increase as temporature increases, but that, in each ease, the value of this increase for a given temperatureintorval is considerably less at high than at low temperatures. The observed variation in the rate of increase of the specific heat of crystallised boron is nearly identical with the observed variation in the rate of increase of the specific heat of crystallised carbon for equal intervals of temperature up to 230-250°; if it is assumed that this identity remains at higher temperatures, then the specific heat of crystallised boron may be calculated. from the chservations made with crystallised carbon, at tomperatures up to about 1000°. The value thus calculated for the specific heat of boren at 1000° is .50. The specific heat of orystalline silicon attains an almost constant value at about 230°. (For more details v. Boron, CARBON, SILICON.)

Leeking at the determinations of the specific heats of solid elements as a whole, it appears clear that the specific heat of any element varies with the temperature, and that the relatiou between the variation of specific heat and that of temperature differs for each element; and, moreover, that the value of the specific heat of an element depends to some extent on the physical condition of the element. But there eeems cortainly to be an interval of temperathre for which the specific heat of an element attains a constant, or nearly constant, value; this tomperature-interval varies for each element, especially for the non-metallic elements with small atomic weights; for many elements it may he appreximately taken as 0° to 100°(C.).

element be said to obey the law of Dulong and

This law may now be etated in a practical form thus :- The atomic heat, i.e. the product of specific heat, at the temperature-interval for which sp. ht. is nearly constant, into atomic weight, of atl solid elements is nearly a constant, the mean value of which is 64. If this is granted it follows that the atemic weight of any solid element is approximately equal to the quotient

spec. heat; provided that the specific heat of the element has been determined for a considerable range of temperature, and, if the specific licat has been found to vary considerably with variations of temperature, that the determinations have been continued until a constant, or a nearly constant, value has been obtained.

Attempts have been made to determine the specific heats of several elements by an indirect method. The method is based on the generalisatien, $\frac{A.C}{n}$ = a constant (about 6.4); where Λ = the formula weight of a solid compound, C=the specific heat of the compound, and n = the number of elementary atoms in the fermula of the compound. This generalisation has been stated in various forms; the earliest appears to be that given by F. Neumann, in 1831: 'The amounts of chemically similar compounds expressed by their formule possess equal specific heats' (P. 23, 1). The statement is sometimes put thus: 'the molecular heat of a solid compound is equal to the sum of the atemic heats of its constituent elements; by 'molecular hoat' is here meant the product of the specific heat of the compound into the mass expressed by its fermula. The form given above, A.C = a constant, is the cutcome of investigations made principally by Garnier (C. R. 35, 278; 37, 130), and Cannizzare (Bl. 1863, 171).

As an example of the application of this generalisation, to find a value for the specific heat of an element in the solid form, let us take Kopp's calculation of the specific heat of solid chlorine (A. Suppt. 3, 321). The data are these: molecular heats (as defined) of metallic haloid salts: ItCl = 12.8, ItBr - 13.9, RI = 13.4; $RCl_2 = 18.5$, $ItI_2 = 19.4$. In each case R represents one atom of a metal the atomic heat of which is 6.4. The atomic heat of solid bromine = atomic heat of solid iodine = 6.6 (approximately). New, as the metallic chlerides, bromides, and iedides, examined are chemically similar, and as the 'melocular heats' of the similar salts are nearly the same, Kopp has concluded that the atemic heat of solid chlorine is approximately equal to 6.4. This conclusion is in keeping with the observed values; thus: RCl (12.8) - R(6.4) =6.4; $RCl_2(18.5) - R(6.4) = 12.1$, and $\frac{12.1}{9} = 6.05$.

Further data are presented by the following 'melecular heats': KClO₃=24.8, KAsO₃=25.3. The argument here is, that as these values are nearly the same, and as the difference in composition between the two compounds is represented by the exchange of Cl for As, it follows that the atemic heat of solid chlorine is approximately

There is, however, considerable doubt whether the material used by Weber was pure boron.

equal to that of arsenic; but the atomic heat of arsenic, as determined by direct experiment, is 8.1, hence the atomic heat of solid oblorine is approximately equal to 6.1.

This indirect method often leads, as might be expected, to several values for the specific (or atomic) heat of an element. Thus, from determinations of the 'molecular heats' of various oxides and other salts containing metals the atomic heat of each of which has been directly determined to be approximately 6.4, the following values for the atomic heat of solid oxygen are arrived at:

From RO 4.6 RO, 3.7 R_2O_3 4.8 ,, KAsO, 4.2,, KClO, 3.5 (assuming at. ht. of Cl = 6) KMnO, 3.8 The mean of these values is 4.1.

The indirect method of finding the atomic heat of an element is undoubtedly useful, but no great stress can he laid on conclusions arrived at by this method only. It is certain that an erroneous conclusion regarding the value of tho atomic weight of an element may be deduced from measurement of the specific heats of solid compounds of that element. For example, Donath determined the specific heat of uranosouranic oxide to bs 0798 (B. 12, 742); assuming the specific heat of solid oxygen to be 0.25 $\left(=\frac{4\cdot\hat{1}}{16}\right)$, the specific heat of uranium was calculated to be .0497; now $.0497 \times 120 = 5.96$, therefore, as analyses of compounds had proved that the atomic weight of uranium is n120, it was concluded by Donath that the atomic weight of uranium is 120. But puro metallio uranium was prepared shortly afterwards, and the specific heat of this metal was directly determined to be 028; now $028 \times 120 = 3.3$, but $028 \times 240 = 6.6$;

uranium (v. regarding this subject, Kopp, B. 19, 813).
The following statements fairly summariso the results of the determinations of the atomic heats of the elements:

hence the atomic weight of uraninm is much

more probably 240 than 120. The larger value

(240) has been confirmed by the preparation

and analyses of two gaseous compounds of

I. Solid elements, 45 in number, the specific heats of which have been directly determined, and the atomic heats of which are all approxinately equal to 64: Li, Na, Mg, Al, P, S, K, Ca, Ti, Mn, Fe, Co, Ni, Cu, Zn, As, Se, Br, Zr, Mo, Ru, Rh, Pd, Ag, Cd, In, Sn, Sb, Te, 1, La, Ce, Di, W, Os, Ir, Pt, Au, Hg, Tl, Pb, Bi, Th, U...(Cr).

II. Solid elements, 6 in number, the specific heats of which have been directly determined, and vary considerably with temperature, and the atomic heats of which appear to be approximately equal to 5.5: Ga (? inaccurately determined), Be, B, C, Si, Ge.

III. Solid elements, 5 in number the specific heats of which have been indirectly determined and the atomic heats of which are probably approximately equal to 6.4: V, Rb, Sr, Cs, Ba.

IV. Gaseous elements; specific heats in solid form very doubtful, and apparently variable: H, (F), N, O, Cl.

It has been already shown that the applica-

tion of Avogadro's law enables a maximum value to be found for the atomic weight of any eloment which forms one or more compounds gasifiable without decomposition. The maximum value thus found for the atomic weight of aluminium was 51.01; but as this value was based on analyses of only throo gascous compounds, it was asserted that the true value was possibly one-half or one-third, &c. of this number. Now, the specific heat of aluminium has been determined to bs 22; hence, assuming the law of Dulong and Petit, the atomic weight of aluminium must be approximately equal to 30 $(30 \times 22 = 6.6)$; therefore the value $\frac{54.01}{4} = 27.01$ is assigned to the atomic weight of this metal. The maximum values assigned to the atomio weights of iron (111.8), copper (126.8), and gallium (138), by the application of Avogadro's law have, in each case, been halved whon determinations have been made of the specific

Various observations on the connexions between the chemical composition and the crystalline form of solid compounds had been made previous to the year 1819, in which year the 'law of isomorphism' was propounded by E. Mitschorlich; this law was subsequently modified and extended, and in 1821 Mitsoherlich stated it as follows: 'Equal numbers of atoms similarly combined exhibit the same crystalline form; identity of crystallino form is independent of the chemical nature of the atoms, and is conditioned only by the number and configuration of the atoms.' Further research has shown that Mitscherlich's statement was too absolute. On the one hand, many solid compounds are known, the atomic compositions of which are very similar, and which, nevertheless, crystalliso not only in different forms, but in different systems, thus:

licats of these metals.

PbCrO, is monoclinic, but PbMoO, is quadratic; AgCl and AgBr arc regular, but AgI is hexagonal; KNO, is rhombic, but CsNO, and RbNO, are hexagonal.

On the other hand, many solid compounds crystalliso in identical or very similar forms, and nevertheless exhibit unlike atomic compositions; thus the crystalline form of the following salts is the same: K.TiP.H.O., CaTiF.4H.O., K.NbOF.H.O., CuNbOF.4H.O., K.WO.F.H.O., CuWO.F.4L.O. Many ammonium salts crystallise in the same forms as the corresponding salts of potassium, but the number of atoms in one formula-weight of these salts is different. It is indeed somewhat difficult to give an exact meaning to the expression 'isomorphous crystals; by this phrase some naturalists mean crystals any one of which is capable of growing in unmodified form when immersed in a solution of any other (Kopp, B. 12, 900 et seq.); others include crystals belonging to the same system but exhibiting very small differences in the measurements of their angles, e.g. the rhombohedral carbonates of magnesium, calcium, iron, zinc, and manganeso; others even include crystals which very closely resemble each other but yet belong to different systems. The fact that the same compound may crystallise in two. or even thres, distinct forms, further complicates the connexion between isomorphism and

chemical composition; thus, arsenious oxide, As,O., and antimonious oxide, Sb,O., both orystallise in regular octahedra and also in rhombio forms; titanium dioxide, TiO, crystallises in two forms belonging to the quadratic system, but exhibiting very different relations of orystalline axes, and also in a third form, viz. rhombic prisms.

The constituents of isomorphous compounds are sometimes themselves isomorphous; e.g. tho double compounds 3Ag,S.Sb,S, and 3Ag,S.As,S, orystallise in identical forms, and the sulphides Sh.S. and As.S. also crystalliso in identical forms. On the other hand, the constituents of isomorphous compounds are sometimes not isomorphous; e.g. the sulphates of magnesium, nickel, and zine, crystallise in rhombic forms, but the oxides of magnesium and nickel crystallise in regular, and exide of zine, in hexagonal, forms. Isomorphism is sometimes not shown in comparatively simple analogous compounds of two elements, while the more complicated analogous compounds of the same elements crystalliso in identical or very similar forms; e.g. many of the simpler compounds of cadmium are not isomorphous with the analogous compounds of the magnesian metals (Mg, Ca, Mn, Fe, Co, Ni, Zn, Cu), but the comparatively complex salts of eadmium, such as CdSO, K2SO, 6H2O, are usually isomorphous with the analogous salts of the metals named. Hence it is necessary to distinguish strict isomorphism as applied to bodies which exhibit the same or nearly the same crystalline form, from the isomorphism of bodies which, although themselves crystallising in different forms, nevertheless combine with one and the same third body to produce compounds into which they enter as corresponding elements or groups, and which crystallise in tho same forms (v. Kopp, Lehrbuch der Physikal. Chemic, 2, 141). The crystalline forms of several elements have been determined, but the statement that such or such elements form an isomorphons group usually means only that analogons compounds of these elements are for the most part isomorphous (v. Isomoneursm).

Notwithstanding the many qualifying clauses with which any general statement of the connexion between crystallino form and chemical composition must, at present, he guarded, it has frequently been found possible to use the knowledge we have of the connexion in question as a guide in researches concerning the atomic weights of elements. In these cases it is assumed that, as a general rule, those masses of two bodies which can mutually replace each other in compounds without change of the crystalline form of the compounds, or in other words those masses which are crystallographically equivalent, havesimilar atomic compositions. By compounds of similar atomic composition is here meant compounds which are very analogous in their chemical relations, and the formula of which contain equal numbers of atoms, or groups of atoms which react through series of changes as if each were a single atom.

Now, if the atomic weight of a specified element is known, and if experiment shows that the mass of this element expressed by its atomic weight is crystallographically equivalent to x unit masses of another element, it follows

that the value of x is very probably the value of the atomic weight of the second element. Thus, the facts that gallinm sulphate formed a double compound with ammonium sulphate, and that this double sulphate was isomorphous with the alums, indicated that the double snlphate in question was a true alum; hence the general formula which expresses the composition of alums expresses the composition of the double sulphate of gallium and ammonium. The formula in question is X,3SO, M,SO, 24H,O, where M = an alkali metal or thallium; but in common alum $X_2 = Al_2 = 2 \times 27.02$ parts by weight of aluminium; and in gallium alum X, was experimentally determined to be 138 parts by weight of gallium. Hence, as two atoms of aluminium were replaced by 138 unit masses of gallium without change of crystalline form, and as the aluminium and gallium compounds were very similar in their chemical relations, the conclusion was drawn that 138 represents the relative weight of two atoms of gallium; therefore the value $\frac{100}{2} = 69$ was deduced for the atomic weight of gallium. This number was afterwards confirmed by analyses of gaseous gallium chloride, and by determinations of the specific heat of the metal. It was at one time supposed by II. Rose (P. 108, 273) that a metal existed closely allied to, but not the same as, niobinm; but Marignac (A. Ch. 60, 257) found that compounds obtained from this hypothetical metal were isomorphous with the corresponding compounds of tin and titanium, and that the groups of atoms SnF and TiF could be replaced by an atom of Rose's 'hyponiobium' without change of crystalline form. Hence Marignac suggested that 'hyponiobium' was a compound; and, because of various reactions, that it was a compound of niobium and oxygen in the proportion expressed by the formula NbO, where Nb has the value 94. If this were admitted it followed that the groups NbO, SuF, and TiF, were crystallographically equivalent in various compounds; but if so, it also followed, from analyses of the various compounds, that one atom of tin (=117.8 parts by weight), and one atom of titanium (=48 parts by weight), were replaced by 91 parts by weight of niobium in isomorphous compounds; therefore the atomic weight of niobium was 94. This value was confirmed by determinations of the relative densities. and by analyses, of the gaseous chloride and oxychloride of niobinm. In this case the comparison of the crystalline forms of compounds led at once to a determination of the atomic weight of an element, to a proof of the nonexistence of a hypothetical metal, and to the recognition that a body supposed to be an element was really a compound. An analogous case is furnished by Roscoe's researches on vanadium; in this case also the study of isomorphism led to the correct determination of the atomic weight of vanadium, and to the discovery that the body supposed to be vanadism was in reality a compound of this metal with exygen (T. 1868. 1 et seq.).

No practical definition of the atomic weight of an element can be given in terms of the data of isomorphism. The foregoing examples serve to show how these data are applied to supplement those gained by the analyses of gaseous compounds, and by determinations of the specific heats, of the elements.

If the stomic weight of caloium is known, then the isomorphism of the carbonates of Mg. Sr. Ba, Pb, Mn, Zn, and Fc, with the carbonato of calcium, helps to fix values for the atomic weights of these 7 elements; the isomorphism of the sulphates of Co, Ni, and Ca, with sulphate of iron gives data from which values may be deduced for the atomic weights of Co, Ni, and Cu; values are found for the atomic weights of Tl and Hg from considering compounds of these elements isomorphous with corresponding compounds of Pb; similarly, Zn and Cd—Fe, Al, and Cr—form many isomorphous compounds; many manganates are isomorphous with selenates and chromates, some chromates are isomorphous with molybdates and tungstates, permanganates are frequently isomorphous with perchlorates and periodates, hence values are found for the atomic weights of Se, Cr, Cl, and I, and also for Mo, and W; from copper we pass to silver through the isomorphism of Cu₂S and Ag₂S; silver leads on to sodium and the alkali metals on the one hand and to gold on the other hand; the compounds RS. and RAs, are isomorphous, hence conclusions can be drawn regarding tho atomic weight of As, and from this the passago is casy to conclusions regarding the atomic weights of P, V, Sb, and Bi; iron is connected with Ti, and this with Si, Zr, Sn, and Th; lastly, given the atomic weight of Pt, Ir, Pd, Ro, Ru, or Os, values can be assigned to the other metals of this group from a study of the composition of isomorphous compounds of these metals. Thus it is seen how helpful is the study of isomorphism in determining the atomic weights of the elements.

These then are the three generally applicable methods whereby values may be found for the atomic weights of the elements: the method founded on the law of Avogadro; the method based on the study of the specific heats of solid elements; and the method which considers tho relations between the chemical composition and the crystalline form of similar compounds. The first of these methods can be applied to determine the atomic and molecular weights of elements and the molecular weights of compounds, but the application is restricted to bodies which are gasifiable without decomposition; the second and third methods can be applied, strictly speaking, only to find values for the atomic weights of solid elements or of elements which form solid compounds.

All the methods are essentially physical; they are based on physical conceptions, and they are to a great extent developed by physical reasoning.

The conception of the molecule of a gaseous element or compound which is implied in the statement, 'equal volumes of gases contain equal numbers of molecules,' is wholly physical. The image of the molecule which this statement calls mp in the mind is that of a small definite portion of matter 'which moves about as a whole so that its parts, if it has any, do not part company during the motion of agitation of the gas' (Clerk Maxwell). It is when this conception is applied to chemical changes that we shanges the parts of molecules do part company;

thus we are led to the chemical conception of the atom, as a portion of matter smaller than the molecule, and either itself without parts, or else composed of parts which, so far ss we know at present, do not part company during any of the changes which the atom undergoes. Thon we proceed to study the properties of these atoms; and among these properties we seem to find two of great importance; the property namely which is expressed in the statement that the atoms of all solid elements, st certain temperatures, have equal capacities for heat; and the property which may be expressed in the statement that identify of crystalline form among compounds is usually accompanied by equality in the number of atoms of which the chemically reacting masses of these compounds are composed.

But here we ask: Are the molecules of isomorphous compounds built up of equal numbers of atoms? Can the physical conception of molecule, which has been gained by the study of gaseous phenomena, be applied to solid com-pounds? And the auswer at present is: it is those small masses of isomorphous compounds which take part in chemical reactions, which as a rule, are composed of equal numbers of atoms. The physical definition of molecule cannot, in the present state of knowledge, be safely applied to solid and liquid bodies. Thus we seem to arrive at two conceptions, and two definitions, of the molecule. On the one side we have tho physical conception, as that of a small mass of a gaseous clement or compound which moves about as a whole, and the parts of which do not part company during the motion of agitation of the gas; and on the other side we have the chemical conception, as that of the smallest mass of an element or compound which takes part in a chemical chauge, and which exhibits the properties of the specified element or compound.

The first of these definitions holds good whether the small particles of a gas are themselves composed of smaller particles, or are chemically indivisible. The volume occupied by a number of gaseous molecules is independent of the numbers of atoms which by their union form these molecules; in one case a gaseous molecule may consist of a single atom (Hg and Cd), in another case a gaseous molecule may be formed by the union of 2 atoms (Hcl), 3 atoms (H₂O), 9 atoms (C₂H₄O), 11 atoms (C₃H₄O₂), or a much larger number of atoms; but in every case, equal volumes of the gases contain equal numbers of molecules. But we know of no single property of liquid and solid compounds which is similarly independent of the number of atoms forming the atomic complex or reacting chemical unit of the compound.

Let us consider the conception of the chemically reacting unit or collocation of atoms a little more closely. We have already seen that the application of the empirical laws of chemical combination could not lead to final determinations of the atomic weights of elements, because these laws could not enable chemists to determine which of several values should be given to the smallest mass of a compound espable of exhibiting the properties of that compound. The value 8, 16, 24, &c. would be assigned to the atomic weight of oxygen, according as the

'atom' of water-that is, in Daltonian language, the smallest mass of water which exhibits the properties whereby water is dietinguished from all other kinds of matter-was assumed to bo 9, 18, 27, &c. times heavier than the atom of hydrogen. But a study of the properties of water leads to the conclusion that the ' atom' of water very probably contains two atoms of hydrogon and one of oxygen, and that the atomio weight of oxygen is therefore more probably represented by the number 16 than by the number 8. Thus, if 9 grams of water react with chlorine or bromine in sunlight 8 grams of oxygen are evolved, and 86.5 grams of a compound of hydrogen with chlorine, or 81 grams of a compound of hydrogen with bromino, are produced; in the former case, the 36.5 grams of the chlorine compound are proved by analysis to be composed of 35.5 grams of chlorine and 1 gram of hydrogen; in the latter ease, the 81 grams of the bromine compound are proved to be composed of 80 grams of bromine and 1 gram of hydrogen; in both cases the whole of the oxygen of the 9 grams of water is removed from combination with the hydrogen and makes its appearanco as free oxygen. Again, if 9 grams of water are acted on by potassium, 5 grams of hydrogen are evolved, and 28 grams of a compound of potassium, hydrogen, and oxygon, containing 8 grams of oxygen—i.e. all the oxygen originally combined with hydrogen in the 9 grams of water—are at the same time produced; if these 28 grams of the new compound are dried. fused, and, while molten, are acted on by potassium, 5 grains of hydrogen are evolved, and 47 grams of a new compound of potassium and oxygen are produced, which 47 grams contain the whole of the oxygen (i.e. 8 grams) originally combined with hydrogen in the 9 grams of water. These experiments prove that the hydrogen in a specified mass of water can be removed from that mass of water in two equal portions, but, so far as those experiments go, that the oxygen in the same mass of water is either not removed at all, or is wholly removed, from comhination with hydrogen. Hence the conclusion is drawn that the smallest reacting mass of water contains one chemically indivisible mass of oxygen, but two chemically indivisible masses of hydrogen. But masses of hydrogen and oxygen are combined in water in the ratio 1:8; hence, if the smallest reacting mass of water is composed of 2 smallest parts, i.e. atoms of hydrogen, and one smallest part, i.e. atom, of oxygen, it follows that the atomic weight of oxygen is at least 16, that of hydrogen being unity, and that the relative mass of the smallest reacting portion. that is the reacting weight, of water is represented by the number 18, not by the number 9.

What value is to be assigned to the reacting weight of marsh gas? Masses of carbon and hydregen combine to form marsh gas in the ratio 3:1; hence the value we are seeking cannot be less, but may be greater, than 4. If 4 grams of marsh gas are acted on by ohlorine, a series of 4 compounds is produced; tho first of these compounds contains ollorine and hydrogen combined with carbon, the masses of carbon and hydrogen being in the ratio 3: 75; the eccond and third contain the same three elements, in the second the carbon and hydrogen are in the

ratio 8: 5, and in the third in the ratio 8: 25; the fourth is a compound of the whole of the earbon originally combined with hydrogen in the 4 grams of marsh gas with chlorine, and contains no hydrogen. If now 4 grams of marsh gas are burnt in a plentiful supply of oxygen 11 grams of carbon dioxide are produced, or if the same mass of marsh gas is burnt in a limited supply of oxygen 7 grams of carbon monoxide are produced; in each case the oxide of carbon formed contains the whole of the carbon originally combined with hydrogen in the 4 grams of marsh gas used. No compound has yet been obtained from 4 grams of marsh gas containing a smaller mass of carbon than was originally present in the marsh gas, i.e. containing less than 3 grams of carbon. The conclusion drawn from these experiments is that the smallest mass of marsh gas which can take part in chemical changes is itself most probably composed of at loast 4 atoms of hydrogen combined with at least one atom of carbon; but if this is granted it followe that an atom of carbon is 12 timee heavier than an atom of hydrogen, and that tha reacting weight of marsh gas is represented by a number certainly not smaller than 16.

Wo have thus determined, on chemical grounds and by chemical reasoning, the following values for the atomic weights of two elements: (H=1) C=12, O=16. Now let us consider a compound of these elements. The simplest formula that can be given to acctic acid consistently with the values H=1, C=12, O=16, is CH.O. If this acid is neutralised by soda, and the sodium salt so formed is analysed. this salt is found to be composed of the same masses of carbon and oxygen, combined with \$ the mass of hydrogen, which were present in the mass of acid used; hence the smallest reacting mass of acetic acid must contain at least 4 atoms of hydrogen. But if this is granted it follows, from the fact that the elements are combined in the ratio C:2H:O, that this smallest reacting mass must also contain at least 2 atoms of carbon and 2 atoms of oxygen, and that the formula expressing the composition of the reacting weight of the acid in quostion must be written C₂H₄O₂. Further evidence in support of this conclusion is afforded by the preparation of thio-acetic acid, which is composed of earbon, hydrogen, oxygen, and sulphur, the earbon and hydrogen being present in the same ratio as in acetic acid, but the oxygen being present in the ratio of 16 to 4 hydrogen (i.e. O:4II), and tho sulphur in the ratio 32 to 4 hydrogen. Now the atomic weight of sulphur is almost certainly 32; hence the simplest formula which expresses the composition of the reacting weight of thio-acetic acid is C₂H₄OS. In this case, ½ of the oxygen of the reacting weight of acetic acid is replaced by sulphur without any further change in the composition of the said; hence, there must be at least 2 atoms of oxygen in the reacting weight in question, because atoms are (by definition) chemically indivisible.

This is an example of the general proposition

that when $\frac{1}{n}$ of a constituent element of the

reacting weight of a given compound can be replaced by another element without any other shange in the composition of the original substance, it follows that the reacting weight in question must contain at least n atoms of the element which has been removed; and that if the atomic weight of the replacing element is known, it is easy to calculate, from the composition of the original substance, tho masses of the other constituents which must be present united with the n atoms of the specified element, and hence to assign a minimum value to the reacting weight of the original substance.

When a formula has been assigned to a compound by such chemical methods as those now sketched, it is frequently possible to argue from this to the formulæ of similar compounds. Thus, the properties and the methods of formation of sulphide of hydrogen show that this compound is similar to oxide of hydrogen (water); but if the reacting weight of water is represented by the formula H.O, that of sulphuretted hydrogen is probably represented by the formula H2S; again, the marked analogies between the sulphide, sclenide, and telluride, of hydrogen suggest that these compounds have similar compositions; but if the first named is H.S, the others are probably H.Se, and ILTe, respec-tively. If these formulæ are admitted, values are at once found for the atomic weights of the three elements, supplier, sclenion, and tellurium. Again, the metal magnesium reacts with water in the ratio of 24 parts by weight of the metal to 18 parts by weight of water, the products of this action being, (1) an oxide of magnesium containing 16 parts by weight of oxygen united with 24 parts by weight of magnesium, and (2) two parts by weight of hydrogen; hence, as the reacting weight of water is represented by the formula H.O. that of maguesium oxide is probably represented by the formula MgO, where Mg = 24 parts by weight of magnesium; and hence the atomic weight of magnesium is probably 24.

The chemical methods for determining the atomic weights of elements then lead to a definition of atomic weight which may be stated thus: the atomic weight of an element is a number which tells how many times greater is the smallest mass of that element found in the chemically reacting weight of any of its compounds than the smallest mass of hydrogen found in the chemically reacting weight of any compound of hydrogen, such smallest mass of hydrogen being taken as unity. The difficulty in applying this definition lies in the vagueness of the expression 'the chemically reacting weight of a compound.' This expression cannot be defined; the illustrations already given indicate the interpretation usually put upon it, and also the methods whereby approximately accurate values are obtained for it in special cases.

The physical conception of molecule is clear, and admits of being put into words which have a definite quantitative meaning; this conception leads to that of the atom, the definition of which may also be put into a quantitative form. But the definition of the molecule is strictly applicable only to gases; hence arises the need of a subsidiary definition. We conceive chemical changes occurring among liquid and solid bodies as occurring among the smallest particles of these bodies which are capable of existing as

bodies in question. These smallest particles we may call the chemically reacting units, or the reacting weights, of the bodies; they are generally called molecules; but if we use this term we must not forget that it is omployed in a somewhat vague manner, and without the strict quantitative signification which is attached to it when we speak of the molecule of a gas.

It seems probable that the mass of the chemically reacting unit of a compound varies, within certain not very wide limits, in different reactions. This mass must of course always be expressed by a whole multiplo of a certain number; but it is probable that the value of the multiple varies. Thus many of the reactions of potassium permanganate can be simply expressed by assigning to the reacting weight of this salt the formula KMnO; but other reactions indicate that this formula should be doubled and written K2Mn2O2. Again, periodic acid generally reacts as if the smallest particle which exhibits the chemical properties of this acid had the mass 228, and were composed of hydrogon, iodine, and oxygen, combined as shown in the formula H₁IO₆; but some of the reactions of periodio acid are more simply explained by doubling the formula, and writing it $\Pi_{i0} \mathbf{I}_{i0} \mathbf{I}_{i0}$ Indeed, even in the case of gaseous elements and compounds, we have sometimes direct evidence to show that the molecular weight of the gas varies with variations of temperature. Consider, for instance, the following data:-

Spec. orav. of Iodine gas (Air = 1).

DI DOI VIII	ATT OF LOWERING COMM	1
Pressure	Temperature	Sp. gr.
760 mm.	448°	8.74
,,	855	8.07
"	C (1275	5.82
"	2 1470	5.06
76 ium.	(1275 1470 1250	4.72

SPEC. ORAV. OF SULPHUR OAS (Air = 1).

Pressure	Temperature	Sp. gr.
760 mm.	520°	6.62
•	660	2.93
,,	860	2.23
" (cf.	Sulamer, vol. iv.)	

RONG ORAN ON ACETIC ACID GAS (Air .: 1)

DLEG ORYA C	M. W. EYIC VCID GVH	(. x x z x) .
Pressuce	Temperature	Sp. gr.
760 mm.	124°	3.20
**	130	3.11
**	160	2.48
	230	2.09
**	280	2 08
**	338	2.08
**	1775	200

SPEC. GRAV. OF NITROGEN TETROXIDE GAS (Air=1).

Pressure	Temperature	Sp. gr.
125 mm.	-60	3.01
138 ,,	+1	2.84
760 ,,	70	1.93
	135	1.60
11 11	185	1.57
yy 10	200	

The density of iodine gas would be 8.77 if the composition of the molecule were represented by I., and 4.38 if the composition of the molecule were represented by I: the numbers given point to the existence of molecules wholes and of exhibiting the properties of the having the composition I, at comparatively low

temperatures, and having the composition I at high temperaturss when the gas is under a small pressure. The numbers given for sulphur gas suggest the existence of molecules S, at temperatures from b.p. to c. 550°, and of molecules S, from e. 650° to c. 1000°; but more recent results throw considerable doubt on the accuracy of this conclusion (cf. Sulphun, vol. iv.) In the case of acctic acid gas, experiments indicate the existence of two different molecules; the data point to the existence of the molecules C₂H₁O₂ (calculated sp. gr. = 2.08, air = 1) at about 230° and upwards, but to the existence of hoavier molecules, $C_3H_0O_3$ (calculated sp. gr. = 3·12, air = 1), at about 120° - 130° . Lastly, the existence of the molecules N_2O_4 (calculated sp. gr. = 3·18) in gaseous nitrogen tetroxide at low temperatures and pressures, and of tho moleculos NO₂ (calculated sp. gr. = 1.59) at higher temperatures, is indicated by the numbers which represent the observed relative densities of this gas. Tho sp. gravs. of some gases slowly decrease as temperature rises until a value is attained which remains constant throughout a considerable interval, e.g. iodine, acctic acid, nitrogen tetroxide, gases; in other cases the sp. gr. remains nearly constant throughout a considerable range of temperature, and then rapidly docroases until another constant value is reached, which again remains constant for a considerable temperaturo-interval, e.g. sulphur gas (v. Dis-SCOLATION, also ALLOTHOPY, and ISOMERISM). But in both classes of gases the data point to the existence, at different temperatures, of more or loss stable melecules, the mass of the heavier of which is a whole multiple of that of the lighter.

The practical conclusions to be drawn from these facts are, that before the molecular weight of a gas can be regarded as satisfactorily determined, observations of the sp. gr. of that gas must bo mado throughout a considerable range of temporature; and that the number which ropresents the sp. gr. in question for such a range of temperature is to he taken as the basis for calculating the molecular weight of the gas, or it may be in some cases the numbers which represent the sp. gravs., each for a considerable temperature-interval, are to be used for finding the different molecular weights of the gas.

If then the mass of the molecule of a gas may have a different value, and therefore the molecule be composed of a different number of atoms, at a high than at a low temperatureand so far as data goes it seems that the mass of the molecule, if variable, is greater at temperatures near the condensation point than at temperatures far removed from this point—it is at least very probable that, if we carry over tho conception of the molecule from gases to liquids and solids, we must be prepared to regard the mass of the molecule of a liquid or solid compound as considerably greater than that of tho molecule of the same compound in the gascons state. But, in practice, when we speak of the molecular weight of a liquid or solid compound we use the term molecular weight with a meaning different from that which we assign to it when we speak of the molecular weight of a gas. In the latter case the term signifies a small mass of matter, itself built up of smaller parts,

which collides with other similar small masses. rebounds, vibrates, but yst remains intact, when a number of these small parts of matter are heated; in the former case the term summarises a number of chemical data in a convenisht form, and asserts that the number of atoms which are so associated as to act in many changes as a chemical whole, is not lsss than a certain specified number.

The chemical formulæ of solid and liquid bodies do not then stand on the same footing as the formulæ of gases (v. Foamulæ). But the question arises: are these collocations of atoms

which we have called reacting chemical units also the reacting physical units of this or that compound? Are the physical constants of compounds conditioned by the masses of these reacting units? If these questions are answered in the affirmative, it is possible that measurements of some physical constant for a series of chemically similar compounds might enable just conclusions to be drawn regarding the relative masses of the reacting units of these compounds. Many measurements of this kind have been made: but no wide generalisation has yet been found which enables us to determine the relative masses of the reacting units of solid and liquid compounds from a knowledge of the physical constants of these compounds. All the generalisations which have been, or which at present can be, ventured upon, are for the most part empirical: the theory of the grained structure of matter has been developed, so far as it has been developed, only for gases; as regards gases, conclusions can be drawn from the fundamental principles of the theory, and these conclusions can be tested by experiment; but as regards liquids and solids, no such general conclusions can be drawn, and the theory can be used as a guide in experimental research only in a wide and general manner. What is wanted now is therefore not only further experimental determinations of the physical constants of series of chemically similar compounds, but a great development of the general theory of the structure of matter, especially in the direction of applying this theory to liquid and solid bodies (v. Molecula theories, also Physical methods). The great difficulty lies in the fact that most of the physical constants of liquid and solid compounds appear to be conditioned both by the nature and number, on the one hand, and by the modes of combination, on the other hand, of the atorus which form the atomic comploxes we have called reacting chemical units. But the kinetic theory of gases has been chiefly developed from the study of properties which are independent of the nature and number, and are conditioned only by the states of union, of the parts of molecules.

But although we must for a time be content with the conception of the ohemically reacting unit of a liquid or solid compound, and although we may at times wistfully contrast this with the clear physical conception of the molecule of a gas, yet there is one well-established chemical generalisation by the application of which values may be obtained for the atomic weights of many elements. This generalisation may be stated thus:—The properties of the elements vary periodically with variations in the atomic

weights of the elements; or thus: - If the elements are arranged in order of increasing atomic weights, the properties of the elements vary from element to element, but return more or less nearly to the same values at certain fixed points in the series. Let the elements be arranged in the order of their atomic weights, from hydrogen to uranium; let them be divided, broadly, into series of sevens; let the second series be placed under the first, the third under the second, and so on; then the elements contained in any one vertical column are called a group, and these in any one horizontal column are called a series. In this arrangement hydrogen is placed in a series by itself, and under it, that is in the same group, is placed the element (lithium) which comes next after hydrogen in order of increasing atomic weight; certain gaps are also supposed to occur in the list of elements, so that an element which immediately succeeds another in order of increasing atomic weights is sometimes placed, not in the group immediately succeeding, but in the group next but one or next but two &c. after that which contains the element with the smaller atomic weight. Thus uranium (210) comes after thorium (232) in order of atomic weights; therium is placed in group IV.; but uranium is placed in group VI. Certain elements are also placed in an eighth group by themselves; and the last member of each series in this group is repeated as the first member of the next series in group I.

The following table shows the arrangement of the elements in accordance with the periodic law. The fermula at the head of each group represents the composition, either of the highest, or of the most characteristic, exide of the elements belonging to that group; in each ease the formula gives the number of atoms of oxygen referred to two atoms of the element.

parts, and to examine the nature of the connexion between the atomic weights, and such measureable properties, of the elomonts, as atomio volume

atomic weight (i.e. the quotient S.G. of solid element)

position in electrical series, fusibility, composition of oxides, chlorides, &c., wave lengths of characteristic lines in the spectra, heats of combustion or of combination with chlorine, &o. &c. The expression 'properties of the elements' is also to be taken as including the proporties of the compounds of the elements; so that the periodic law asserts that e.g. the melting-points of similar compounds (say of chlorides) vary periodically with variations in the atomio weights of the elements.

The periodic law will be disoussed in detail in the article with that heading; meanwhile suffice it to say that the law rests on a firm basis of well-established facts of diverse kinds. We shall here make use of this law to establish values for the atomic weights of one or two typical elements.

At the time of the publication of Meudelceff's first memoir on the periodic law no elements were known which could be placed in group III. series 4 and 5. Calcium (10) and titanium (48) were known; zine (65) and arsenic (75) were known: calcium and zine evidently belong to the group which comprises magnesium, strontium, cadminm, and barium; titaninm must be placed in the same group as earbon, silicon, and tin; and arsenie could not be separated from phospherus, vanadium, and antimony. Hence two gaps occurred in group III. (series 4 and 5), and one in group IV. (series 5). From considering the difference between the values of the atomio weights of consecutive elements, (I) in

				GR	OUPS.			
Series	I,	т	111.	IV.	v.	V1.	VII,	VIII.
S.	It _a ()	R _i O _i	R_zO_3	R ₂ O ₄	R,O,	R,O,	R ₂ O ₂	$[R_sO_s]$
1 2	H = 1 Li = 7	 Be = 9	B=11		N=:14	O=16	P=19	
3	Na=23 K=39	Mg. =24 Ca = 10	A1=27 Se=44	Si=28 Ti. 48	P-31 V=51	S=32 Cr=52	C1 = 35.5 Mu = 55	Fe=56 N1=58.6 Co=59 Cu=63
δ	(Cu=53) Rb=85	Zu = 65 Sr = 87	(la=69 Y=89	Ge=72 Zr=90	As=75 Nh=91	Se=79 Mo=96	Br=80 (? 100)	{Rh=104 Ru=104.5 Pd=106 Ag=108
7 8	(Ag = 108) Cs = 133	Cd 112 Ba = 137	In=11: Ia=139	Sn .118 Ce=110	Sb=120 Di=141	Te_125 7 149	1 = 127 7 150	? 152—156 4 Elements i
9 10	? 4 Ele ? 170	naents 156 to ? 172	162 ? Yb=173	? 178	Er = 166 Ta = 182	? 167 W=181	? 169 ? 190	(Os=191 Ir=192.5 Pt=194 Au=197
11	(Au = 197)	Hg = 200 ments 220 to	T1=201	Pb = 207 Th = 232	Bt == 208 7 237	? 2 Elements U = 240	212 to 220 7 7 245	

periodic connexion between the atomic weights and the properties of the elements, it is necessary to break up the phenomena connoted by the phrase 'properties of the elements' into bers of groups I., II., and III., and for the first

In order to establish the existence of a | the same series the average value of this difference is about 2 in series 3, 4, and 5 -and (2) in the same group-the average value of this difference for the first, second, and third mem-

and second members of groups IV., V., VI., and VII., is about 15, and for the succeeding members of these groups it is about 23—Mendeléeff assigned the value 44 as approximately that of the atomic weight of the unknown element in serios 4 of grenp III., and the value 69 as approximately that of the atomic weight of the unknown element in series 5 of the same group. Mendeléeff also predicted many of the properties of these two unknown elements from considering the positions they occupied in the 'periodio arrangement of the elements. Thus, the relations of the unknown element with atomic weight 44 to aluminium should be similar to these between (1) calcium and magnesium, (2) titaninm and silicon, (3) vanadium and phosphorus, (4) chremium and sulphur; again the relations between (1) beryllinm and calcium, (2) carbon and titanium, (3) nitrogen and vanadium, (4) oxy gen and chromium, (5) fluorine and mangans ,, ought to be similar to the relations between boren and the unknown element. As regards the properties of the second unknown element with atomic weight 69, it was known that (1) zine is more like magnesium than it is like calcium, (2) arsonic more resembles phosphorus than it does vanadium, (3) selenion shows closer analogies with sulphur than with chremium, and (4) bromine and chlorine are more like each other than either is like manganeso; hence, it was argued, the unknown element (69) will resemble aluminium moro closely than it resembles the other unknown element (44), and more closely than the second unknown clement itself resembles aluminium. The relationships indicated were of course studied in detail by Mendeleeff. Thus, take the pairs of consecutive elements in series 3 and 4; the resemblance between any of these pairs (Na, K; Mg, Ca; Si, Ti; P, V; S, Cr; CI, Mn) is less marked in the higher than in the lower groups. Or, take the two elements in each group belonging respectively to series 3 and 5; the resemblance between any of these pairs (Na, Cu; Mg, Zn; ...P, As; S, So; Cl, Br) is more marked in the higher than in the lower

The relationships examined by Mendelecti were those between atomic weights, fusibilities, atomic volumes, composition of oxides chlorides and other compounds, acid or basic character of oxides, power of forming double salts and composition of these salts if formed, &c. &c. As a result of his study of theso relationships, Mendeléeff tabulated many properties of the two anknown elements. Since the memoir of the Russian naturalist was published, several new elements have been discovered; some of the properties of two of these elements will now be compared with the proporties which Mendeléef asserted ought to characterise the elements belonging respectively to series 4 and 5 of group III.

Mendeleeff's Eka-aluminium (III.-5).

Atomic weight about 69. Readily obtained by reduction. Melting-point low. Sp. gr. = 5.9. Not neted on hy air. Will decompose water at a red heat. Slowly nitacked by acids or alkalis. Will form a potassium alum more soluble,

but less easily crystallisable, than the corresponding aluminium salt.

Oxide = El2O2. Chloride = El2Clat

Gallium.

Atomic weight = 69.

Readily obtained by electrolysing alkaline

M.P. = 30·15°. Sp. gr. = 5·93. Non-volatile, and but superficially oxidised in air at bright red heat.

Decomposes water at high temperatures. Soluble in hot hydrochleric acid, scarcely attacked by cold nitrio acid; soluble in oaustic

Ferms a well-defined alum.

Chlorid $= Ga_{s}Cl_{s}$. Oxid $= Ga_{s}O_{s}$.

Mendeleeff's Eka-boron (III.-4).

Atomic weight about 41.

Oxide Eb₂O₃ soluble in acids; sp. gr. about 3.5; analogous to but more basic than Al₂O₄; less basic than MgO; insoluble in alkalis.

Salts of Eb colourless, and will yield gelatinous precipitates with KOH, K2CO3, Na.11PO,

Sulphate, Eb. 3SO, will form a double salt with K.SO., probably not isomorphous with the

Chleride EbCl, or Eb2Cl, sp. gr. about 2, less volatile than Al₂Cl₆.

Scandium.

Atomic weight = 44.

Oxide Sc.O.; sp. gr. = 3.8; soluble in strong acids; analogous with but more decidedly basic than Al.O.; insoluble in alkalis.

Solutions of Se salts colourless and yield gelatinous precipitates with KOII, K.CO., and Na HPO

Sulphate, Se. 3SO, forms a double salt, not an alum, Sc. 3SO, 3K. SO,

Gallinm and scandium are, therefore, the elements which Mendeléeff named cha-aluminium and cka-boron, and many properties of which were accurately and in detail tabulated by him, while the elements were yet maknown.

Much discussion has of late been carried on, and a great deal of experimental work has been done, regarding the value to be given to the atomic weight of beryllium. Chemists are agreed that the value in question is either (in round numbers) 9 or $9 \times 1\hat{3} = 13.5$; if the former value is adopted, beryllium must be placed in group II. series 2; if the latter value is preferred, the metal must find a place between earbon and nitrogen. If the former value is adopted, the formula of beryllinm oxide becomes BeO; if the latter value is preferred, the formula of this oxide must be written Be₂O₃. The periodic law is a guide in the solution of this problem. Briefly, the law directs us to study the properties of the element itself and the composition and properties of its compounds; to compare these with those of elements which must come in the same group and the same series as beryllium; to eccupare the relations between beryllium and these other elements with the relations which have been established between elements occurring in positions similar to that occupied by baryllinm and the other elements in question; and to adopt

that value for the atomic weight of beryllium which best harmonises with the outcome of this study. There can be no doubt that the value which best learmonises with the results of this study is 9; hence the atomic weight of beryllium is almost certainly 9. This result is confirmed by the application of the law of Dulong and Petit, and also of the law of Avogadro; for the specific heat of beryllium at about 500° is nearly constant and is approximately represented by the number 62 ($62 \times 9 = 5.6$), and the vapourdensities of beryllium chloride and bromide show that the formulæ of these compounds, as gases, are BeCl, and BeBr., respectively (Be = 9).

The atomic weight of tellurium had for long been supposed to be greater than that of iodino (127); but if this were so tellurium must be placed in group I. sories 9; that is to say, in a group which contains the alkali metals. This position cannot be defended; moreover, every chemist knows that tellurium exhibits marked analogies to sulphur But if tellurium is to find a and sclenion. place in group VI. the value to be given to its atomic weight must be greater than 120 and less than 127. In 1883 Branner undertook an experimental criticism of the methods whereby the atomic weight of tellmium had been determined by different chemists. Branner proved that these methods almost necessarily gave too large values; he also made very careful determinations of the atomic weight of the element by two new methods, and obtained a series of numbers varying from 121.94 to 125.4, with a mean value of 125. The periodic law has, thorefore, prevented chemists from finally adopting an erroncous value for the atomic weight of tellurium, notwithstanding the great weight of authority which was in favour of regarding that value as correct.

These examples will suffice to show how the periodic law is used as a guide in determining what multiple of the combining weight of an element is to be adopted as the atomic weight of that element. Incidentally, these examples also impress us with the extreme importance of the constants which we call the atomic weights of the clements. Given this constant for a new element, and we may, to a considerable extent, predict the properties of the element and its compounds. The periodic law also enables values to be given, if not to the molecular weights, then certainly to the reacting weights of compounds; because the position of an eloment in a group and series determines the formulæ of the salts of that element, and, as we assume the atomic weights of the other elements in these salts to be known, therefore determines the relative masses of the chemically reacting units of these salts. There are at least one or two elements in each group which form some gasifiable compounds; the molecular weights of these compounds are therefore known; hence conclusions may tentatively be drawn regarding the molecular weights of similar compounds of other elements in the same group. But no great stress must be placed on such reasoning as this. Aluminium and indium occur in group III. (series 3 and 7), these metals exhibit fairly marked analogies; yet the molecular formula of gaseous aluminium chloride is Al₂Cl, while that of gaseous indium chloride is InCl.; thallium Vot. I.

belongs to the same group as aluminium and indium (series 11), yet the formula of the only chloride of thallium which is stable as a gas is TICI.

There is then at present one generally applicable method for determining the molecular weights of gaseous elements and compounds; this method springs ont of the application of the generalisation of Avogadro to chemical changes occurring between gascous elements. The application of the generalisation in question leads to practical definitions of the terms molecular weight and atomic weight. In addition to this method there are three others which servo to determine, more or less accurately, the values of the atomic weights of the elements; and two of these are also employed to find the relative masses of the small particles of solid and liquid compounds which take part in ohemical changes.

The methods founded respectively on the laws of Avogadro, Dulong and Petit, and Mitscherlich, are essentially physical methods; they are outcomes of the physical theory of the grained structure of matter. The applications of this theory to chemical phenomena which have been considered in the present article have been treated in a purely empirical manner. But it is possible to deduce the law of Avogadro from the first principles of the theory in question. The theory assumes that the temperature of a gas represents the mean kinetic energy of the molecules of that gas; hence, if M and M1 represent the masses, and V2 and V,2 the mean squares of the velocities, of the molecules of two gases at the same temperature, it follows, from the laws of energy, that $MV^2 = M_1V_1^2$.

But if the pressures of the two gases are equal,

 $MNV^2 = M_1N_1V_1^2$;

where N and N, represent the number of moleenles in unit volume of the two gases; because, according to the theory, the pressure of a gas on the walls of the containing vessel is an effect of the impacts of the molecules of the gas, and this depends on the number and velocity per unit of time of these molecules. From these oquations it follows that

 $N = N_1$;

that is, when two gases are at the same pressure and temperature the number of molecules in unit volume of either gas is the same. But this is the law of Avogadro.

Neither the law of Dulong and Petit, nor the law of isomorphism, can as yot be satisfactorily deduced from the first principles of the molecular theory. We know very little, if anything, of the structure of gaseous molecules; and of the molecule: phenomena of solids our accurate knowledge may be said to be almost nothing (v. Accaecation, States or, p. 87; also Molecular STRUCTURE OF MATTER, THEORIES REGARDING; also Physical METHODS APPLIED TO CHEMICAL PHENOMENA).

The atomic weights of all the known elements have been more or less accurately determined; but only fourteen elements have been gasified, and hence the molecular weights of only fourteen elements have been determined. The molecules of the greater number, but by no means of all,

these elements are most probably (v. remarks on 840 regarding the molecules of hydrogon, &c.) and two atoms; they are diatomic, he following table shows the classification of the following tables are classification of the following tables are classifications.

Atomicity of Elementary Molecules (the temperatures are approximate).

Monatomic	Diatomi c	Triatomic	Tetratomic	Hexatomic
odium otassium ino admium lercury dine at c. 1500° 'Bromine at c. 1800°) ntimony at c. 1700°	Hydrogon Chlorine Bromine Iodine at 200°-1000° Oxygen Sulphur at 800° and upwards Selenion at 1200° and upwards Tellurium Nitrogeu Phosphorus Arsenio heat	Oxygen as ozone Selenion at 700°-800°	Phosphorus Arseuic (both at temps. nearly up to white heat)	Sulphur at 450 550° (very doubtful v. Biltz a Meyer, B. 21 2013)

The following table presents the data available for calculating the molecular weights of the lemontary gases :--MOLECULAR WEIGHTS OF ELEMENTARY GASES.

I.	11.	111.	1V.	v.	I,	JI.	117.	IV.	₹,
Name of element	Relative density air=1	Temp. of observation	Density × 28°87	Molecular weight	Name of element	Relative density air=1	Temp, of observation	Density × 28.87	Molecular weight
Hydrogen	06926	0°	2	2	" Bromine	5.51	1000	159-9 լ	159.5
*Soilium	'87	1200^-1500°	25.5	23	18 ,,	5.38	100°	155.3	1000
Nitrogen	0.9713	0.	28:04	2802	19 17	4.43	abt 1500°	117:9	?
Oxygen	1 106	abt. 1400°	31.94	31.92	20 Seleniou	5.68	abt. 1400°	161.1	157.6
, ,,	1.10563	00	31.92	3132	21 11	6.37	abt, 1000°	183.9	?
"(ozone)	1.658		47.86	47:88	20 11	7:67	860,5	221.4	236 4
Sulphur	2.23	860°	64.1		23 Mercury	6.96	abt. 1000°	200.93	
, ,	2.24	10400	64.6	63.96	24 19	6.98	446°	201:5	199-8
٠ "	2.17	abt. 1400°	62.6		26	7 03	4242	203.0	1000
Zine	2:38	abt. 1400°	68.7	61.9	26 11	6.7	8820	193.4	ĺ
* Chlorine	2.45	200°	70:73		" Indine	8.8	250"-150"	2540	
٠.,	2.61	abt. 1000°	75:35	70.71	.* ,,	8.72	1850	251.7	
·•	2:44	abt. 1200°	70.72			8.70	4479	251.2	253.03
* Cadmium	3.94	abt. 1000°	113.7	1121	30 ₃₁	8.72	abt. 1000°	251.7	203 08
Antimoay	9.78	1640°	136-1	120	a) ,,	8.84	2500	255.2	1
• Phosphorus	4:35	500°	125.6		212	8:55	665°	246.8	ļ
٠ ,	4.50	abt. 1000°	129.9	123 84	22 ,,	5.87	abt. 1100°	1694	1
* Arsenic	10-2	860°	294.5	000.4	23 ,,	4:76	abt. 1500°	137:4	[? 126.58]
•	10.65	644°-668°	307.4	299-6	34 Tellarium	9:08	abt. 1400°	262.1	255
0 ₈	6.53	1430°	188.5	[2119:81	l				1

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    Scott, Pr. E. 14, 410.
    V. Meyer, B. 12, 1426.
    Soret, C. R. 61, 941; 64, 901.
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The following table presents a summary of the atomic weights of the elements and of the evidence upon which each value is based :-

Rognault, C. R. 20, 975.

Regnault, Lc.

Regnault, Lc.

N. Meyer, R. 12, 1126.

Sorce, C. R. 61, 941; 54, 90

V. Meyer, B. 12, 1112.

Mensching a. Meyer, B. 19, 3295.

Ludwig, B. 1, 232.

V. Meyer, B. 13, 400.

129 d. B. 15, 2774 (mean of 5 experiments).

Deville a. Troost, C. R. 49, 239.

Bilts a. Meyer, Z. P. C. 4, 249.

Mitscherlich, A. 12, 159.

Mitscherlich, Lc.

V. Meyer, B. 13, 406.

 ¹⁰ Crafts, C. R. 90, 183.
 ²⁰ V. Meyer, B. 13, 1107, 1110 (mean of 6 experiments).
 ²¹ Dumas, A. Ch. (2) 33, 337.
 ²² Mitscherlich, Lc.
 ²³ J. Meyer; a. Meier a. Crafts, B. 13, 868 (mean of 7 extrements). "Y. Meyer; a Meier a. Crafts, B. 13, 868 (mean of 7 experiments).

Dumas, Lc.

Y. Meyer, B. 13, 396.

N. Meyer, B. 13, 1115.

Devillo a. Troost, Lc.

Id. 13, 1010.

Biltz a. Meyer (Z. P. C. 4, 249) have obtained values which point to a mol. w. for phosphorus between P, and P, for bismuth between Bi, and Bi, and for thallium as Tl, at very high temperatures.

ATOMIC WEIGHTS OF THE ELEMENTS.

4	ij	Ħ	1Υ.	V. Atomic Weight		ΛΙ.	_ v11.	WIII.
Rement	Frincipal com- pounds vapour den- sities of which have been determined	Specific heat: how determined	Isomorphism: compounds compared [see Note A, p. 361]	(1) (2) By vapour By sp.heat density method method		Compounds analysed &c. in order to find com- hining weight of the element [see Note B, p. 361]	Com- blning weight [Note B, p. 361]	Remarks [see Note C, p. 361]
Hydrogen	H. HCl. 11Br. Hl. H.S. H. Se. H. Te. H.N. H.P. H.C.	H	1	-				
Lithium	&c. None	heat abnormal?] Directly		1	10-2	· Lithium chloride	7.01	
Beryllium.	BeCl	Directly: sp. heat waries unuch with tempera-	-17	80-3	80.6	2 Beryllinm sulphate	75.	
Boron	BF., BCI., BBr., B(CH.),	H	- az pas	10.3	10.9	Borax; boron chloride	3-66	
Carbon	CH. CH.F. CH.C. CH.Br. CO. CO. COCI. COS. CS. COCI. COS. CS. COCI. COS. CS. CH.O. C.H.O.		Direct! 7:50. heat varies CN compounds with those of much with tempera- F, Cl. Br, and 1	11.97	11:97	• Diamond burnt to CO,	\$6. **	
Nitrogen	NH, NO, NO, NO, NO, NO, NO, NO, NO, NO, NO		NH, compounds with those of alkaii metass	14-01	ı	Ammoninm chloride;	4.0	
Oxygen	OH, ON, OC, OCI,P, O,C, O,S, O,Os, &c.	Indirectly: very un- decided (from sp. heats of various compounds)	ı	15.96	1	* Synthesis of water	7-98	
Fluorine	FH. F(CH,), F,B, F,Si, F,P, &c.	Indirectly: very undecided [from sp. heat	FH. F(CH,), F,B. Indirectly: very un. Metallic floorides with analogous F,Si, F,P, &c. of code (from sp. heat compounds of Ci, Br, and I	19-1		Sodium finoride;	19.	
Sodium	None	Directly		1	53	Sodium chloride	 	
Magnesium .	£	*	other alkall metals Mg compounds generally with those of Zn. Mn, and Fe (in	1	ŧ.	• Magnesium sulphate, do. chloride	12	
Aluminium . AICI,	AICI,		ferrous salts) With Cr. Mn, and Fe in B, C, and derivatives	27.03	27-03	* Ammonia - alum; sluminium hromide; solution of alu- minium in soda	2004	
Entertier 368 and 374. Be. Nilson and 1 B. Berzellus, P. C. Dumes and S. Marchan, P. C. Dumes and S. Marchand, J. Pr. 33, 188	"Li, 7. W. Mailet, Am. S. [2] 22, 349. Stat, Nouselles, Remarka, 588 and 574. " Be. Nilson and Perference, R. 13, 1451. " B. Berrellus, P. 2, 139. Deville, A. Ch. [3] 55, 189. " C. Dumas and Sust, A. (A., [3], 5. Edimann and Sust, A. (A., [3], 5. Edimann and Sust, P. 20, 20, 20, 20, 20, 20, 20, 20, 20, 20,		N. Stas, Rapport, 50, 87, 92, and Nouvelles Recherches, 57, 281. O. Refunant and Marchand J. pr. 26, 468. Durna, A. O., [3] St. Durye, C. [3] 28, 291, 301. Durna, A. O.,	87, 92, nd J. pr. 26, 291, 301,	and Nour 468. Dur Durnes, 4.		Pelonze, C. Jacquelain, 56, 310. Du. J. W. Mallet.	Na. Pelonze, C. R. 20, 1047. Dumas, A. Ch. [7] 184. Stas, Rapports, 78, and Journal or Recherchs 348. "Mg. Jacquelain, C. R. 30, 106, and A. Ch. [31 33, 308. Behr, L. Fry. 54, 310. Dumas, A. Ch. [31 58, 189.

atomic and molecular weights.

VIII.	Remarks [see Note C, p. 361]		•					Sc. The atomic weight of this metal is mostprobably 14:68 x3= 4; if this is so, the naride is Sc. 9, and is analogous with the oxides of the earth metals					
ji j	hining weight [Note B, p. 361]		10.32	15-99		33-04	19-95	14.66	13	17-07	50.5	27.5	21.95
. TA	Compounds analyses occinionate to find com- bining weight of the element [see Note B, p. 361]	" Silicon chloride and hromide	synthesis of phos- phorus pentoxide	sulphide; reduction of allersulphate by	"Poussin chlorate; synthesis of silver chloride	" Potassinm chloride,	14 Calcinm chloride;	salcium carouste sulphate	Titaninm chloride and bromide	va Vaosdinm pentoxide,	Potassinm dichro-	mitte Maganese chloride; manganoso - man- ganic oxide; man- ganous oxalate. Sil-	Synthesis of ferrio oxide; reduction of ferricoxide; analysis of ferrous and ferring
	(2) 3y sp. heat method	28.3	30-36	31.93	ı	39.04	33-9	1	\$	1	# 65 20	55	0. 12.
v. Atomic Weight	By vapour light density method	28:3	30-96	31.33	35-37	39-04	1	1	\$	51.3	F.09	2	55.9
IV.	Isomorphism: compounds compared [see Note A, p. 361]	With G, II, Zr, Sn, and Th in compounds of type BO,	Phosphates with vanadates and arsenates; organic compounds of P with those of N, As, and Sb	With Se compounds; with Te compounds of type R'Te. Salts of H,SO, with those of H,GrO,	and Alter, with analogous com- pounds of Br and I	K compounds with those of	other alkali metals Ca compounds with those of Sr,	as, and in some cases rn as and compounds with those of other earth metals)	TiO, and some derivatives with analogous, compounds of	Vanadates with phosphates and	Salts of H.Cro, with those of H.Mno, and H.Teo.; Cr.o. with Al.O. Mn.O. and Fe.O.	Mn,O, with Al,O, Gr.O, and Fe,O, R,MnO, with B,GrO, and R,TeO, RMnO, with RGIO,	Fe,O, and derivatives with Al,O,, Cr,O, Mn,O, and derivatives. Some Fe salts with those of NI Co, and Cu
m.	Specific heat: how determined	Directly: sp. heat varies much with tempera- ture	Directly	r	Indirectly: donbiful from comparison of various baloid compounds]	Directly	=	Sp. heats of some com- pounds determined	Directly	Sp. heats of one or two	Directly: (? too low)	" (?too high)	Directly
TI .	Principal com- pounds vapour den- sities of which have been determined	SiP. SiCl. SiI., Si(CH.), SIH,CI, Si,OCI	Ph. Pol. Pil. PP. Pol. Psc. PP. Pol. Psc.	SH, SO, SO. SOCI.	CIR. CI (CE,) CIT. CI.Zh. CI.HK. CI.HC. CI.HK. CI.K. CI.K. CI.K. CI.K.	KI **, **.	None	R į	ਸਾਕ,	. VCI., VOCI.	cro <u>ci</u> , cra, cra,	MnCl,	• • • FeCls Falls
1 ·	Element	Bilicon	Phosphorus .	Sulphur	Chlorine	Potassium	Calcium	Scandium	Titanium	Vanadium	Chromium	Manganese .	Iron

ATOMIO AND MOLECULAR WEIGHTS.

		31	A'1') JE	TO Y	TIA.	D WOL	ECU	LAI	MEIGH	174 507
	·		9			- 80	26		12		P. C., 31, 391. W. Hampe, Pr. 13, 352. Baubigny, C.R. 17, 312. Gsy-Lusso and Thenari. G. 4. 37, 449. Barseline, P. R. 18, 184. Edmann, Berneine's Labbouch (5th ed.), 3, 113. P. A. Ferre, d. Ch. [3] 10, 103. Marigna, dr. 26, 13, 10, 5, 193. Leon de Budbaufran, C. Proc. 1894-87, 81. P. G. Winkley, Jr. 1865, 177. Proc. 1894-87, 81. P. A. G. Winkley, Jr. 1865, 177. Proc. 1894-87, 81. P. M. 1910, P. M. 1911, P. M. 1911, P. M. 1911, P. M. 1911, P. Staphovedder Rederate, 184, 110 and 112. Goodeffroy, A. 181, 339. Piccard, J. pr. 86, 453
ž	8	63.3		32.74	23-3	18-08	24.97	39.4	79.15	69 62 63 63 64 64 64 64 64 64 64 64 64 64 64 64 64	191. W. Hi. 184. Erdi. 3. A. Favre, 3. A. Favre, Winkler, W. Waller, W. W. Waller, W. W. Waller, F. F. Kessli F. F. Kessli Bansen, Bansen, Bansen,
** Nickel chloride; nickelous oxide; strychnine nickel cyanide; brucine-	* Amonius obait oyanie; phenii- amonium cohait cyanie; strychnine- and brucine cobait	24 Reduction of copper oxide; electrolysis of copper sulphate	** Synthesis of zinc oxide	Solution of zinc in	"Oxidation of the metal amenda	" Germanium tetra-	** Arsenic bromide; do. chloride, do. oxide	** Reduction of selenion dioxide; reduction	** Synthesis of silver bromide; potassium	s. Rubilium chloride	
989	2	63.3	6.19	65.43	6.69	72-35	0. .#	18.8	79.75	1	10. Berlin, Siev Myffen, 17. 8 lawsck, 17. 8 lawsck, 17. 8 lawsck, 17. 8 lawsck, 17. 18. 18. 18. 18. 18. 18. 18. 18. 18. 18
ı	1	63.2	6.19	63-48	6.69	79.32	74.3	78.8	21.62	7. 7.	[3] 12, 55, 17, P. 95, 17, P. 95, 15, P. 95, 15, P. 98, 15, P. 98, 15, P. 98, P. 149, P. H. Lee, B. H. Lee, B. H.
Ni with Co compounds: some Ni compounds with those of Fe (ferrous salts)	Oo with Ni compounds: some Co compounds with those of irou (ferrous salts)	Most Cu compounds with those of Ni and Co; some with Fe (ferrous) compounds. Cu and	Ag compounds or type had Zn compounds with those of Mg and Mn		Ga-slum with other alums	1	As compounds with those of Sh and Bi ; organic compounds of As with those of N, I', and Sb ; arsenates with phosphates and	Se with S compounds	Bromides, with analogous com- pounds of Cl and I	indirectly: doubtful Rb compounds with those of [from compounds with those of compounds with those of other alkali metals]	** Cr. E. Pilvot. 4. (A. [3] 19, 530. Berlin. 4. 5. * Cr. E. Pilvot. 4. (A. [3] 19, 530. Berlin. 4. 5. * Cr. E. 1.31. Fr. Kesider, P. 84, 210. Siewert. Edischrif für die gesummten Naturwieurschaften. 15, 530. S. Ball. Bawsch. P. 107, 53, 44. Berreins. 4. 50, 420. Svanberg and Norbin, 4. 50, 432. Erhenn. and Marchin. 4. 57, 122. E. Rivot. 4. 78, 214. Dumas. 4. Ch. [3] 55, 155. Swanberg and Norbin, 4. St. Nivot. 4. 78, 214. Dumas. 4. Ch. [3] 55, 155. Swanberg and Norbin, 4. 50. Swanberg. 4. Ch. [3] 55, 155. Swanberg. 4. Ch. [3] 55, 155. Swanberg. 4. Ch. [3] 55, 155. Swanberg. 5. 50. Weesley, 5. 8, 182. Erhmann and Marchand.
Directly		2	2		Directly [? tco low]	Directly	£ .	•	r	Indirectly: doubtful [from comparison of some compounds with these of other sikuli metals]	mas, A. Ch. [3] oung, C.J. Si, Sig. cittor, A. Ch. [3] 18, and Nouvelles , and Nouvelles 54, 210; 76, 219. Nerre, A. Ch. [3]
• • • Ni(CO),		ದಿ,ದ್	ZnCl, Zn(CH,),, Zn(C,H,),		Ga,೮1,, GaCI,, GaCI,	GeBr., GeCl., GeS	AεH,, AεCl, Aεl, As(CH,),Cl, Aε,0,, &c,	Sett., Sev., Sed., SecilBr	BrH. Br(CH.), Br,Cd. Br,B. Br,Sh,	RPCI. Rbi	 B.Si., Feloure, C. R. 20, 1047. Dums., A. Ch. [3] Sh.183. J. Sekiel, A. 120, 94. Thorpen. Young, C.J. 51, 575. B.S. 131. Dumas, A. Ch. [3] 55, 174. B.S. 132. Pumas, A. Ch. [3] 55, 174. C. Stas, Reports 53, 47, 44, 118, and Nowelless Redering 768. Redering, T. S. Stas, Rapports, 69, 91, 118, and Nowelless Redering 76. Redering, S. Rapports, 69, 91, 118, and Nowelless Redering and Marchad, A. 44, 216; 52, 210; 76, 219. Berlan, C. R. 17, 318. W. Stas, Silson, P. 4, 114, J. Flerre, A. Ch. [3] 58, 170. Barrela, C. R. 17, 318. W. B. S. Nikon, P. A. 11, 144, J. Flerre, A. Ch. [3] B. W. Thorpe, C. 44, 106. B. 407. Thorpe, C. 44, 106.
Hiokel	Cobalt	Copper	Zino		Gallium	Germanium .	Arsenic	Selenion	Bromine	Rubidium	18 B. Pelouze, 18 J. Pelouze, 18 J. Schiel, J. Pelouze, 18 J. Stav. Krg. C. Stav. Krg. C. Stav. Krg. Recherch, Stav. Recherch, Stav. Recherch, Stav. Recherch, Stav. Bereitt Erfman and Masserska, C. R. If Serveta, C. R. If Servet

ATOMIC AND MOLECULAR WEIGHTS.

Mement Po		1	١٧.		1	AT.	1	AUI.
-	Principal com- pounds wapour den- stries of which have	Specific heat: how determined	Isomorphism: compounds compared [see Note A, p. 361]	Atomic (1)	١	Compounds analysed &c. in order to find com- bining weight of the	Com- bining weight	Remarks [see Note C, p. 261]
	Deen determined			density	method	[see Note B, p. 361]	[Note B, p. 361]	
Strontium	None	Indirect; s doubtful (c mparison of compounds of Sr, Ca, and Ba)	Sr compounds with those of Ca and Ba, and with some Ph salts	1	1	2 Strontium chloride	43.65	Sr. The atomic weight of stron- tinm must be taken as 43.65 x 3 m 87.3 if the formules of the salts are to become analogous to those
Yttrium	•	Sp. heats of a few com- pounds determined	Yt compounds with those of other earth metals	ı	1	Synthesis of yttrium sulphate	29-87	of the Ba and Ca salts 17. Atomic weight probably = 29-87 × 3 = 89-6 because of analogy of Yt salts with those of
Zirconium Za	zrO1.	Directly [? toe low]	ZrO, with TiO, ThO, SnO, and SiO,	2	2	** Zirconium chloride; potassium - zirco-	\$	the earth metals
Miobium N	NbCl., NbOCl.	ı	Nh with Ta compounds. Nb fluorides and oxyfluorides with	46	1	num inoride * Niojum chloride; potassium niohium	31.33	
Molybdenum M	MoCI.	Directly [? too high]	Mo do, do, Mo with W compounds; some salts of H,MoO, with those of H,CrO, Mo with Nh fluorides	95-8	8.58	oxyfluoride Molybdennm dichlo- ride, tetrachloride, and pentachloride	19.16	
Buthenium . Bn0.	tho.	Directly	Most Rn compounds with those	\$.T01	101.4	* Potassinm-rutheninm	25-38	
Rhodium	None		Most Rh compounds with those	1	102:7	Potassium - rhodinm	25-55	
Palladium	ř	•	Most Pd compounds with those	1	106:3	Palladium chloride	26.58	
Silver A	Aga		Some Ag compounds with those of Na and other aikali metals.	107-66	107-68	Silver chlorate, bro- mate iodate:synthe-	107-66	
			Ag with Cu compounds of type R.O. A few Ag and Au			sis of silver bromide and iodide		
Cadmium CdBr,	dBr,	•	Some Cd compounds with those	112	112	" Cadmium bromide	99	
Indium In	Incl., Incl., Incl	•	Some In compounds with those	113.4	113.4	Synthesis of indium	37.8	
Tin Sn	SnCl., Sn.Cl., SnCl.	•	Sno. with Tio, Zro, and Tho.	117-8	8./11	Synthesis of staunic	58.5	
Antimony . Sp	SbCl., SbBr., Sbl., Sb(CH.), Sb.0. SbCl.	A	Sh compounds with those of As	120	120	Antimory bromide: reduction of anti- mony oxide; also analysis of antimory sulphide	đ	

			ATOM	IO	AND	M	OLEC	ULAR	WEI	GHTS.	
			Ba. Atomic weight is 68.4 × 2=136.8, because of analogies between sales of the Sr and Ca				Er. This metal belongs to the carth group, hence the atomic	WEIGHT IS MAKEN BEND A X-A ELDO. 10. For similar reasons to those which apply in case of So, Y, and Er, the atomic weight of Ytter. bium is supposed to be 3 times the	combining weight = 118		** Ba. Marignae, A. 68, 218. Dumas, A. Ch. [3] 55, 137. [2] La. Marignae, A. Ch. [13], 67. Chev. Bl. 60, 212; [2] La. Marignae, A. 65, 214, 13, 67. Chev. Bl. 60, 212; *** Ch. Marignae, A. 65, 218. H. Bunrig, J. pr. [2] 13, 250. *** Robinson, P. F. 37, 130. Branner, C. J. Tran. 1884, 879. *** Di. R. Branner, C. J. Tran. 1882, 68, 18. *** Di. P. T. Cheva, C. R. 21, 331. Nilson, B. 13, 1446, 179. *** To. Marignae, A. Sappl. [4] 331. *** W. Roscoe, C. K. 54, 61, 73.
:	126.53	132.7	£-89	46-17	9-94	\$	(47·6)) 55·3	67-63	19-09	30.6	Marignac, Marignac, Brauner, Marignac, 1300, Pr. 1300, Pr. 18 P. T. Clev P. T. Clev Nilson, B. Marignac, Roscoe, C.
** Oxidation of teiln- riuo: potassium- teilurum bromile; synthesis a teiluriom sulplace teiluriom sulplace	** Silver iodate: silver iodile: synthesis of do.	" Cæsium chloride	* Barium chioride	"Lanthanum sulphate;	late Ccrium oxalate; do. chioride; do. sui-	* Dilymiam oxide and	** Erhium sulphate	* Ttterbium snlphate	** Potassium - tantalum fluoride : amno- nium-tantalum flu-	** Reduction of tungstic oxide; tungsten berachloride	
125	126-53	I.	ı	138-5	133-9	144	(342)	ı	1	153.6	11, 28; C. N. 292. Buosen, 292. des Antimon. 13, 1; 11, 13, 704. Brauner, 2, 189, 193. 3, Allen, Ass. S.
125	126-53	135.1	I	ı	ı	1	1	1	162	183.6	P. 4m. 4. 14, 8; 102. 154. s 4(onjectral) P. 4m. 4 runs. 1879. Johnson 185.
Most Te compounds with those of S and Se	lodides with analogous com- pounds of Cl and Br	Cs componeds with those of other metals of aikalis	pounds with those of obtained with those of contractly and obtained of contractly. Ba compounds with those of Ca [Comparison of cone. and Sr		Most La compounds with those of Ce. Di. Er, and Yt.	Some compounds of these metals with Ca compounds		? A few Tb compounds with those of other earth metals]	Ta with Nb compounds	W with Mo compounds. Some salts of H, WO, with those of H, CrO, and H, TeO.	** Cd. O. W. Huntingtoo, P. Am. A. 17, 28; C. N. 41, 20. *** Th. C. Winck'er, J. pr. 94, 8; 102, 282. Buosen, P. 181, 2. *** Sh. S. Barbeller, Cife and Journal of Antimors (Berlin), 1860, J. P. Cook'e, P. Am. A. 13, 11, 13, J. Bonartz, B. 18, 199, 20. *** The Walls of Cook'er, J. Cook'er, J. A. 13, 11, 17, 13, J. Bonartz, B. 18, 199, 198, C. J. Trunt, 1879, 704. Branney, B. 18, 5005, Bursen, P. A. 119, 1, Johnson E. M. Barsen, P. A. 119, 1, Johnson E. Allen, Am. S. [2], 35, 94, R. Godeffoy, A. 184, 185.
Directly	£		pounds with those of other alkali metals] Io-lirectly : doubtful [comparison of com-	Directly			Sp. heats of a few com- pounds determined	Ř	ı	Directly	as. A. Ch. (3) 185, 191, Stringer, B. 9, 1540, Entgan, A. 1540, 189, 344, R. 107, 394, 189, 946, ert a. Rodbé, A. 250, ett a. A. A. 250, 44; and Nourelles
Tellurium TeH., TeCl., TeCl.	IH, ICI, I(CH ₃), I ₃ P., I ₃ Hg, I ₃ P. I ₃ As, I ₄ Si, I ₄ Al ₂ .	3; 50 50 50 50 50 50 50 50 50 50 50 50 50	None			R	á.	a	TaOL.	WOCI, WCI, WCI.	Marignac, A. 106, 168. Dumas, A. Ch. [3] 55, 191, Cleve, B. 6, 145; Itanmesister, E. 9, 156; Hermann, J. Pr. 31, 77. Marignac, A. Ch. [3] Marignac, Pr. 5, 480. P. Lieschit, B. Krempe, A. 169, 384, Claus, P. 65, 138, Joly, C. R. 107, 984, 108, 945, Berrelins, P. 13, 437. Seubert a, Kobbe, A. 200, Berrelins, P. 13, 437. Seubert a, Kobbe, A. 200, Sian, Rapsort, 38, 43, 44; and Nourelization, P. 18, 184, 184, 189, 184, 806
Tellurium	Iodine	Cæsium	Barium	Lanthanum .	Cerium	Didymium .	Erbium	Ttterbium .	Tantalum	Tungsten	Sr. Marigna S. Y. Cleve. B S. Zr. Herman S. Nr. Marigna S. Nr. Marigna S. Mo. P. Licel F. Hu. P. Licel J. H. Berzelin S. R. Berzelin S. Sas. B. Ber

	ATT.	Remarks [See Nove C, p. 361]																		** Bi. Schneider, P. 82, 303, Dumas, A. On. [3] 58, 175; Marginas, A. Ph. [3] 13, 15, 5, 138. Edwe, Pr. 22, 488, Schneider, Apr. [7] 39, 3710, 5, 138. Thy Nitson, P. 15, 327. Krits a. Nitson, B. 50, 1883, S. Th.
	VII.		p. 361]	47.24	48.125	48.575		65.08		6 .66		203.64		103.2		.e.69		57-97	09	thneider, f. 19. [2] 30, iison, f. 15. [2]
	VI.	Ü	[see note B, p. 361]	Se Osmium tetroride	Potassium . inidium	Potassium - platinum	and original	5. Gualite, &c.	the and bromide	do. oxide	4. Strithogic of the	nitrate		Synthesis of lead ni-	is Courthood.	Intlinits ox 10 515	muthous chioride	Thorner Pu.phate	" Craniunacetate; do.	
	V. Atomic weight	By varour By sp. hent density method		150.3	192.5	2.561		197	103.6		- 603-61			F-902	808		23:-87	5	0.50	Tpe a. Lanrie, 1. 392. Syan.
	Atomic	By varous density	mentod	190-3	I	1		1	139.8	2	203.64	!		* 90:	808		231-57		042), 355. Tho d. <i>J. pr.</i> 31 . [3] 18, 344
	E:	Isomorphism: compounds compared [see Note A, p. 361]			Os. Ir. and Pt compounds	Pd		Some Au compounds with those of Ag. A few An compounds	with those of Ni and Fe	ko	Tl compounds with those of	type RG: Tl compounds of type TiG with those of alkeli	Some Ph with m	many Pb with Cu and Hg com-	Li compounds with those of	As and Sh	Tho, with Sio, Tio, Sno, and	Some If compounds as trees	C.O. with those of Al, Cr, Mn,	4. C., 15, 337. Levol, A. Ch. [3] 30, 355. Thorpe a. Laurie, C., 5, 15, 555. Kritss, R. S., 203. W. H.G. Erlemann a. Murchand, J. pr. 31, 332. Svan. W. A. Joseph A. Star, A. Ch. [3] 18, 345. W. C. Croftes, T. 19, 35, 577.
	LIL	determined	Directly	f 100 a					•		R		•	ł	R)3 W.11a:ber- 1212. Javal,
F	Principal com-	sities of which have been determined	0.0	None		_		•	[HgCl], HgCl	(C.C. 18)			Poct, Pb(CH,),		BiC', BiF,		T.BCI.	· · TCI., UBr.		** Os. Devil'e a. Dodorry, 4. (74, [3] 56, 405 ** Ir. K. Sculert, B. II, 174, 7. ** Ir. K. Sculert, B. Ist, 865; 4. 207, 20. W. Haiber- ** Ir. Berrelius, Lehrbuch (5th ed.), 3, 1212. Javal.
-	i	Tiente in	٠.	•	Platinum		Gold		Mercury	Thallium	•		Lead Pbcn, Pb(CH,),	, in the second	· · unmard	Thomas		Oranium 1		*Os. Devil'e a. I. * Ir. K. Seulert, * Pt. K. Seubert, * And J. 17, 2962.

Notes to Table of Atomic Weights.

A. As the method based on isomorphism of compounds is chiefly used as a means of verifying values assigned to atomic weights by other methods, no numbers are given in column IV., but merely an indication of the various compounds which have been compared crystallographically, and on which arguments for or against a given value for the atomic weights in column V. have been, or may be, based.

B. This column (VI.) is not to be regarded as containing anything like a complete summary of the processes employed for determining the combining weights of the elements; only the more important processes are indicated references are given to the original papers.

By combining weight is here meant the smallest mass of the element which combines with 8 parts by weight of oxygen, I part of hydrogen, or 35.5 parts of ohlorine.

A complete account of all researches on this subject will be found in A Recalculation of the Atomic Weights, by F. W. Clarke [Part v. of the Constants of Nature published by the Smithsonian Institution], and also in Die Atomgewichte der Elemente, by L. Meyer and K. Scubert

[Leipzig, 1883].

C. When the atomic weight given in column V. section (2) is a multiple of the combining weight in column VII., no number being given in section (1) of column V., it is to be inferred that, besides the argument drawn from the value of the specific heat of the element in question, there are other chemical reasons for adopting the special multiple which appears in V. (2); these reasons may be broadly described as based on analogies between salts of the given element and salts of other elements the atomic weights of which have been established by the two leading physical methods. M. M. P. M.

ATOMICITY. Term used to denote number of atoms in any specified gaseous molecule, usually in the molecule of an element.

ATOMIC VOLUMES v. PRYSICAL METHODS; sect. VOLUMETRICAL.

ATRACTYLIC ACID C ... H ... S .O ... Potassium atractylate K.A" occurs in the root of Atractylis gummifera, from which it may be extracted by boiling 70 p.e. alcohol (Lefranc, Bl. [2] 11, 499; J. Ph. [4] 9, 81; 10, 325; 17, 187, 263; C. R. 67, 954; 76, 438). Boiling potash hydrolyses it, forming valerio acid and so-called (\$)-atractylic aoid, C20 H38 2O16, which is further split up into H2SO4, valerio acid and atractylin.

ATRACTYLIN C20 H30O6. From '(B)-atractylic acid' by boiling with aqueous KOII. White gummy substance, with sweet taste, v. sol. water and alcohol, insol. ether and aqueous NaCl. Forms a violet-red solution in warm H₂SO₄. Boiling KOH forms atractyligenin and a saccharine substance.

ATRANORIC ACID C, H, O. [190°-194°]. Extracted by ether from certain lichens Lecanora atra, Stereocaulon vesuvianium, Cladonia rangiformis). Trimetric crystals; a:b:c=1:398:306; sl. sol. alcohol, cold ether, and benzene; m. sol. hot benzene; sol. alkalis forming a yellow solution. Heated with water in a sealed tube it splits up into atranorinic acid, C.H.O., and atrarie acid, C. H.O. (Paterro, G. 9, 279; 10, 157; 12, 256; Coppola, G. 12, 19).

ATRANORINIC ACID C,H,O,. Formed by heating atranoric acid (q. v.) with water. Needles, m. sol. water, sol. alcohol and ether. Its alkaline solutions are yellow. Its aqueous solutions give a green pp. with AgNO, a brownish-green colour with Fc.Cl., and blood-red colour with bleaching-powder (Paterno, G. 12, 256).

ATRARIC ACID C10 H10O2. [141°]. Produced by heating atranoric acid with water (Paternò, G. 12, 257). Iridescent laminæ, sl. sol. water, m. sol. alcohol and ether. Its alkalino solutions are colourless. It gives a brownish pp. with AgNO, and no colour with Fe Cla.

ATRIPAÏC ACID CallaO12 Gad. [98° when hydrated). An acid obtained from the sugar eanc (Savary, C. C. 1884, 968).

Ethyl ether (184°-188°).

ATROGLYCERIC ACID C,II,O, i.e.

CII. (OH). CPl1 (OII). CO. II. [146°]. a\$-Di-oxy-a-phenyl-propionic acid. From a\$-di-bromo-aphenyl-propionic acid and oxcess of alkali (Kast. A. 206, 30). Crystalline aggregates, sol. water and ether. Salts.—CaA',... BaA',.. Nitrile CH.OH.CPh(OH).CN. [57°]. From

benzoyl-carbinol and HCN (Plöchl a. Blümlein, B. 16, 1292).

ATROLACTIC ACID v. a-Oxy-a-PHENYL-PRO-PIONIC ACID.

ATROLACTYL-TROPEINE C.H., NOC, H.O., Pseudo-atropine. [121°]. Crystalline solid. Very similar in physiological action to atropine. Formed by the action of dilute HCl on tropine atrolactate.

Salts. - Mostly soluble. - B'HClAuCl3: sparingly soluble tables. The picrate also forms sparingly soluble tables (Ladenburg a. Roth, B. 15, 1027; A. 217, 87).

ATRONENE C₁₆H₁₄ i.e. C₀H₄ CH₂ - CH CPhH.CH

Phenyl-naphthalene dihydride. (325° i.V.) Formed, together with atronio acid, by the dry distillation of (a) - or (B) - iso-atropic acid (Fittig, A. 206, 51). Liquid. Chromic acid oxidises it to o benzoyl-benzoio acid.

Atrenene sulphenic acid C₁₆H₁₈SO₈H. [130°]. Needles; v. sol. water.—BaA'₂.—CaA'₃2aq.
ATRONIC ACID C₁₂H₁₄O₂ i.e.

Coll, Colling CH. (?) Phenyl naphthalens di-hydride carboxylic acid (i). [164°]. Formed, together with atronene by the dry distillation of (a)- or (B)-iso-at: pic acid (Fittig, A. 206,

4(i). Prisms; insol. water, sol. alcohol and glacial HOAc. Salta.—CaA'.6aq.—BaA'.4aq. Iso-atronic acid C.,H₁,O₂. [157]. Obtained by heating (a)., or (β)., iso-atropic acid with conc. H₂SO₄ (Fittig, A. 206, 86). Leaflets, insol. water, sol. alcohol, ether, and glacial HOAc.

Salts.—CaA', BaA', Gaq.
ATRONYLENE SULPHONIC ACID

C16H11.SO3II. [c. 258°]. Formed by heating (a)-, or (β)-, iso atropic acid or iso-atronic acid with 9 pts. conc. H₂SO₄ at 90° (Fittig, A. 206, 60). Prisms (from 50 p.c. acetic acid). Insol. water, v. sol. alcohol. The aqueous solutions of its salts when exposed to sunlight deposit small

needles of atronyl-sulphone $C_{is}H_{jo}SO_{2}$. [193°].

ATROPIC ACID C₈H₈O₂ i.e. CH₂:CPh CO₂H.

a-Phenyl-acrylic acid. M. w. 148. [107°].

(203°) at 75 mm. S. 111 at 19°.

Fornation.—1. By boiling atropine with baryta (Bichter, J. pr. 11, 33; Kraut, A. 128, 282; Fittig a. Wurster, A. 195, 147).—2. By heating atropine with funning HCl at 120° (Lossen, A. 138, 230).
3. By the action of HCl on ethyl-tropic acid, CH_xCPh(OEt).CO.II, obtained from acetophenone chloride by alcoholic KCN and saponification (Ladenburg a. Rügheimer, B. 13, 2041).—4. By action of boiling NaOH upon α-chloro-β-phenyl-propionic acid which is formed by the action of HCl upon acetophenone cyanhydrin or by heating α-οxy-α-phenyl-propionic acid with saturated HClAq at 130′ (Spiegel, B. 14, 237; 1354).

Properties.—Needles (from water) or monoclinic prisms (from alcohol); v. sol. CS₂.

Reactions.—I. Chromic acid mixture forms benzoic acid.—2. Potash fusion forms phenylacetic acid.—3. Sodium-analyam reduces it to a-phenyl-propionic acid.—4. Furning HCl forms a-chloro-a-phenyl-propionic acid, which is converted by aqueous Na.CO, at 120° into tropio acid, CH_(OII).CPhH.CO.H.—5. ClOH forms chloro-tropic acid CH_(OII).CPhCl.CO.H (Ludenburg a. Rügheimer, B. 13, 376).—6. Cold conc. Il BrAq forms both a., and B., bromo-a-phenyl-propionic acid; at 100° it forms only B-bromo-a-phenyl-propionic acid.—7. Bromine forms CH_Br.CPhBr.CO.H.

Salts.—Neutral atropates are not ppd. by manganous salts (difference from ciunamates).—CaA'₂5aq (K.).—CaA'₂2aq (L.).

(a)-Iso-atropic acid C18H16O1 (?). [2370].

Preparation.—(a), and (8), isoatropic acids are both formed when atropic acid is heated alone or with water; they may be separated by orystallisation from 50 p.c. acetic acid.

Properties.—Crystalline aggregates, sl. sol. boiling water, sol. alcohol. Chromic acid gives anthraquinono and o-benzoyl-benzoic acid. V. also Atronene, Atronic acid, and Atronylene sulfing acid.

Salts.--CaA" 2aq.--BaA" 21aq. Ethyl ether Et.A" (180°).

(B) Iso-atropic acid $C_{is}H_{ie}O_{i}$. [206°]. Rectangular tables (from water). More soluble in water, alcohol, and HOAc than the (a) acid. Gives the same reactions as the (a) acid.

Salts.—CaA" 3aq (Fittig, A. 206, 34; B. 12, 1739; compare R. Meyer, A. 219, 290).

ATROPINE C, H23NO, i.e.

CH_OH_CPhH_CO.O.CH_CH_CH_CH_NMe. Daturine. [114°]. S. 33.

Occurrence.—Together with hyoscyamine in all parts of Atropa belladonna (Geiger a. Hesse, A. 5, 43; 6, 44; Mein, A. 6, 67), and in the seeds of Datura stramonium (Geyger, A. 7, 272; Planta, A. 74, 245).

Formation.—Crystalline tropine tropate has no action on the cyes, but when treated with dellydrating agents, such as ZnCl., Ao.O., or HCl, atropine is formed. It is best to evaporate frequently with very dilute HCl at 100° (Ladenburg, A. 217, 78; B. 12, 912; 13, 104,

909; C. R. 90, 921). Tropine contains hydroxyl which is etherified by tropic acid:

CH_a(OH).CPhH.CO₂H+HO.CH₂.CH₄.C₂H,NMe=CH₄(OH).CPhH.OO.O.CH₄.CH₂.C₄H,NMe+H

Preparation.—Dry belladonna leaves are digested for three days with cold water, the extract is evaporated, and after uixing with Na₂CO₂ the syrupy liquid is agitated with benzene. The benzene solution is decanted off and shaken with dilute sulphuric acid. The acid liquid is rendered alkaline with Na₂CO₂ and the solution agitated with chloroform; the extract is filtered and, after addition of light petroleum, allowed to evaporate spontaneously, when the atropine separates out first, the mother liquors containing another alkaloid (Pesci, G. 10, 426).

Properties.—Needlos (from dilute alcohol). Sl. sol. water; v. sol. alcohol, and chloroform, m. sol. ether. The solutions are alkaline to test-paper, and taste bitter. Its salts enlarge the pupil of the eye. '05 to 2g. is a fatal dose. Three drops of a 1 p.c. solution of (artificial) atropine enlarges the pupil to the maximum extent. Atropine overcomes the stoppage of the heart's action produced by musearine.

Reactions. - I. When evaporated to dryness with fuming IINO, a residue is left which is turned violet by alcoholic KOII.—2. Chromic acid mixture forms benzoic soid.-3. A solution in HCl gives with gotd chloride an oily pp. that quickly changes to lustreless crystals which melt under water or, when dry, at 136° .-4. Tannin gives, in very dilute neutral solutions a white pp., sol. HCl .- 5. Potassio-mercuric iodide gives a white cheesy pp.-6. I in KI gives a brown oil which solidities after some time .- 7. Picric acid gives, in somewhat dilute acid solutions, a crystalline pp. -8. Conc. H.SO. gives, on warming, a pleasant odour .- 9. Cyanogen gas passed into an alcoholic solution gives, after some time, a red colour. -10. Chtoride of iodine forms a dark yellow pp., sol. on warming, and separating out on cooling in brown crystals (Dittmar, B. 18, 1612).-11. Dc. composed by hot baryta-water or cold cono. HCl into tropic acid and tropine (Kraut, A. 128, 280; 133, 87; 148, 210; Lossen, A. 138, 230).—12. Hot cone. HClAq at 120° gives tropine, tropicacid, atropic acid, iso-atropic acids, and (at 180°) tropidine .- 13. With NaNO, II,SO, and subsequently NaOll a violet colour is developed .- 14. Glacial HOAc and H.SO. produce on prolonged warming a greenish-yellow fluorescence (Flückiger, Ph. [3] 16, 800).-15. H.SO, and KClO, give a greenish-blue colour (Vitali, Ph. [3] 12, 459).

Salts.—B'HAuCl, [135°—137°].—B'_HPtCl_ [208°].—B'IICl 2HgCl_2 (Gerrard, Fr. 24, 601). B'HIL, brown prisms (Jörgensen, Z. 5, 673).—B'HIL,—B',H_SO_4: needles, got by adding an othereal solution to an alcoholio solution of H_SO_4 Valerate B'C₂H₁₀O₂ \(\frac{1}{2}\text{aq} \) [42°] (Callmann, T_2 , T_3 , T_4 , T_5 , 69).

Additionat References.—Günther, J. 1869, 781; Fr. 8, 476; Lefort, Ph. [3] 2, 1029; C. C. 1873, 797; Brunner, B. 6, 96; Newark, C. C. 1872, 536; Guliolmo, Fr. 2, 404; Ludwig, Ar. Ph. [2] 107, 129; Schmidt, A. 208, 196.

Ethyl-atropine C,H,EthO, Formed by action of Ag,O on its hydriodide, B'HI, obtained

by heating atropine with Etl at 100° (Lossen,

A. 138, 239). A syrup, sol. water.

Apoatropine C, H₁₁NO₂. [60°-62°]. Formed by treating atropine with HNO₃ (Pesci, G. 11, 538; 12, 60). Prisme; sl. sol. water, v. e. sol. alcohol. Decomposed by baryta-water at 100° into tropine and atropic acid. Does not enlargo the pupil. Salts .- B'HAuCl, [180°] amorphous .-- B'H.SO, 5aq.

Pssudo-atropine v. ATROLACTYL-TROPEINE. Hydro-apc-atropins

CH, CHPh.CO.O.CH, CH, C, H, NMo. by the action of nascent hydrogen on apo-atropine (Pesci, Atta d. Acad. dei Lincei, 5, 329). Oil. Decomposed by baryta-water at 100° into a-phenyl-propionic acid and tropine. a crystallino compound with HgCl... Neutral KMnO, oxidises it to 'homo apo atropino' C16H21NO2, an alkalino oil which forms the following salts: B'CO., B'.H.PtCl., B'HAuCl., B'HAuCl., B'.H.SO., xaq, B'.H.PtCl.; its hydrochloride gives white pps. with tannin, Mayer's reagent, and HgCl₂; and it gives a blood-red colour with fuming HNO₂; heated with baryta-water it gives a-phenyl-propionio acid and tropigenine (Pesoi, G. 12, 285, 329; Merling, B. 15, 289).

Homo atropino v. l'henylglycolyl-tropeïne.

Meta-atropine v. TROPINE.

ATROPYL-TROPEINE C, II, INO. Anhydroatropine. Obtained by heating tropino hydrochloride with atropic acid and HCl (Ladenburg, A. 217, 102; B. 13, 1085). Oil.—B'HAuCl.; small needles

ATROXINDOLE v. o.AMIDO. g.PHENYL-PROPI-

ONIO ACID, p. 179.

AURANTIIN C. 112, O12 4aq. [171°]. S. 33. $[a]_1 = -64.57^{\circ}$. A glueoside in the flowers of Citrus decumana (E. Hoffmann, B. 9, 691). Yellow monoelinic prisms. Bitter taste. Gives a brownish-red colour with Fe₂Cl₆.

AURATES. Auric hydroxide (=Au2O3.H2O) reacts with HNO3Aq to form the compound Au(NO.) .. HNO. 3H.O; from this several basic nitrates and one or two sulphates of gold may be produced (Schottländer, A. 217, 312). But Au,O, H,O also dissolves in KOHAq, and on evaporation at a gentle heat and finally in vacuo, crystals are obtained, which when dissolved in water, recrystallised, drained, and dried in vacuo, are eaid to have the composition Au,O, K.O.311.0 (=Au₂O₄K₂3II₂O). Aurate of potassium is very soluble in water, forming an alkaline liquid which is easily decomposed by organic com-pounds and by heat; the solution gives pps. with solutions of various metallic salts, e.g. CuCl.; these pps. are said to be anrates, but very little is known about their composition. An aurate of ammonium of indefinite composition, known as fulminating gold, is formed by pptg. a solution of a gold salt by excess of NII3Aq, and boiling in NH,Aq; or by digesting Au,O,H,O in (NH),SO,Aq (Premy, A. Ch. [3], 31, 480; Figurer, id. [3] II, 34I). M. M. P. M. AURIC ACID. The hydrated oxide of gold

An,O,H,O is sometimes called auric acid because of its salt-forming reactions with alkalis (v.supra). This compound is best prepared by adding NaOHAq to very dilute AuCl₃Aq, in the ratio NaOH:AuCl, (the AuCl, Aq should be prepared in about the ratio AuCl.:800H2O); warming till the liquid is dark brown; adding Na2SO,Aq;

allowing pp. to settle; washing repeatedly by decantation, and then on a filter, until the washinge are free from H₂SO, and HCl; boiling pp. with conc. HNO, Aq; and again washing free from acid (Thousen, Th. 3, 391). According to Krüss (B. 19, 2546), Au.O. II.O is better prepared from Au.Cl.Aq by ppg. with magnesia alba, boiling with dilute HNO.Aq, washing with water, and drying over P₂O₃. Au₄O₃.H.O is easily soluble in HBrAq and HClAq, with production of much heat, and formation of AnCl, HAq and AuBr, HAq respectively (v. Gold). M. M. P. M. AURINE v. Rosolic Acid.

AUSTRALENE v. TURPENTINE OIL.

AXIN v. Age, p. 87.
AZAMMONIUM COMPOUNDS. Compounds obtained by oxidising mono-alkylated o-amidoazo compounds, or by heating azimido-compounds with alkyl iodides followed by moist Ag.O (Zincke a. Lawson, B. 20, 1173). Thus $C_a\Pi_3N!N.C_{1a}H_a.NII.C_b\Pi_i$ gives, on oxidation with chromic acid, $C_{a_i}\Pi_{i_1}N_{i_2}O$ which might be expected to have the formula:

 $C_{i_0}\Pi_* \overset{N(OII).C_v\Pi_*}{\underset{N \longrightarrow N.C_oII_*}{|}}.$ Naphthalene · di · phcnyl · azammonium hy-

 $C_{zz}H_{16}N_{z}$.OII i.e. $C_{10}H_{a} < N(OH) \cdot C_{d}H_{z}$ (?).

The chromate is obtained by the exidation of benzene · azo · phenyl · (β) · naphthylamino

C₁₀H₀ | (?) with K₂Cr₂O₇ and acctic acid. It is converted into the chloride by boil-

ing with alcohol and HCl till all the chromic acid is reduced. The hydrate is obtained from the chlorido by Ag.O. Its solution has a greenish fluorescence, is strongly alkaline, and tastes bitter; it decomposes on evaporation.

Salts.-Like the base, they have a greenish fluorescence in aqueous and alcoholic solution, and a bitter taste - C22H16N3Cl: glistening prisms, v. sol. alcohol, less in water; it forme sparingly soluble double chlorides with SnCl2, ZnCl₂, IIgCl₂, &c.—(C₂ Π_{16} N₃Cl)₂PtCl₄: sl. eol. erystallino pp. — $C_{22}\Pi_{16}$ N₃,IISO₃: glistening transparent needles or prisms, v. sol. hot alco hol, sl. sol. cold water. — C., II, N, NO, : long tlat glistening needles, v. sol. hot alcohol, sl. sol. water. — $(C_{\omega}H_{16}N_3)_{\omega}Cr_{\omega}O_{\tau}$; long yellow needles, sol. acetic acid, v. sl. sol. water.— $\times (C_{\omega}H_{16}N_3)C_{\omega}H_{\omega}(NO_2)_3O$; [243°]; small yellow needles, v. sl. sol. water (Zincko a. Lawson, B. 20, 1172).

AZARONE C₁₂H₁₄O₂, [59°], (296°), S.G. 13 1:165; and 1:0743. Obtained by distilling the rhizomes of Azarum europæum with steam (Boutleroff a. Rizza, Bl. [2] 43, 114). White crystalline body, sl. sol. water, sol. alcohol, etber CCl, and acetic acid. Combines with Br, forming C₁H₁₀Br₂O₂.

AZAUROLIC ACIDS.

Ethyl-azaurolio acid C.H.N.O. [142°]. From ethyl nitrolic acid (2 g.) by the action of water and sedium amalgam. The yield (25 g) is bad. Formed also by reducing di-nitro-ethane (V. Meyer a. Constam, A. 214, 330; B. 14, 1455). Properties .- Orango-red prisms (from aloo-

hol). M. sol, hot alcohol, sl. sol, ether, v. sl. sol, water, chloroform, benzone or light petroleum. Alkalis form a deep orange solution. At 142° it melts, with decomposition, leaving a liquid which, after solidifying, melts again at 133°.

Reactions .- 1. Ammoniacal solution gives with AgN(), a brown pp., and with salts of Zn or Pb, yellow pps. A solution of the ammonium salt deposits, on evaporation, needles of the free acid. 2. Heat, dilute acids, nascent hydrogen, and ammonia all convert it into ethyl-leucazono: 2C₂H₄N₂O + H₂O = C₄H₄N₂O + O + NH₂O the oxygen converting another portion of ethyl-azaurolio acid into acctic acid, N, and N.O.—3. K.Cr.O. and Il, SO, give acctic acid and CO.

Constitution. - Its formation from ethylnitrolio acid indicates the group McC<N, and the presence of Me.C is shown by the production of acetic acid on oxidation. Ethyl-azaurolic acid would then be Mo.CII NO or, more probably, McC11(NO).N:N.C11(NO)Me or perhaps McC(NO): N.NH.CMc: NOH.

Ethyl-leucazone C,11,N,O. [158°]. Formed together with nitrogen, N.O, and hydroxylamino by heating othyl-azaurolic acid with diluto HCl (M. a. C.). Satiny needles. Reddens litmus. Combines with acids, bases, and salts. V. sol. alcohol or water, insol. ether. Its aqueous solusteon or water, inside each or the state of the steon is turned red by Fe₂Cl₃ and oxidised to acetic acid by K.Cr.O. and H.SO.

Salts.—B'.H.SO.; [161-5°]; prisms, sol.

ordinary [90 per cent.] alcohol.—Ba(C,H₂N₃O).

—C.H.N.OAgNO.;

Propyl-azaurelie acid CaH, No. [127.5°]. From propyl-nitrolic acid by sodium-amalgam. AZELAIC ACIDS C,H,O.

n-Azelaïo acid

CO.H.CII..CH..CH..CII..CII..CH..CII..CO.H. [118°]. Formed by reducing butyro-furome acid with H1 and P (Tönnies, B. 12, 1200). Slondor needles (from eliloroform).

Azelaïo acid C₄II₁₆O₄. Anchoic acid. Lepargylic acid. Mol. w. 188. [105°]. (above 860°). S. 108 at 12°; S. (ether) 1.88 at 11°.

Formation .- 1. By the oxidising action of HNO, npon Chinese wax (Buckton, C. J. 10, 166), eork, oleïe acid (Laurent, A. Ch. [2] 66, 154), coeoa-nut oil (Wirz, A. 101, 265), castor oil (Arppe, A. 120, 288; Ganttner a. 11ell, B. 14, 560, 1545), and enniuoic acid (Krafft, B. 11, 1415).—2. From oleie and KMnO, Aq (Saytzeff,

J. pr. [2] 33, 301). Preparation. Custor oil is oxidised by HNO. (S.G. 1.25). 11cptoic soid is distilled off with steam, and the hot residual liquor poured off from a heavy nitrogenous oil. On ecsuberic and azelaic acids crystallise. On cooling, suberic acid is got by washing the mixed acids with ether, which dissolves azolaic acid as well as oily impurities. The other is evaporated, the residuo dissolved in boiling water and NaCl added. Oily matter then separates and azolaïo acid erystallises from the brine (Dale a. Schorlemmer, C. J. 35, 684; cf. Ganttner a. Hell, B. 14, 1545).

Properties.-Large thin plates, not volatile with steam. Sol. water, alcohol, and ether. Nitrie acid oxidises it to succinic seid. It does not give a homologue of suberone when distilled

with slaked lime, hence its constitution is probably not analogous to that of the homologous suberic acid.

Salts.—K₂A": small plates.—K₃A"2aq: needles.—K₁A".—Na,A"aq: soluble plates.—Na₂HA"₂.—(NII₄)₄A": large plates.—(NH₄)IIA".—BaA"aq: S. 65 at 16°; 628 at 100°.—SrA"aq.—CaA": crystalline powder; S. 100°.—StA" aq.—CaA": crystalline powder; S.
185 at i7°; 193 at 100°.—MgA" 3aq.—MnA" 3aq: sleuder needles; S. 206 at 14°; 103 at 100°.

— NiA" 6aq.—CoA" 6aq.—ZnA": crystalline pp.; S. 226 at 12°.—PbA": white pp.; S. 206 at 14°.—CuA".—Te(OII)A" 2aq.—CdA".

Ethyl ether Et,A". (260°). Decomposed on belling

AZIDINES. Compounds related to hydrazines (q. v.) in the same way that amidines are related to amines. Thus the hydroelderide of phenyl-hydrazine acting on a solution of benzimido-ether in absolute alcohol producee dark red needles of di-phenyl-benz-azidine, Ph.C(N11.NPhH): N.NPhH; while form-imidoether gives, when similarly treated, di-phenyl-formazidino CH(NH.NPhH):N.NPhH [185°] (Pinner, B. 17, 182, 2002).

AZIMIDO- COMPOUNDS. This term was originally applied by Griess (B. 15, 1878) to the compounds obtained by the action of nitroue acid upon ortho-diamines, which are probably

of the form R" NH, and was extended by Zincke (B. 18, 3134) to compounds, probably of the form R''()NR', got by oxidising o-amidoazo- compounds (hydrazimido- compounds). The first formula may also be written R" < NH >N, which stands to $\mathbf{R''} <_{\mathbf{N,Cl}}^{\mathbf{NH_2}}$ in the relation that diazobenzene anilide stands to a mixturo of

diazobenzene chloride and aniline.

phenylene-diamine sulphate and KNO, (Ladenburg, B. 9, 222). Needles (from benzene).

Formed by passing nitrous acid into a solution of nitro-o-phenylene diamine (Hofmann, Pr. 10, 406). Long white prisms, v. sol. alcohol and ether, sl. sol. cold water. Nitrous acid does not affect it, hence it does not contain NH2.

Salts.—C.H.KN,O.,—C.H.AgN,O.,. affected by boiling HCl, or KOH. Azimido-toluene C,H,N, i.e.

[83°]. (823°). From aqueous tolylene-o-diamine sulphate and KNO₂ (L.). Prisms containing C.H. (from toluene). V. sol. alcohol, m. sol. ether and boiling water. Not affected by HClAq or

at 160°. Salts,-B'HCI: decomposed by water. B'H,PtCl.

Acetyl derivative C,H,AcN, [1809]. From acetyl-o-tolylene diamine, HCl, and NaNO₂ (Boessneek, B. 19, 1758). Needles.

(B)-Azimide-benzeic acid - C,H,N,O, ЙH

From the hydrochloride of di-amido-benzoio acid, CO2H.C6H2(NH2)2 [1:3:4] and KNO2 (Griese, B. 2, 436). Also by the action of boiling potash upon C₁H₃(CO₂H)(NO₂)(NH.CO.NH₂) [1:3:4] and upon $C_sH_2(\tilde{CO}_2H)(\tilde{NO}_2)(NH.CO.NH_2)$ [1:4:3] (Gricss, B. 15, 1880). This seems to indicate the symmetrical formula. Short needles, containing water of crystallisation; v. sl. sol. water. Not attacked by warm fuming IINO,.

(γ)-Azimide-benzoic acid. Prepared similarly from CO.H.C.11, (NH.), [1:2:3]. Long hair-like needles (G.).

[73°]. From the hydrochloride of o-amido-phenyl carbamic ether and KNO₂ (Rudolph, B. 12, 1295)

Phenyl-azimide-naphthalene

$$C_{1e}H_6 \stackrel{N}{\underset{N}{\bigvee}} N.C_eH_5$$
, [108°]. Fine white needles.

Soluble in hot acetic acid, sparingly in alcohol and benzeno. Formed by oxidation of benzeneazo-(β)-naphthylamine with CrO₂ in acctic acid solution. It is not attacked by strong II, SO, acotie anhydride, or reducing agents (Zincke, B. 18, 3136).

o-Oxy-phenyl-azimido-naphthalene

silky needles. Easily soluble in alcohol and benzone. Formed by oxidation of an alkalino eolution of o-oxy-benzene-azo-(B)-naphthylamine with lead peroxide. It is not attacked by strong H2SO4, acetic anhydride, acetyl chlorido, or redueing agents (Z.).

p-Oxy-pbenyl-azimido-naphthalene

or white needles. Soluble in hot alcohol and hot acetic acid, sparingly in benzene. Formed by exidation of an alkaline solution of p-oxybenzene-azo-(β)-naphthylamino with lead peroxide. It is not attacked by reducing agents.

Acetyl derivative thin silky N.C.II.(OAc) [165°]; platee (Z.).

Telyl - azimido - teluene C, H, N,

$$O_rH_0 < \prod_{N} NC_rH_1$$
, [125°]. $V.D. = 7.23$ (for 8.32).

Formation.—I. By exidation of p-toluene-o-azo-p-toluidine [I:4] C₆H₁McN₂, C₆H₂Me(NH₂) [4:1:2].—2. By heating the imide of o-diazo-toluene-azo-toluene or by boiling ite acctic acid solution, Na being evolved .- 3. Together with an

Control of

amido-phenol or diamine by reducing with SnOl. the azo compounds obtained by combining o-diazo-toluene-azo-tolnene with phenole or aminee (Zincke a. Lawson, B. 19, 1455; 20, 1178).

Properties.—Thin colourlese plates, benzene, hot alcohol, and hot IIOAc. attacked by hot Ac.O, cold H.SO., or reducing agents.

AZINES. Compounds of the typo

sented by the formula

and the nomenclature of other azines may be gathered from two examples (Hinsberg, B. 20, 21).

Phen-naphthazine
$$C_{\bullet}\Pi_{\bullet}$$
 N
 $C_{10}\Pi_{\bullet}$

Tolu-naphthazine $C_{\bullet}\Pi_{\bullet}$
 N
 $C_{10}\Pi_{\bullet}$

Derivatives of quinoxaline

are called quinoxalines, e.g. tolu-quinoxaline −CH \parallel , and naphtho-quinoxaline, N-CH

Formation .- 1. From o-diamines and oquinones or o-di-oxy- compounds .- 2. By oxidation of a mixture of o-diamino and phenole (e.g. (β)-naphthol).—3. By boiling cortain azo-dye. stuffs, derived from secondary amines, with dilute acids; e.g. sulpho-benzeno-azo-phenyl-(\$). naphthyl-amine yields naphtho-phenazine and sulphanilic acid C₂H₄(SO₂H) N₂,C₁₆H₆,NHC₆H₅ $C_{i4}H_{i} < \stackrel{N}{N} > C_{i}H_{i} + C_{i}H_{4}(SO_{3}H).NH_{2}$ (Witt, B.

20, 571).-4. By fusing o-quinones with ammonium acetate, or by heating them with alcoholic NH, (Japp, C. J. 51, 100).

Properties.—Weak crystallino bases; their

salts being decomposed by water. Reduced by SuCl, to hydrides, R NH R', whence ForCl,

regenerates the original azine. Azine-ammenium bases. Compounds of the

. They have also been called form X" ŇŔ(OH) Azonium bases (Witt, B. 20, 1183), a term previously applied by Fischer to quaternary hydra-

zine derivatives.

AZINSUCCINIC ACID

(CO₂H)₂C₃H₂:N.N:C₂H₂(CO₂H)₂. Crystalline solid. Very solubls in water and alcohol. Not decomposed by acids or alkalis. The ethers of this acid are obtained by the spontaneous decomposition of the ethers of diazosuccinio acid C₂H,N₂(CO₂R)₂ on keeping, half the nitrogen being evolved. A'Ba₂: sparingly solublo yellowish-white powder.

Tetra-methyl-cther A'Ms,: [150°]; silky white prisms; easily soluble in het water and alcohol, sparingly in cold water, alcohol, and het ether; not volatile with steam (Curtius a. Koch, B. 18, 1299).

AZO-ANILINE v. Amido-benzone-azo-anilino under Azo- compounds.

AZO-BENZENE v. Benzene-azo-benzeno under Azo- compounds.

TRIAZO-BENZENE and its derivatives v. Diazo-benzene imide and its derivatives.

AZO-BENZOIC ACID v. Carboxy-benzeneazo-benzoie acid under Azo- compounds,

AZO- COLOURING MATTERS.

History.—The series of compounds comprised under this class contains one or more diatomic groups 'N:N' linking together acid or basic arematic radicles. Of a very large number of azocompounds known to science only a certain proportion are of technical value, and these are manufactured in large quantities owing to their importance as colouring matters. The first aze-compound introduced into commerce was the exalate of amidoazobenzene (aniline yellew, v. benzene azo-aniline), C,H, N, C,H, NH, manufactured in 1863 by the firm of Simpson, Manle & Nicholson, by the action of nitrous gas on aniline dissolved in alcohol. Three years later triamideazobenzeno, NH2.C6H4.N2.Č6H4(NH2)2 (v. Amido - benzene - azo - phenylene - diamine under Azo-compounds), was manufactured at Manchester and introduced under the name of 'Manchester brown' ('Bismarck brown,' 'Phenyleno brown, 'Vesuvine'). This compound was prepared by the action of nitrons acid upon mphenylone-diamine. In 1875 diamidoazobenzene, C_sH_s, N₂, C_sH_s(NH₂)₂ (v. Benzene-azo phenylenedismine), was discovered independently by Caro and Witt (B. 10, 213, 350) and introduced into commerce by the latter under the name of 'chrysoidine.' These basic colouring matters were soon followed by acid azo compounds of grsater technical value (B. 10, 1378, 1509), and numerous patents have since that time been taken out, the general mode of preparation being similar in principle to that first employed in 1870 by Kekulé and Hidegh (B. 3, 233). These are the chief points in the industrial history of these compounds; the chemical history will be referred to under the individual compounds. Azobenzene, C.H., N.-C.H., (v. Benzene azo-benzene), which may be regarded as the prototype of the azo-compounds, has been known since 1834 (Mitschsrlich, A. 12, 311), but the industrial development of these products is largely due to the researches of Griess upon the diazo-compounds (A. 106, 123; 113, 201; 117, 1; 120, 125; 121, 257; 137, 39). The theoretical visws which have led to the adoption of the fermulæ at present generally received have been developed chiefly by Kekulé (Z. 1866, 2, 309, 689).

Formation,-1. Compounds of the azobsn-

zene type are produced by the action of mild reducing agents, such as alcoholic potash, alcoholic KOH and zinc dust, iron and acetic acid, or sodium stannito, upon the corresponding nitro-

bodies: $2C_aH_3NO_2 + 4H_2 = C_bH_3N_aC_cH_3 + 4H_aCO_2$. By the oxidation of the corresponding amido-derivatives by potassium permanganata, bleaching powder, obronie acid, hydric peroxids, &c., thms: $2C_aH_3NH_4 + O_2 = C_aH_3N_aC_cH_3 + 2H_aO$ (Glaser, Z_c , [2] 2, 308). This method is applicable to the sulphonic acids and other derivatives of the amido-compounds: $2C_aH_3(CH_3)llSO_3NH_2 + O_2 = N_2(G_aH_aCH_3)llSO_3)llSO_3NH_2 + O_3 = N_2(G_aH_aCH_3)llSO_3llSO_3NH_2 + O_3 = N_2(G_aH_aCH_3)llSO_3NH_2 + O_3 = N_2(G_aH_3)llSO_3NH_2 + O_3 = N_2(G$

3. By the action of cthyldichloramine (Tscherniak, B. 9, 147) upon certain aromatic anines (c.g.p.-tolnidine): 2C,H,NH,+C,H,NCl,=C,H,NH,HCl+HCl+C,H,N,C,H, (Pierson a. Heumann, B. 16, 1048).

4. By the action of nitroso-compounds upon aminos and phenols: C_0H_3 , $NO + \Pi_1N.C_0H_3 = C_0H_3$, $P_1C_0H_3$, $P_2C_0H_3$, $P_1C_0H_3$, P_1C_0

5. From axoxy-compounds by intramolecular transposition, such as the formation of oxyaxobenzene (henzene-axo-phenol) by warming axoxybenzene with strong sulphuric acid

 $C_c\Pi_sN>0$ – $C_s\Pi_sN_2$. $C_s\Pi_s$.OII (Wallach a. Belli, B. 13, 525).

6. By the action of phenylhydrazino upon the naphthoquinones: $C_{10}\Pi_0O_2 + \Pi_1N.NIH.C_0H_3 = C_{10}\Pi_0O_2 + \Pi_2N.NIH.C_0H_3 = C_{10}\Pi_0O_2 + \Pi_2N.NIH.C_0H_3 + H_2O$. The naphthoquinonehydrazide then undergoes intramolecular transposition with the formation of an azocompound: $HO.C_{10}\Pi_0N.N.C_0H_3$. The compound thus obtained from (c)-naphthoquinons is identical with beuzene-azo-(a)-naphthol, while (8)-naphthoquinone gives a compound which is isomeric and not identical with benzene-azo-(b)-naphthol (Zincke a. Bindewald, B. 17, 3026).

Preparation.—The method in general use for the preparation of azo-colours is that depsnding upon the rendiness with which diazo-compounds react with animes and phenols. In practice it is not necessary to isolate the diazo-salt, but the amido-compound which is to be converted into a diazo-salt is treated with the necessary quantity of sodium nitrits and acid to diazotise the NH₂ group, and the solution of the diazo-salt is then mixed with the solution of the amine in acid or the phenol dissolved in alkali. During the process of diazotising, the solution containing the amide-compound must be kept well cooled, as the diazo-salts are very unstable,

especially in aqueous solution. Most aromatic amido- compounds lend themselves to this reaction, the azo- colears of commerce being produced by the action of diazotised amines, amide-sulphonic acids, or amido-carboxylic acids, upon amines, phenols, amido-sulphonic and oxy-snlphonic neids. A selection of typical reactions illustrating the formation of those compounds by the present method may be here

conveniently discussed:

(a) When a salt of diazobenzeno acts upon anilino the first product of the reaction is always diazobenzencanilide (diazoamidobenzene): $\underline{\mathbf{C}}_{\mathbf{e}}\mathbf{H}_{\mathbf{b}}\mathbf{N}_{\mathbf{e}}\mathbf{C}\mathbf{l} + \mathbf{C}_{\mathbf{e}}\mathbf{H}_{\mathbf{b}}\mathbf{N}_{\mathbf{e}}\mathbf{N}\mathbf{H}_{\mathbf{e}} = \mathbf{C}_{\mathbf{e}}\mathbf{H}_{\mathbf{b}}\mathbf{N}_{\mathbf{e}}\mathbf{N}\mathbf{H}_{\mathbf{e}}\mathbf{C}_{\mathbf{b}}\mathbf{H}_{\mathbf{b}} + \mathbf{H}\mathbf{C}\mathbf{l}.$ The latter compound was first produced by the action of nitrous gas upon aniline dissolved in cold alcohol (Griess, A. 121, 258). When allowed te stand in the presence of anilino and aniline hydrochloride, or when acted upon by hydrochlorie acid or unstablo metallio chlorides in the cold, the diazobenzeneanilide is slowly converted into the isomeric amidoazobenzeno, C.H., N. C. II , NIt .. This last compound is manufactured on a commercial scalo for the preparation of other nzo-colours and the indulines (q. v.). In the case of other amines in which the para-position with respect to the NII, group is open (or the ortho-position in the unplithalene series), the transformation of diazo- into amidoaze- compounds takes place with great readinoss. Thus, by the nction of nitrous acid npon o- and m- toluidine, m-xylidino and the naplithylamines, antidoazo-compounds are at once formed.

(b) The salts of the propagatic diazo-compounds react with primary and secondary monanines, but not with tertiary monamines of the fatty series. The resulting products are diazocompounde, such as C_0H_1 , N_2 , $NH.C.H_5$ (diazobenzeno-ethylamide) and C_0H_3 , N_1 , N_1 , N_2 , N_1 , N_2 , N_1 , N_2 , N_3 , N_4 , (diazobenzenediniethylamide). The diazo-compounds of this class do not furnish icomerio azo- compounds (Baeyer and Jäger, B. 8, 148). True azo- compounds of a mixed aromatic and fatty type are produced by the action of diazocompounds upon the sedium derivativee of nitro-hydrocarbons of the fatty series, e.g.

C₂H₂,N₂,N₃ + NaC₂H₁,NO₂ =
C₃H₂,N₂,C₄H₁,NO₂ + NaNO₂
(formation of benzeneazonitroethane, V. Meyer

and pupils, B. 8, 751, 1073, 1078; 9, 384; 12,

(c) Diazo-compounds always furnish true azocompounds with secondary and tertiary aromatic monamines, such as C.H., N., C.H., NMc. (benzene-azo-dimethylaniline) by the notion of diazobenzene upon dimethylaniline (Griess, B. 10,528), and CaHs. N. CaH NILCaH (benzene-azodiphenylamine) hy the action of diazobenzene upon diphenylamine (Witt, B. 12, 259, and C. J. **85**, 185)

(d) The action of diazo-compounds upon aromatic diamines is typified by the action of diazobenzene-chloride upon m-diamidobenzene to produco chrysoïdine: C_uH_1 , N_2 , $C_l+C_uH_4$ (NH_2) = C_0H_3 , N_2 , C_0H_3 (NH_2), HCl. Nitrous acid gives with ortho-diamines a class of substances termed

by Griess azimido- compounde:

 $C_{s}H_{4}(NH_{2})_{2} + HNO_{2} = C_{0}H_{4} < N > NH + 2OH_{2}$ (Hofmann, A. 115, 249; Ladenburg, B. 11, 219; azotoluene-azorecordin by the action of diazotised

Rudolph, B. 12, 1296; Griess, B. 15, 1878). Metadiamines give under the same circumstances compounds of the type of triamidoazo-benzene (Bismarckbrown)2C₂H₄(NH₂)₂ + HNO₂ = (NH₂)C₄H₄, N₅, C₅H₃(NH₂)₂ + 2H₂(O(Caro a. Grices Z. 1867, 278). By acting with an excess of HNO₂ upon p- and m. C₆H₄(NH₂)₂ in acid colution with suitable precautions both these diamines can be diazetised and give aze- colours when combined with phenols (Griess, B. 17, 607

and 19, 317).

(e) The formation of azo- colours from diazocompounds and phenols as illustrative of the general method of technical production is shown by the following typical examples: C, H, N2.NO. + C_sH_sONa - C_sH_sN_sC_sH_sOH + NaNO_s (benzeneazophenol from diazobenzeue nitrate and sodium phenato: Kekulé and Hidegh, B. 3, 233). By a similar reaction would be formed such compounds as benzeneazoresorein, C_sH_s,N_s,C_sH_s(0H)_s (Typke, B. 10, 1876), benzeneazo-(a)-naphthol, C_sH_s,N_s,C_sH_s,OH_s, dre. (ibid. 1580). Azosulphonic acids are formed by the action of diazosniphonic acide upon phenols, or by the action of diazo- salts upon phenolsulphonic acids, thus, for example, n-snlphobenzene-azo-β-naphthol (Poirrier'e 'Orange No. II.'), HSO4.C.H4.N2.C10HB from diazo-benzene sulphonio acid and eodium (B)-naphtholato; benzene-azo-(B)-naphthol disniphonie acid, Cells. N. C10H4(HSO2) OIIs from diazo-benzene and (9)-naphthol sodium dienlphonate ('Orange G,' Meister, Lucius a. Brüning), or p-sulphobenzene - azo - (β) - naphthol disulphonic acid, HSO_3 , C_6H_4 , N_2 , $C_{(0)}H_4$ (HSO_3), OHB from diazotised sulphanilic acid and (B)-naphtholdisulphonic acid sodium salt (Meldola, B. 13, 942).

Classification and Nomenclature.—Azo-com-

pounds may be described as primary, secondary, tertiary, &c. according as they contain 1, 2, 3, &c. N_2 groups. The modes of formation above described have been illustrated by reference te primary azo- compounds, all of which (with the exception of the mixed aromatic and fatty compounds) may be regarded as derivatives of nzobenzene, C_aH_aN_aC_bH_b, benzeneazonaphthalene, C_aH_aN_aC_bH_b, and azomaphthalene, C_aH_bN_aC_bH_b, or generally, in the case of phenolic azo-compounds, as R.N_aPl, where R may etand for C₆H₃, C₁₀H₂, C₆H₄, HSO₃, C₆H₁, NO₂, &c., and Pl for C₆H₄, OH, C₁₀H₆, OH, C10 II s(IISO3)OII, &c. It will be found convenient to write the formulæ of azo- compounds so ae to represent the order of introduction of the radicles. Thus, C_cH_s,N_xC_sH_s,OH indicates the product from diazobenzene and phonel, and would be described as benzene-azo-phenol; $C_6H_1(HSO_4)N_2.C_{10}H_4.OHa$ is p-sulphobenzene-azo-(a)-naphthol, obtained by the action of diazotised-enlphanilie acid upon (a)-naphthol; while C_aH₂.N₂.C_{1a}H₃(HSO₃)OHa is the isomerie benzene-azo-(a)-naphtholmonosulphonic acid obtained by the action of diazobenzone upon (a) naphtholmonosulphonic acid. The eame rulee may be followed with the more complicated types of azo-compounde; thus, for inetance $C_cH_s.N_u.C_cH_s.N_u.C_cH_s(OH)_2$, benzeneazobenzeneazoresoren obtained by the action of diazotised amideazebenzene upon resorcin; in like manner CaH, Na.C.Ha.Na.CaH, (OH) maybe called benzenebenzene azoamidotoluene upon resorcin; and C_aH_a,N_a,C_aH_a , benzeneazodioxybenzeneazobenzene by the successive introduction of two diazobenzene-groups into resorcin. Secondary and tertiary azo compounds of the types $(R.N_a).Pl''$, and $(R.N_a).Pl'''$ have been termed by Wallach disazo and trisazo-compounds (B.15, 22 and 2812; v. also Heumann, ibid. 813).

Secondary and tertiary azo-compounds .- The typical secondary azo-compound of the typo (R.N.),Pl"is the so-called 'phonolbidiazobenzene, (C₆H₅,N₂) :: C₆H₅.OH, discovered by Griess (B. 10, 628). Compounds of this class are formed by the successive introduction of two diazotised radicles into a phonol. The resorcin secondary azo-compounds have been especially studied by Wallach (B. 15, 22; Wallach a. B. Fischer, ibid. 2814), and the crosol compounds by Nölting a. Kohn (B. 17, 351). Analogous amido compounds of the type (R.N.), C.Il.(NII.). have been obtained by the introduction of diazotised radioles into chrysoidine (Griess, B. 16, 2028). Secondary azo: compounds of the type R"(N₂.Pl)₂ havo been obtained by Wallach hy acctylating one amido group in a diamine, diazotising the acctdiamido compound, and combining with a phenol so as to produce a compound of the type B" < N...1'1.Ac' The acetyl group is then removed, the NH2-group diazotised, and the diazocompound R" N L' again combined with a phenol (B. 15, 2825 and Wallach a. Schulze, ibid. 3020). The metadiamines appear to lend themselves most readily to this method. The most direct method of preparing secondary azocompounds of this class is by diazotising both amido groups in a p or m-diamine, and then acting with the product upon a phenol or phenolsulphonic acid (Griess, B. 17, 607 and 19, 817). Another method of obtaining secondary azo compounds of these types is by diazotising a p-nitramido- compound and combining with a phenol or secondary or tertiary amine so as to produce NO, R", N, Pl or NO, R", N, R", NR... The mitro-group is then reduced, the amido-axocompound diazotised and again combined with a phenol, &c., thus producing R"(N2Pl)2, 106 and 47, 657). If the nitro azo- compound ia combined with a monumine and the nitrogroup reduced, a diamidoazo-compound of the type NH. R".N.R".NII, is produced in which both amide groups can be diazotised and combined with phenols so as to form tertiary azo-compounds, I'l.N., R".N., R".N., Pl (ibid.). Secondary azo. compounds of these types are also formed by reducing p-nitracetauilide, diazetising and combining with a phenol or annine so as to produce axo compounds of the type $C_vH < N_{r_v}P_1$. The acetyl group is then removed, the autido-azo- compound diazotised and again combined with a phenol (Nictzki, B. 17, 343 and 1350).

Secondary azo-compounds of the type IV.N...IV.N...P are obtained by diazotising amido-azo-compounds and combining the diazo-azo-salt with phenols. The typical compound,

C₂H₄,N₅,C₂H₄,N₇,C₄H₄,OH, was first prepared by Caro and Sohranbe (B. 10, 2230) and several members of this group have since been introduced into commerce under the names of 'Biebrich scarlet,' 'Croceine scarlet,' &o. These scarlets are of considerable technical importance, especially the latter, which are prepared by the action of diazotised amidoazobenzene-sulphonic acid and its homologues upon (B)-naplithol (a)-sulphonic acid. By the action of diazotised amidoazobenzene upon chrysoidine a tertiary azo-compound of the type [N. 18] N₂, N₃, N₄, N₅, N₆, N₆

K¹N₂, I²N₃ C₄H₄(NH₄)₂ is formed (Griess, B. H. N₂ C₃H₂(NH₄)₂ is formed (Griess, B. H. N₂ C₃H₄(NH₄)₂ is formed (Griess, B. H. N₂ C₄H₄(NH₄)₂ is formed of some importance have recently been prepared by diazotising diamidodiphenyl (benzidine) and its homologues and combining the tetrazosalt with phenols, annues, and their sulphonic acids; of these colouring matters *Congo red, C₄H₄, N₄ C₆H₄(HSO₂)NH₄a, may be mentioned as a typical example. Tertiary azo-compounds derived from triphenyl-carbinol are obtained by diazotising the rosanilines and combining with phenols (Meldola, C. J. 47, 668).

Constitution of Azo-compounds. When s diazo- compound combines with an amine or phenol to form an azo- compound the Ng group invariably takes up the para position with respect to the NH2, NiIR, NR2, or HO, if this position is open. The proof of the constitution of azo- compounds is furnished by the products which they yield on complete reduction. Thus, amidoazobenzene, C₆H₅, N₂, C₆H₄, NH₂, gives on reduction aniline and p-phenylene-diamine; chrysoidine gives aniline and (1, 2, 4)- triamido-benzene (Witt, B. 10, 658). The constitution of some of the azo- compounds obtained by combining diazosulphonic acids with phenols has been established by Gricss (B. 11, 2191). The constitution of the azo-naphthol colours has been determined chiefly by Liebermann and his pupils (B. 14, 1310 and 1795; Jacobson, ibid. 1791; also Witt, C. J. 35, 179). When diazocompounds act upon substituted phenols, such as salicylic acid, the N_z-group also takes up the para-position with respect to the HO-group (P. F. Frankland, C. J. 37, 746); thus (a)naphthaleneazosalicylic acid gives on reduction a-C,0H,.NH, and C,H,(HO)(NH,)CO,H [1:4:2] When the para-position with respect to the NII, or 110-group is already occupied the N₂group enters the ortho position (Nölting a. Witt, B. 17, 77; Liebermann a. Kostanecki, ibid. 130, 876; Griess, ibid. 338; Nölting and Kolin, ibid. 351). This law holds good also when the para position is occupied by an azogroup, as in the formation of (C.H. N.); C.H.OH from benzeneazophenol by the introduction of another C.H., N.-group (Nölting a. Kohn, B. 17, 368). In the case of the (8)-naphthalene derivatives the N .. group also enters the ortho (a)position with respect to the hydroxyl or amidogen group. Thus (\$\beta\$) - naphthol orange, HSO, C, H, N, C, H, OH, gives on reduction sulphanilic acid and (a) amido-(\$\beta\$)-naphthol, the latter yielding (8)-naphthoquinone on oxidation (Liebermann, B. 14, 1310). Similarly the azoderivatives of (8) naphthylamine yield o-diamidonaphthalene $(NH_2:NH_2=a_1:\beta_2)$ on reduction (Lawson, B. 18, 798 and 2422). The fact that such compounds as bonzeneazo-(β)-naphthol are insoluble in cold aqueous alkalis has led Liebermann to the suggestion that these compounds no longer contain HO, but possess the formula C_sH_s,N_cNil_sC_{to}Π_s (B. 16, 2858). A similar conclusion has been suggested by Meldola with respect to the (β)-naphthylamine derivatives, for which he proposes the formula R.N. NH C_{to}H_s, thus indicating a relationsh p to the azimidocompounds above referred to (C. J. 45, 117). This conclusion is supported by the investigations of Zineke and his pupils (B. 18, 3125, 3132 and 3142). This author has shown that these (β)-naphthalene, as well as other orthoamidoazo-compounds, can be exidised to com-

position of the B.Nz-groups with respect to the

HO as well as upon the formation of bodies of the oximido-type. This appears to be the case

especially with the azo-derivatives of resorcinol (Wallach, B, 15, 22 and 2814).

General properties and technology. The individual azo- colours will be described under their respective headings, so that it will be only nocessary here to give a brief account of the general characters of these colouring matters. The parent azo-compounds, C_aH_a , N_a , C_aH_b , C_aH_b , N_a , $C_{10}H_{7}$, $C_{10}H_{7}$, N_a , $C_{10}H_{7}$, although possessed of colour are not of any value as dyestuffs: it is only when acid or basic radicles are present in one or both aromatic nuclei that true colouring matters are formed. The colours produced by these compounds are of various shades of yellow, orange, brown, red, scarlet, indigo-blue and violet. No pure blue or green azocompound is known. Of the basic primary azocompounds, amidoazo-benzene, chrysoidine, and 'phenylene brown,' show a gradation in shado from yellow through orango to brown, last compounds are now alone of technical value. Among the acid azo- colours a regular gradation of shade is also observed with the increase in molecular weight. This is well seen in the colours produced by the action of diazobenzene and its homologues upon the naplitholsulphonic acids, the lowest members of the series being orange, and the bighest members scarlet of an increasing shade of blueness. The acid primary azo- compounds are dyed from acid baths and have great affinity for silk and wool, but do not readily dye cotton without the use of mordants. Primary azo colours as a ruie dia-Vol. I.

solve in strong sulphnric acid with a red or orange colonr. Azo- colonrs can be produced directly in the fibre of cotton by impregnating the latter with a phenol and a diazo compound, and then developing by means of an acid (T. Holliday, S. C. I. 1, 204). A similar process has been described by Grässler (S. C. I. 4, 262) who uses the nitrite in the form of an ammonium salt or in combination with ammonium salts, so that on heating the impregnated fabrio the nitrous acid is liberated in the presence of the amine and phenol, and the colour is developed without the use of acid. Oxyazo-compounds, which are not sulphonic acids, and which are insoluble, can be made soluble by warming them in aqueous or alcoholic solutions of bisulphites. soluble double compound is thus formed which is decomposed on heating with the liberation of the colouring matter so that the process is especially applicable for calico printing (Meister, Lucius a. Brüning, B. 17, 452). The secondary azocolours derived from diazotised amidoazobenzene and its homologues in combination with naphtholsulphonic acids are scarlets of greater tinctorial power and purity than the primary azo- scarlets, and possess a certain affinity for coffon, especially the 'croecine scarlets' already referred to. The secondary azo-colours derived from diazotised benzidine ('Congo red' series) and its homologues are reds or yellows which also possess a certain affinity for cotton, but many of these colours are too sensitive to acids to be of much use technically. The stability of these colours is increased, however, by using the higher homologues of benzidine. It is only among secondary azo- compounds that true shades of violet and indigo blue are found (Nietzki, B. 17, 344; Meldola, C. J. 47, 665). Some of these blue shades are now met with in the market, a typical compound of the class being a salt of ditolyltetrazo-(A)-naphtholdisulphonic acid, C_1H_6 , N_2 , $C_{16}H_2$ (HSO₃)(OH) β (Azo - blue of F. C_1H_6 , N_2 , $C_{16}H_3$ (HSO₃)(OH) β

G.H., N.C., L.H. (1180.) (OH) Azo-blue of F. Bayer & Co.). Secondary azo-compounds dissolve as a rule in strong sulphuric acid with a violet, blue, or green colonr. The tertiary azo-compounds have but little tinetorial value. For observations on the absorption spectra of azo-colonrs see papers by H. W. Vogel (B. 11, 623), Landauer (B. 14, 391), and Stebbins (Am. 6, 117 and 149).

AZO- COMPOUNDS. (V. also Azo- Golouring MATTERS and Di-Azo- compounds.) The nomon-elature of these compounds is based on the following rules. Imagine N, displaced by H, and the compound to break up accordingly, one H going to one half and one to the other. If both the resulting compounds are aromatic, name the compound richest in carbon according to the rules followed in this dictionary, and profix azoto the name. Before azo- write the name of the hydrocarbon from which the other half of the molecule is derived. Then prefix all the substituents of the latter so that they may follow one another in this order: chloro-, brome-, iodo . cyano-, nitro-, oxy-, amido-, sulpho-, and car-boxy-. When both halves of the azo- compound contain the same number of atoms of carbon, then the half which contains CO.H is put last. If neithor contain CO,II, the one containing BB

SO,H comes last; if neither contain this, pre-ference is given to OH, NH₂, NO₂, Br, or Cl, in succession.

If one half of an azo-compound is derived from a fatty hydrocarbon the name of this half is written last.

The compounds obtained by the action of diazo- compounds upon (8)-naplithylamine, (8)naphthol, or more generally upon amido- or oxycompounds in which the p-position is already occupied, are described as if they were ordinary azo- compounds. It is, however, not improbable that they may have a different constitution. Thus the compounds obtained from diazo- compounds and (B) - naphthylamine behave like diazoamides in their decomposition by acids into (8)-naphthylamine, a phenol, and nitrogen; but act on reduction like amido-azoderivatives giving (1:2) - naphthylene - diamine and the amine. Three views may be taken with regard to the constitution of these bodies: -(1) That they are true diazonmides; (2) that they are o-amido-azo-derivatives; (3) that they are hydrazimido- compounds, i.e. dihydrides of azimido- compounds.

The first hypothesis easily explains their decomposition by acids; and their reduction to naphthylene-diamine might be accounted for by assuming an intermediate change into the amide-aze- compound. However, the fact that a different isomeric body is formed by diazetising (3)-naphthylamine and combining it with the other amine does not agree with this view, eince compounds of the form X.N...NIIY and Y.N. NHX are always found to be identical. The other reactions of these bodies are also inconeistent with this hypothesis. The hypothesis that the compounds are o-amido-azo- derivatives does not readily account for their behaviour on oxidation, which tends to show that they do not contain an NH2 group.

The third hypothesis leads to the formula

It is strongly supported by the fact that on oxidation these bodies give rise to white crystalline azimido compounds, which probably have

the constitution
$$C_{10}H_0$$
NR, analogous to

Griess's azimidobenzene C.H. NH obtained

by the action of n'trous acid on o-phenylenediamine. The corresponding compounds from (8)-naphthol and diazo-salts may also be obtained by the action of hydrazines upon (8) naphthoquinone, and hence may be represented in a similar manner by the formula

The formation of hydrazimido-compounds from a diazo-salt and (6)-naphthylamine can be explained by assuming the intermediate formation of true diazo - amides C10 II, NH.N:NR, which by taking up H₂O (or HCl) would form R'.

C₀H₁,NH₂N(OH),NHROC₁₀H₂,NH₂NH₂NH₂NH₃NH₄N(OH)R, azo to and by again eliminating H₂O it might give the salt withwdrazimido-compound. The decomposition by

acids might also be explained as a change inverse to the above, resulting in the formation of the diazo-amide, which would then be decomposed. The ortho-amido-azo- derivatives of the benzens scries would also seem to be similarly constituted, for on oxidation they also give colourless crystalline azimido- compounds.

Apparently opposed to the hydrazimidohypothesis, is the bohaviour of these bodies towards nitrous acid, which converts them into diazo-compounds. In many points, however, these diazo-compounds greatly differ from the ordinary diazo- compounds, and their constitution may possibly be expressed by the formula:

HO.N.N. , which would represent their form-N.N.R'

ation from hydrazimido- compounds. SnCl, or SO, these diazo-compounds are not reduced to hydrazines but to non-basic stable bodies which probably have the constitution

The o-diazo-imides readily lose N-N.R N2 on heating and are converted into azimido-

N.R' identical with those compounds R" obtained by oxidation of the o-amido-azo- compounds. On the hydrazimido- hypothesis the

body derived from diazo-benzene and phenyl-(β)-naphthylamine would be represented by the N.C.II, formula C10H which agrees with its

 $_{N_{s}HC_{o}H_{s}}^{H}$ which agrees with its Thus cone. IICl removes aniline, reactions.

forming an azine $C_{10}H_{4} \langle \downarrow \rangle$ C,H, while oxi-.

dising agents produce a powerful ammonium base $C_{10}H_{\circ}N_{\circ}(C_{\circ}H_{\circ})_{\circ}OH$ (Meldola, C.~J.~45,~107; Nölting a. Wilt, B.~17,~77; Lawson, B.~18,~796, 2422; Sachs, B. 18, 3125; Zincke, B. 18, 3132, 3142; Zincke a. Lawson, B. 19, 1452).

DESCRIPTION OF AZO- COMPOUNDS.

Acetophenene-azo- (β) -naphthel [4:1] C.H. (CO.CH.) - N2-C10H. (OH). Formed by diazotising p-amido-acetophenone and combining the diazo- compound with (\$)-naphthol in alkaline solution (Klingel, B. 18, 2695). Slender red needles. V. sol. alcohol, sl. sol. ether, insol. water. Dyes silk a dull red.

o-Amide-benzene-azo-aceto-acetic acid C₆H₄(NH₂)-N₃-CH(CO.CH₃).CO₂H. [157°]. Orange-red tables. Easily soluble in acctic acid, alcohol, ether and chloroform, sparingly in water. Formed by reduction of o-nitro-benzene-azo-acetoacetic acid with FeSO, and NH, (Bamberger, B. 17, 2420).

p-Amide-benzene-p-aze-aniline C₄II₄(NH₂)—N₂—C₆II₄(NH₂). p-Azo-aniline. Di-amido-azo-benzene. [235°]. Long flat yellow

needlos. Easily solublo in alcohol, sparingly in benzene and ligroine.

Preparation .- 1. The mono-acetyl derivative of p-phonylene-diamine is diazotised and combined with aniline, the anilide thus obtained is dissolved in aniline and warmed with aniline hydrochloride by which it is converted into the mono-acetyl derivative of amido-benzene-aso

aniline, which is finally saponified.—2. From its di-acetyl derivative which is formed by the action of zinc and ammonia on acetyl-p-nitro-anilins (Mixter, Am. 5, 282).

The mono-acid salts are green, the di-acid red. B"H₂Cl₂: nsedles.

Mono-acetyl derivative C₁₂H₈N₂(NII₂)(NIIAc). [212^o], glistening golden yellow plates. Its salts are red (Niotzki, B. 17, 345).

Di-acetyl derivative [282°] (M.). Amide-benzsns-axo-aniline

[4:1] C,H₄(NH₂)-N₂-C,H₄NH₂ [1:4]. [142°]. Obtained by reducing nitro-benzene-azo-nitro-benzene [206°] with alcoholic ammonium sulphide (Janovsky, M. 6, 460). Minute plates (from alcohol). Salt.—B'II,Cl.. This body is probably identical with the preceding, the melting-point having perhaps besn misprinted.

m-Amido-benzens-m-azo-aniline

[8:1]C_uH₁(NH₂) - N₂ - C_uH₁(NH₂) [1:3]. From the nitro-compound by ammonium sulphido (Janovsky, M. 6, 458). Minute flat yellow needles with green lustre (from alcohol).

Amido-benzene-szo-benzene-p-snlphonic acid [4:1] C_aII₄(HSO_a)—N:N—O_aII₄NI, II₄ [1:4] S. ·0144 at 22°. Yellowish-white microscopic needles or scales (containing aq). Nearly insoluble in water, alcohol, ether, and chloroform. Formed by the action of diazo-benzene-p-sulphonic acid on aniline (to extent of about 30 p.c.). Propared from benzene-azo-anilino and fuming II₂SO₄ at about 70°. On reduction with tin and IICl it gives p-phenylene-diamino and sulphanilic acid.

Salts.—A'NH,*: orange - yellow plates.— A'.Ba 6aq: sparingly soluble trimetric orange needles.—A',Ca 2aq (Griess, B. 15, 2184).

Amido-benzene-azo-benzene p-snlpbonio acid [4:1]C,H₁(SO,H1)—N:N—C,H₁,NIL,[14]. S. 0196 at 22°. Pearly plates (containing aq). Formed by reduction of the sparingly soluble nitro-benzene-azo-benzene-p-sulphonio acid with ammonium sulphide. On complete reduction with and HGl it gives p-sulphanilio acid and p-phenylene-diamine, and hence ought, according to theory, to be identical with the preceding compound; this, however, does not appear to be the case.

Salts.—KA'aq: yellow trimetric plates.—
*NaA': needles.—BaA'6aq: largo glistening
monoclinic needles.—CaA'4aq: yellow pearly
plates.—PbA'.: monoclinic plates (Janovsky,
B. 16, 1488; M. 4, 279, 652).

Di-amido-bsnzene-aze-benzens-p-sulphonic acid (NI₂)₂C₂H₃-N₂-C₂H₄,SO₂H. Chrysoïdine sulphonic acid. Brownish-red glistening needles. Sl. sol. water.

Formation.—1. By combination of p-diazobenzsno-sulphonic acid with m-phsnylene diamine.—2. By sulphonation of chrysoidine. On reduction it gives sulphanilic acid and (1:2:4)tri-amido-benzone (Griess, B. 15, 2196).

m-Amido-benzsns-p-azo-di-msthyl-aniline [8:1] C₈H₄(NH₂)—N₈—C₈H₁NMc₂ [1:4]. [166°]. Golden lamina, sol. alcohol. Formed by the action of warm dilute H₂SO₄ on its acetyl derivative [184°] which is obtained by mixing diazotissd acctyl-m-tolylene-diamine with dimethylaniline (Wallach, A. 234, 263).

p-Amido-benzsns-azo-dimethylanilino [4:1] NH₂.C₈H₄.N₂.C₈H₄.NMe₂ [1:4]. [188°]. By warming the alcoholic solution of the nitrocompound NO₂.C₆H₄.N₂.C₆H₄.NMs₂ with ammonic sulphide (Meldola, C. J. 45, 107).

Properties.—Brick-red needles (from dilute alcohol). Insol. boiling water, but forms yellow solutions in most solvents. Reduced by zinc dust and HCl it gives dimethyl: p-phenylene-diamine and p-phenylene-diamine. On adding acetic acid to an alcoholic solution, the liquid turns green, on dilution with water it becomes red. In conc. H.SO₄ the solution is orange. A very dilute solution mixed with nitrous acid and exposed to the air turns blue (test for nitrous acid).—B"(HCl)_PtCl.

Methylo-iodide.—Insoluble brown scales, Acetyl derivative.—Orango needles (from alcohol). [217°].

p.Amide-bsnzsne-azo-(β).naphthol [4:1] NH_x,C_zH_y,N:N.C_{1z}H_y,OH [a:β]. Formsd by reducing the corresponding nitro-compound with annonium sulphide and annuonia. Forms a crimson solution in H_zSO₄ (Meldola, C. J. 47,668).

p-Amido-benzens-azo-(a) naphthol [4:1] NH₂.C₃, II, N₂.C₄, II₄.OH [a:B]. Formed by reducing para-nitro-benzene-azo-(a)-naphthol in hot dilute NaHO with ammonium sulphide; ppd. by HCl (Meldola, C. J. 47, 662). Dark brown powder; sl. sol. hot water, v. sol. hot alcohol. Forms a red solution in conc. H.SO..

p-Amido-benzsns-azo-(\$\beta\$) - naphthol di-sul-pbonie acid. Acetyl derivative C.H.(NHAC)-Nz-CieH.(OH)(SOzH)z. Goldon glistening plates. Searlet red dyc-stuff. Formed by diazotising the mono-acetyl derivative of p-phenylene-diamine and combining it with (\$\beta\$)-naphthol (\$\beta\$)-di-sulphonic acid (modification insoluble in alcohol). By saponification of the acetyl group it yields a bordeaux-red dye stuff. By diazotising the latter and combining it with (\$\beta\$)-naphthol di-sulphonic acid a blue colouring matter is produced (Nictzki, \$\beta\$, 17, 344).

p-Amido-benzene-aze-(a)-naphthylamine [4:1]NH_...C,H_.N.,.C_,H_.,NH...[1:4]. [160°]. Formed by reducing NO...C_aH_.,NH...[1:4]. [160°]. Formed by reducing NO...C_aH_.,N.,.C_i.H_s.NH., with aqueous animonio sulphido (McIdola, C. J. 43, 432). Ochreous needles. Readily soluble in alcohol, acetone, benzene and chloroform. Its salts form crimson aqueous solutions; excess of acid throws down the neutral salts.—B"(HCl)_*PtCl_*.

p. Amido-benzene-aze-o-oxy-bsnzeic acid [4:1] NIL_C_1L_-N__ C_4H_(OH)(CO_III) [1:4:2]. From the sodium salt of p-nitro-benzene-azo-salicylic acid by reduction with animonium sulphide. Colonrloss need s. Sl. sol. boiling water. Its alkaline solutions are yellow. Glacial HOAc forms a crimson solution. Blackens at 219°-220° C. (Meldola, C. J. 47 667).

m-Amido-benzene-szo-phenol
[3:1] C₆H₁(NH₂) - N₂—C₆H₂OH [1:4].
[168°]. Brownish yellow scales. Obtained by
saponification of the acetyl derivative.

Acetyl derivative C₁H₄(NHAc).N_xC₆H₄OH [c. 208°]. Prspared by diazotising the mono-acetyl derivative of mphenylene-diaminoand combining it with phenol. (Wallach, B. 15, 8020).

p.Amide-benzene.azo-pbsnel [4:1] NH₂C,H₂,N:N.C,H₂,OH [1:4]. [181^o]. Obtained by heating p-nitro-benzsns-azo-phenol with ammonium sulphide. Brown scales (from

p-Amido-benzene-azo-diphenylamine

[4:1] NH, C, H, N2. C, H, NHC, H, [1:4] [0. 91°]. Got by reducing the nitro- compound by ammonio

eulphide (McIdola, C. J. 43, 440).

Properties.—Sl. sol. in boiling water, v. sol. alcohol, acctone, chloroform and benzene, forming yellow solutions. IICl added to the alcoholic solution turns it first green, then rod. Solution in cono. II, SO, is violet, turned red by

Salts .- Form crimson aqueous solutions.

Dye wool orango.

m-Amido-benzene-azo-m-phenylene-diamine [8:1] C₆H₁(NH₂)-N₂-C₆H₃(NH₃)₂ [1:2:4]. Bismarck brown. [137°]. Ppd. by adding NaNO₂ to a neutral solution of m-phenylene-diamino hydrochlorido (Caro a. Griess, Z. 1867, 278). Brown plates. Sl. sol. in hot water; v. e. sol. alcohol and ether. Dyes wool brown. Absorption spectrum (Hartley, C. J. 51, 180). Salts.—B"2HCl.—B"H₂PtCl_o.

p-Amido-benzene-azo-phenylene-diamine *[4:1] C_aH_a(NII_a)-N_a-C_aH_a(NH_a), [1:2or3:4]. From benzene-azo-benzene by nitration and roduction (Janovsky, M. 6, 466).

p-Amido-benzene-azo-resorcin

[4:1] NH₂.C₆H₄.N:N.C₆H₃(OH)₂.[1:2:4]. By dissolving p-nitro-benzene-azo-resorein in diluto NaHO and warming with anmonium sulphide (Meldola, C. J. 47, 661). Its alkaline solutions are red; its acid solutions are pale orange. -B', II, SO,: silvery scales. - B', H, PtCl,

p. Amido-benzene azo-m-xylidine (4:1) NIL.C.H.,N.C.II.,Ne.NII. [I:3:5:2] [I63°). Formed by reducing NO.C.II.,N.C.II.Mc.NII. with aqueous ammonio eulphide (Meldola, C. J. 43, 432). Golden ecales (from water). V. sol. alcohol and benzene.

Salts.—The acid salts are very soluble in water. Excess of HCl forms an amorphous brown pp. of B"2HCl.—(B"2HCl)PtCl... Amido-thiophene-azo-benzene

C.H.N.,C.SII.,NH,. Formed by adding diazobenzeno chloride to a tolerably concentrated solution of thiophenine hydrochloride.-BIICL aq: yollow needles, solublo in water and alcohol (Stadler, B. 18, 2317).

Amido - thiophene - azo - bonzene - p - sulphonio acid [1:1] C₆H₁(SO₇H),N₂,C₁SH₂,NH₂. Formed by combination of diazo-benzone-p-sulphonic acid with thiophenine. Yellow needles, red when dry. Sl. sol. water and alcohol. Dyes silk yellow. (Stadler, B. 18, 2318).

Amido-thiophene-azo-naphthalene

C₁₀H.,N.,C.(SH.,NH₂. Forned by adding (a)-diazo-naphthaleno chlorido to a solution of thiophenine hydrochloride. The hydrochloride forms microscopic red needles, sparingly solublo in water and alcohol (Stadler, B. 18, 2318).

m-Amido-tolucne-azo-aceto-acetic acid (4:2:1) C_sH_s(CH_s)(NII₂).N₂-CII(CO.CII₂).CO₂H [162°]. Red glistening needles. Formed by reduotion of m-nitro-tolucne-azo-acoto-acetic acid with FeSO, and NII, (Bamberger, B. 17, 2421).

Amido tolnene azo amido cresol [4:3:1] C₄II₃(CH₃)(NII₃),N₂,C₄H₂(CH₃)(NII₃)(OR), Oxyazo-toluidine [212°]. Small dark red needles, V. sol. alcohol and ether, sl. sol. water. Formed from amido toluene azoxy toluidine

water); v. sol. alcohol.—B'_H_PtCl_ (Meldola, | C_H_Me(NH_1).N_2O.C_H_Me(NH_1) by intramole C. J. 47, 658). cular obange by heating it with H.SO, at 110°. It is reduced by SnOl, to tolylene-diamine C.H.3Mc(NH.)2 [I:2:4] together with diamidoeresol (Limpricht, B. 18, 1405).-B"H2SO4-B"2HCl.-B"H2PtCl.

Amido tolnene azo (8) naphthol

[6:3:1] $C_0H_3(CH_3)(NH_2)-N_2-C_{10}H_4OH$ [1:2]. Acetyl derivative

C₆H₃(CH₃)(NHAc)-N₂-C₁₀H₆OH [276°]. Insol. water, sl. sol. alcohol, m. sol. a mixture of alcohol and chloroform. Red colour. Formed by diazotising the mono-acetyl derivative of (1:2:4)-tolylene-diamino and combining it with (β)-naphthol (Wallach, B. 15, 2830).

Amido-tolnene-azo-nitro-ethane.

Acetyl derivative C₁₁II, N₀3 i.e. [6:3:1] C₆H₃Mc(NIIAc).N₂.CH(NO₂).CH₃. [143°]. From C₆H₃Mc(NHAc).N₂Br and NaCII(NO₂)CH₄ (Wallach, A. 235, 250). Red needles (from alcohol-ether); v. sl. sol. water, insol. ligroin.

Amido-tolnene-azo-o-toluidine [4:3:I] $C_6H_3Me(NH_2)-N_2-C_6H_3Me(NH_2)$ [1:4:3] [197°]. Formed by reducing nitro-o-toluidine C_nH₂Mc(NO₂)(NH₂) [1:4:2], or the corresponding azoxy-compound, by sodium amalgam in alcoholio solution (Limpricht, B. 18, 1406; Graeff, A. 229, 350). Long red noedles (from alcohol) or small yellow needles (from water). Sparingly soluble in water, easily in alcohol and other.

Salts.—B"H,SO,: Slender reddish needles
-B"2HCl.—(B"2HCl)PtCl,—B"2HBr.

Amido-toluene-azo-p-toluidine [6:3:1] C.H.Me(NH.)—N.—C.H.Me(NH.) [1:6:3]. [159°]. Red needles. Sl. sol. cold, v. sol. hot, wator; v. sol. alcohol. Prepared together with the hydrazo-compound by the prolonged action of sodium amalgan on an alcoholic solution of nitro-p-toluidine (Buckney, B. 11, 1453).

Di-amyl-amido-benzene-azo-di-amyl-aniline [4:1] $(C_bH_{11})_2N.C_bH_4-N_2-C_bH_4.N(C_bH_{11})_2$ [1:4] Di-amyl-aniline azyline. [115°]. Red pointed erystals. Sol. hot alcohol. Formed by passing NO through an alcoholio solution of di-amylanilino. Salts. — B"(C₀H₂(NO₂)₃OII)₂: small yellow orystals. Periodide B", I₆: small black erystals with violet reflex (Lippmann a. Fleissner, B. 15, 2142 and B. 16, 1419)

Bonzene-azo-aceto-acetic acid

Call N-N2-CHAc.CO2H. [155°]. Yellow leaflets (from alcohol). Prepared by the action of a solution of diazobenzene nitrate on an alkaline solution of acctacetio ether (V. Meyer, B. 10, 2076).

Salts.—A'K: yollow glistening leaflets.
Bah', Pbh', Cuh', and Agh' are yellow pps.
Ethylether [75°]. Light yellow crystals;
very readily saponified (Züblin, B. 11, 1417).

Benzene azo acetono C.H.N., CH. CO.CH. [149°]. Glistening yellow prisms or needies. Peculiar characteristic smell. Only slightly soluble in hot water, and in aqueous alkalis.

Formation.—I. By heating benzene-azo-aceto-acetic ether with a dilute alcoholic solution of NaOH.—2. By heating benzeno-azo-acto-acetic acid to 170°-180°, CO₂ being evolved (Richter a. Münzer, B. 17, 1928).

Benzene-azo-acetophenone C.H., N., CH., CO.C., II. [129°]. Slender golden needles. V. sol. hot alcohol and hot acetic acid. Formed, together with benzene-azo-benzoylacetic ether, by adding a solution of diazobenzene chloride to an iced alkaline solution of | it forms benzene-ano-di-phenyl-thio-urea (q, v), benzoyl-acetic ether (Bamberger a. Calman, B. 18, 2563).

Benzene-o-azo-aniline (?)

[123°]. Formed by C.H.-N.-C.H.NH. (?). reducing benzone-o-azo-nitro-benzene with ammonium sulphide (Janovsky, M. 8, 61; yellow crystals with blue reflex (from dilute nicohol). The salts are less soluble in water than those of the p-compound.

the p-compound.

Benzene-azo-aniline C₁₂H₁₁N₃ i.e.

C_cH₃—N₃—C_cII,NH₃ [I:4]. Anido-azo-benzene.

Mol. w. 197. [126°]. (above 360°).

Formation.—1. By reducing benzene-azo-mitro-benzene [137°] with ammonium sulphido (G. Schmidt, Z. 5, 417; Janovsky a. Erb, B. 18, 1186).—2. Together with bromo-aniline by the action of bromine vargur nanon aniline. the action of bromine vapour upon aniline (Kekulé, Z. [2] 1, 688).—3. By action of mineral acid (one molecule or less) upon diazo-benzene anilide (v. DI-AZO- COMPOUNDS) (Martius a. Gricss, Z. [2] 1, 132; Friswell a. Green, C. J. 49, 746). Preparation .- 1. Diazo - benzene - anilide is

dissolved in 2 or 3 times its weight of aniline, 10th its weight of aniline hydrochloride is added at the ordinary temperature, and the mixture is kept for an hour at 30° to 40°; after standing for twenty-four hours at the ordinary temperature, sufficient HCl is added to combine with the free aniline and the amidoazo-benzene base precipitates, or it can be obtained as hydrochloride by adding more HCl. The yield is nearly theoretical (Witt a. Thomas, C. J. 43, I13; Staedel a. Bauer, B. 19, 1953). - 2. A eone solution (of rather less than 1 mol.) of NaNO, is added to (1 mol. of) aniline hydrochloride dissolved in (5 or 6 mols. of) aniline at 30°-40°, kept at e. 40° for 1 or 2 hours, and then at the ordinary temperature for 12 hours; completed as above, the yield is nearly theo-

Properties .- Yellow crystals; separates from benzene with benzene of crystallisation (W. a. T.); orange prisms with blue reflex (from alcohol). V. sl. sol. hot water, m. sol. ether and alcohol. Its salts are decomposed by water; they dya wool yellow. Tin and HCl give anilino and vphenylene-diamine. Combines with Et1 forming the hydriodide of CoH1(NIL.).N2.CoH1,NHEt. Thediazo-compounds of benzene-azo-aniline and its sulphonic acids are used for the preparation of scarlets by combination with the naphthols and their sulphonic acids.

Salts. - B'HCl: steel-blue needles or scales (from boiling HClAq). -B'2H2PtCl6. -B'HNO3. -B',H,SO, -B',H,C,O, Acetyl derivative C,H,N,C,H,(NHAc).

[142°]. Yellow silky crystals. On reduction with alcoholic ammonium sulphide it gives acetyl-amido-hydrazobenzeno (Schultz, B. 17, 463; Berju, B. 17, 1400; C. C. 1884, 571). Reactions.—1. By boiling henzene-azo-ani-

lins with 10 pts of HCl (S.G. 1-12) it is completely decomposed in a few hours with formation of p-phenylene-diamine, aniline, chlori-nated-hydroquinones, NH, and colouring matters (Wallach a. Kölliker, B. 17, 395) .- 2. An aleoholic solution of bromine forms a dibromoderivative [152°] which is reduced by tin and HCl to aniline and p-phenylene-diamine (Berju, an ethereal solution of sinc ethyl it reacts form-B. 17, 1400).—3. With phenyl thio-carbinide ing ethane (1 vol.), ethylene (3 vols.), and a

together with some benzene-azo-di-phenyl-thiourea-azo-benzene (v. dis-azo compounds).—4. Carbonyl chloride forms benzone-azo-di-plienylurea-azo-benzene (Ph-N2-CaH4.NII)2CO (Berju, B. 17, 1404).-5. Benzone-azo-aniline hydrochloride (2 mols.) heated with acctone (I mol.) at 150°-160° under pressure forms a base C1, H1, N3, [205°]. Yellow needles; v. sol. alcohol, other, and acids. By tin and HCl it is reduced to a base of melting-point [185°]. Dilnte solutions of the salts have a blue fluorescence. Salts .-B'H2SO4: slender soluble needles or small mono elinic prisms. - B'H.Cr.O.; orange-yellow needles. -B'H2Cl2PtCl4: flat yellow needles, sl. sol. cold water (Engler a. Schestopal, B. 20, 180).

Benzene-azo-benzene C₁₂H₁₀N₂ i.e. C_LH₂—N₂—C_LH₃. Azobenzene. Mol. w. 182. [68°]. (293°). V.D. 6·5 (calc. 6·3). S. (alcohol) 8·5 at 16° (Moltchanoffsky, J. R. 1882, 224). S.V. 220·4 (Ramsay).

Formation.—1. By treating nitro benzene with alcoholic potash (Mitscherlich, A. 12, 311; Schmidt a. Schultz, A. 207, 328), sodinn-amalgam in presence of alcohol and acetic acid (Werigo, A. 135, 176; Alexejeff, Z. [2] 3, 33), iron (3 pts.) and aectic acid (1 pt.) (Noble, A. 98, 253), or with zinc-dust (Alexejeff, Bl. [2] 31, 684).— 2. By oxidising aniline with squeous KMnO₄ (Glaser, A. 142, 364), red-hot PbO (Schichuzzk, J. R. 6, 245), H₂O₂ (Leeds, C. N. 41, 210; B. 14, 1382) or bleaching powder (Schmitt, J. pr. [2] 18, 195) .-- 3. It is the chief product of the action of sodium on p-brome-aniline (Auschütz

a. Schultz, B. 9, 1398; cf. Claus, B. 15, 315).

Preparation.-1. By heating nitro-benzone with a solution of sodium stannite prepared by dissolving the theoretical quantity of SnCl₂ in an excess of aqueous NaOH (Witt, B. 18, 2912).— 2. 400 grms. of NaOH (98 p.c. powdered) are hoiled with 2000 c.e. of ordinary alcohol till most has dissolved; 500 g. of nitrobenzene nro slowly added to the boiling solution, and the formation of azoxybenzene completed by 2 or 3 hours' cohobation. 200 grass, of zinc dust are then slowly added and the boiling continued for n day with occasional shaking. The alcohol is distilled off on a salt bath, warm water addod, the insoluble portion filtered off, washed, freed from Zn(OII), by HCl, and extracted with alcohol; the filtrate on cooling deposits the azobenzene in splendid large plates; good yield.

Properties. - Trimetric plates (Jeremejeff). Crystallises with C_bH_a from benzene. Its absorption-spectrum has been described by Hartley (C. J. 51, 176).

Reactions .- 1. Pa sed through a red-hot tube it yields benzene and diphenyl (Ferko, B. 20,664). 2. Chromic acid in acetic acid at 200° forms benzene-azoxy-benzene. - 3. Nitric acid forms Ph.N., C.H., (NO.) [1:4], Ph.N., C.H., (NO.) [1:2], [4:1] C.H., (NO.), N., C.H., (NO.) [1:4], [1:3] C.H., (NO.), N., C.H., (NO.) [1:3], and a nitro-

benzene - azo - di - nitro - benzeno. — 4. Alcoholio ammonium sulphide reduces it to hydrazobenzene (Hofmann, Pr. 12, 576) .- 5. Hot conc. HCl, HBr, or HI reduces it to benzidine, other products being formed at the same time .-6. Alcoholio SO, produces benzidine.—7. With an ethereal solution of zinc ethyl it reacts formproduct which, when treated with water, gives aniline. 80 g. of azo-benzene gives 70 g. of aniline. The reactions are probably:—

(a) $PhN_2Ph + 2ZnEt_2 = 2NPhH(ZnEt) + 2C_2H_4$

(b) $PhN_{a}Ph + 4ZnEt_{2} = 2NPh(ZnEt)_{2} + 2C_{2}H_{a} + 2C_{2}H_{a}$

And then, on adding water: (a') NPhII(ZnEt) + II,O = NPhH₂ + EtH + ZnO (b') NPh(ZnEt)₂ + 2H₂O - NPhII₂ + 2EtH + 2ZnO (Frankland a. Louis, C. J. 37, 560). - 8. Aldahyde at 200° forms a compound CasH...N.O. [164°]. In presence of chloride of zine, aldehyds condenses with it to benzylidene-bonzidino Ph.CH:N.C.H, C.II, N:Cll.Ph (?) [239°] (Barzilovsky, J. R. 1885, 366). 9. Bromine forms mono-bromo- derivatives, a di-bromo- dorivativo, [205°] (v. Bromo-benzene azo-neomo-nenzene), and a tetra-bronio - derivative, C, II, Br, N2,

[0. 320°] (Werigo, A. 165, 200).

Combinations. — C₁₂H₁₀N₂C₈H₈ [38°]. —
(C₁₂H₁₀N₃)₂ 3HCl: unstable yellow crystals. —
(C₁₂H₁₀N₃)₃ 3HBr: unstable red crystals, got by passing IIBr into a solution of azobenzene in CS_2 — $C_{12}\Pi_{10}N_2HBr_3$: crystals, formed by adding bromino to a solution of the preceding body in ohloroform.—C₁₂H₁₀N₂Br_a: red prisms, got by adding excess of bromino to a solution of benzenc-azo-benzeno in chloroform (Werigo, A.

Benzene-aze-benzane sulphonic acid

C.H. - N. - C. II SO. II [1:4]. [127°]. From benzene-azo-benzeno and fuming ILSO, (5 pts.) at 130° (Griess, A. 131, 89; 151, 208; Janovsky, M. 2, 221; 3, 237; B. 15, 2576). Orange-red plates (containing 3aq). Sl. sol. alcohol and ether. Potash-fusion converts it into K,SO, and benzenc-azo-phenol. Ammonium sulphido followed by mineral acid converts it into diamido-di-plienyl sulphonio acid. Nitric acid Janovsky, M. 3, 508).

Salts. KA' 2aq. BaA', AAA'.

Chloride C₁₂I₁₈N₂SO₂Cl. [82°]. Orange

clumps (from other).

Amide CuH,N,SO2NH2. Powder (Skanda-

roff, Z. [2] 6, 613).

Benzens-azo-bonzana disulphenie acid O.H. - N. - C. H. (SO.II) [1:2:1]. Formed, to-

gether with s-m-, and s-p-, sulpho-benzene-azo-bsnzeno sulphonic acids by heating benzeneszo-benzeno (1 pt.) with pyrosulphuric acid (4 pts.) at 150° (Janovsky, M. 3, 237). Very dsliquoscent needles. Isomerides of this acid are described as sulpho-beuzens-azo-benzens

sulphonic acid.

Banzens-azo-banzeie acid

C_eH_a,N_a,C_elI₄(CO₂H) [1:4]. Azo - benzene - p - carboxylic acid. Obtained by saponification of carboxylic acid. its nitrils by boiling with KOH. Long glistening brown prisms. Sol. alcohol, other, and warm benzeno. Salts.-A'K: very soluble brownish-yellow needles .- A'Ba: brownish-yellow needles; sol. alcohol, sl. sol. water (Mentha a. Heumann, B. 19, 3023).

Nitrile Call N. C. H. (CN) [1:4]. p-Cyano-azo-benzene. [101°]. Formed by diazotising benzanc-azo-anilins and allowing the solution of diazo-benzene-azo-benzene chloride to drop into a hot solution of CuSO, and KCN. Brown noedles. V. sol. ether, benzene, and warm alcohol, insol. water. Sublimable (M. a. H.).

Bensene-aze-bensoyl-acetic acid C_zH_zN_z.CH(CO.C_zH_z).CO_zH. [141°]. Long yel-low needles. V. sol. alcohol, ether, and acetic acid. Its ethyl-ether is formed, together with benzene-azo acetophenone, by adding a solution of diazobenzens chlorids to an iced alkaline solution of benzoyl-acetic ether. By boiling with dilute NaOH it is converted into banzensazo-acetoplicnono C₆H₃,N₂,CH₂,CO.C₆H₃ (Bamberger a. Calman, B. 18, 2563).

Benzene-azo-benzylidane-aniline

C,H,-N2-C,H,N:CH.C,H,Benzylidene-amidoazo-benzene. [125°]. Orange plates. Formed by the action of henzaldehyds on benzene-azoaniline. By HCl it is resolved into its constituents (Berjn, B. 17, 1403).

Benzans-azo-o-bromo-benzene

C, H, N2 C, H, Br [1:2]. Bromo - azo - benzene. [87°]. Glistening plates (from alcohol). Sl. sol. cold alcohol. By the action of bromino (I mol) upon a warm acctic acid solution of benzensazo-benzeno (I mol.) a mixture of o, m, and p, niono-bromo-benzene-azo-benzeno is obtainsd; they can be separated by their different solubilities in alcohol. By complete reduction they give aniline and o., m., or p-bromaniline.

Banzane-azo-m-bromo-banzane

C.H., N., C.d., Br [1:3]. [56°]. Yellowish-gresn pearly plates. V. sol. alcohol, ether and acetone. On nitration it gives orange needles of C., H., Br (NO₂)N₂ [123°] (Janovsky a. Erb, B. 20, 359).

Benzsne-azo-p-bremo-bsnzene

C.H., N., C.II Br [1:4]. [82°]. Is the chief product of the bromination of benzenc aze-benzenc in acetic acid. Orango yellow play with limable. V sol. alcohol, ether to solve the corresponding hydrog which needles of C.H. and the di-nitro-derivation of the di-nitro-derivation of the passing Benzens-azo-nuow Benzens-azo-nuow azo-resorcin and by including the passing azo-resorcing the passing azo-resorcing the passing azo-resorcing the passing azo-resorc duct of the bromination of benzenc azo benzene

Benzens-aze-p-chioro-benzenetann a. Floissnet C_aH₄.—N₂.—C_aH₄Ol [1:4]. Ch [89°]. Yellowish-brown plateid ...mes in

brown needles. Easily soluble iffether, benzeue, and hot alcohol, sparingly in cold alcohol.

Preparation .- 100 grms. of benzene-azo-aniline hydrochloride are suspended in 2 litres of water and 220 c.c. of conc. HCl, and diazotised by slow addition of a conc. solution of 20 grms. sodium nitrite. After standing for soms time the diazo-azo-benzeno solution is filtered and slowly added to a boiling solution of 40 grms. Cu,Cl, in 360 c.c. of conc. HCl, and boiled for some time; the greyish-black pp. is treated with conc. HCl and then with dilute NaOH to remove impurities, and extracted with hot alcahol; the alcoholic solution after treatment with animal charcoal deposits the benzene-azoohloro-benzeno on oooling in glistening brown plates; the yield is 38 p.o. of the theoretical (Heumann a. Mentha, B. 19, 1686).

Reactions.-When its alcoholic solution is allowed to stand in the cold with SnCl, and 2 drops of H.SO, it is converted into a chloro-di-amido-diphenyl NH₂,C₂H₄,C₅H₅Cl.NH₂. The

latter body is not formed, however, by heating the hydrazo- compound, previously prepared, with HCl, but benzene azo-chloro-benzene, chloro-aniline, and aniline are formed inetoad. On nitration with tuming HNO₃ benzene-azo-ohloro-benzene gives p-chloro-benzene-azo-p-nitro-benzene C₆H₄Cl.N₂C₆H₄(NO₂). By treatment with tuming sulphuric acid it is converted into p-chloro-benzene-azo-benzeno p-sulphonic acid C_eH₄Cl.N_{..}C_eH₄(SO₃H) (Mentha a. Heumann, B. 19, 2970).

Benzene-azo-o-cresoI

C₆H₅—N₂—C₆H₈(CH₅)(OH) [1:3:4]. [130°]. Glistening yellow plates. V. sol. alcohol, ether, chloroform, and benzene; sl. sol. hot, insol. cold, water. Dissolves in dilute alkalis with a yellowish red colour. Obtained by the action of diazobenzene chloride on an alkaline solution of o-cresol. It readily gives a dis-azo-compound when treated in alkaline solution with a further quantity of diazobenzene chloride.

Acetyl derivative: [82°]; yellow tables,

v. col. alcohol, other, and benzenc.

Benzoyl derivative: [111°]; small yellow needlee, v. sol. other, acetone, and hot alcohol (Liebermann a. Kostanccki, B. 17, 130; Nölting a. Kohn. B. 17, 363).

Benzene-azo-m-cresoI

C₆H₃-N₂-C₆H₃(CH₃)(OH) [1:2:4]. [109°]. Yellow needles. V. sol. alcohol, ether, ehloroform and benzenc. With alkalis it forms yellowish-red salts. Obtained by the action of diazobenzene chloride on an alkaline solution of m-cresol. It readily combines with another mol. of a diazo-compound to form dis-azo-compounds (Nölting a. Kohn, B. 17, 366).

Benzene-azo-p-cresol C.H. -- N2 -- C. 112 (CH2) (OH) [1:5:2]. Orange yellow plates. Soluble in dilute alkalis. Dyce a canary yellow. Formed by the action of diazo-benzene chlorido on an alkaline solution of p-cresol. It does not yield a dis-azo-compound.

Acetyl derivative Ph-N₂-C,H₃(OAc): [68°]; yellow needles, v. sol. aloohol, ctbcr, ohloroform, and acetone.

Benzoyl derivative Pb — N₂—C,H_e(OBz): [113°]; yellow needles, sol. ether, benzone, and hot aloohol (Mazzara, G. 9, 425; Liebermann a. Kostanecki, B. 17, 130; Nölting a. Kohn, B. 17, 852).

Benzene-azo-p-oresoI-sulphonic acid $C_aH_a-N_a-C_aH_a(CH_a)(OH)(SO_aH)$ Small reddish brown tables or needles. Easily soluble in water, sparingly in alcohol. Formed by the action of diazobenzene oliloride on an alkaline solution of p-cresol sulphonic acid [14:2]. A'Na: soluble reddish brown plates, dyes wool an orange yellow (Nolting a. Kohn, B. 17, 357; cf. Stebbins, A. C. J. 1, 465; 2, 263).

Benzene-azo-y-cumenol C.H. N. C.H.(CH₃₎,OH [1:3:5:6:2] [94°]. Glietening brown prisms. In email quantities it can be distilled undecomposed. Insoluble in alkalis. Formed by combining diazo-benzene chloride with \u03c4-cumenol [70°]. On reduction it yielde aniline and amido-v-comenol [167°] (Liebermann a. Kostanecki, B. 17, 886).

Benzene - azo - di - ethyl - amide - benzoio acid C.H. -N. -C.H. (NEt.) CO.H[1:4:2]. [125°]. From diszo-benzene nitrate and di-ethyl-m-amido-benzeic acid. Red crystals with viclet lustre

(from alcohol). Insol. water, al. eol. alcohol and ether. Salte: BaA'₂.—AgA' (Griose, B. 10, 526). Benzene-azo-othyl-(β)-naphthyl-amine

C,H,-N2-C,H.NHEt. [103°]. Red needles. Soluble in alcohol &c. with an orange-red colour, insol. water. It forms bluish-violet salts with acids. Formed by heating ethyl-(B)-naphthylnitrosamine with an acetic acid solution of aniline at 100°. Also produced by combining

diazo-benzene with ethyl-(β)-naphthylamino.

Nitrosamine C_gH_g, N_g,C_{tg}H_g, NEt(NO): [97°]; red crystals (Henriques, B. 17, 2669).

Benzene-azo-ethyl-phenylene-diamine C_aH_z, N_a, C_aH_z(NH_a)NHEt. Ethyl-chrysoidine. Dyes a redder shade than ordinary chrysoïdine. Formed by combining ethyl-m-phenylenodiamine with diazo-benzenc. - B"HCI: reddishbrown needles with violet-blue reflection, soluble in water and alcohol with an orange colour .-B",H,Cl,PtCl,; insolublo red pp. (Nölting a. Stricker, B. 19, 517).

Benzene-azo-indoxyl C,II,-C(OII) C₆H₅-N₂-N<

Red needles or thick orange prisms. [236°]. Sol. alcohol and alkalis, v. sl. sol. water. Formed by the action of diazo-benzene chloride on indoxyl (Baeyor, B. 16, 2190).

Benzene-azo-methane C_eH₅.—N₂.—CH₃. Azo-phenyl-methyl. (c. 150°). Yellow oil of peouliar odour. Very volatile, and readily distils with steam. Formed by oxidation of s-phenylmethyl-hydrazine C. II, NH.NH.CII, with HgO

(Tafel, B. 18, 1742).

Benzene-azo-methazonio aoid Ph-N2-C2H3N2O3. [164°]. From diazo-benzene nitrate and an aqueous solution of sodium methazonate (Kimich, B. 10, 141). Orange

needles (from alcohol); insol. water. - Na, A"2ail. --BaA" aq.

Benzene-azo-di-methyl-amido-benzolo acld Ph-N₂-C_eH₂(NMe₂)CO₂H [1:4:2]. From diazobonzene nitrato and di-mcthyl-m-amido-benzoic aoid (Grices, B. 10, 527). Orange nodules.

Benzene-azo-methyl aniline O.H., N. C.H., NHMe. Methyl-amido-aso-ben-zene. [180°]. Red needles. Sol. alcohol. Formed by heating benzene-azo-aniline with Mel.— B'HCl: violet needles.

Acetyl derivative CaH, N2.OaH4.NMeAc: [189°]; yellow silky needles (Berju, B. 17, 1401).

Benzene-azo-di-methyl-aniline

C.H., N., C.H., NMe., Disnethyl-amido-aso-ben-sene. [115°]. Yellow plates. Preparation.—1. A solution of 74 pts. of

NaNO₂ (100 p.c.) and 40 pts. of NaOH in 540 pts. of water is slowly added to a cooled solution of 100 pts. of aniline, 130 pte of dimethylaniline, and 230 pts. of HCl in 360 pts. of water Griess, B. 10, 525: Möhlau, B. 17, 1490).—2. By heat-ing benzene-azo-aniline with McI (Berju, B. 17, 1402; C. C. 1884, 871).

Methylo-iodide B'Mel. [1740]. Plates:

Benzene-azo-methyl-ketole

O.H.-N.-C.H.N. [116°]. Yellow needles, Formed by the action of diazo benzeno-ohloride upon methyl-ketole in aqueoue alcoholic eclution in presence of codium-acetate (Fischer, B. 19,

Bensene-aze-methyl-phenylene-dismine C₂H₂,N₂·C₂H₃(NH₂)NHMe. Methyl-chrysoldine. Minute orange prisms. Dyee a somewhat rodder shade than ordinary ohrysoldine. Formed by combining diazobenzene with methyl-m-phenylene diamine (Nölting a. Strioker, B. 19, 549).

Benzene -azo-(a)-naphthol

C₄H₃-N₂-C₁₀H₄OH. (a)-Naphthoquinonephenyl-hydrazide. [206°]. Sinal needles with bluo reflection (from alcohol) or flat needles or plates (from benzene).

Formation.—1. By combining (a)-naphthol with diazobenzene.—2. By the action of phenyllydrazino hydrochloride upon (a)-naphthoqui-

none suspended in acetic soid.

Reactions.—Dissolves both in acids and alkalis. Bromine forms two bromo-derivatives [185°], and [196°] (Margary, G. 14, 271). Salts.—A'K: orystalline pp. The hydro-

chloride, hydrobromide, and sulphate form bluish or greenish glistening needles, sparingly soluble in water, alcohol, and acetic acid (Typko, B. 10, 1580).

Methylether A'Me: [83°]; brown crystals.

Methyl ether A'Me: [83°]; brown crystals.

Ethyl ether A'Et: [100°]; long needles.

Acetyl derivative G, II, N (OAc) [128°];
small brownish-red needles (Liebermann, B. 16,
2858; Zincke a. Bindewald, B. 17, 3026).

Benzene-o-azo-(a)-naphthol

$$C_{16}H_{\bullet}$$

$$\downarrow_{N_{\bullet}H_{\bullet}G_{\bullet}H_{\bullet}S}^{Oa}$$
 $\uparrow_{N_{\bullet}H_{\bullet}G_{\bullet}H_{\bullet}S}^{Oa}$
 $\uparrow_{N_{\bullet}H_{\bullet}G_{\bullet}H_{\bullet}S}^{Oa}$

quinone-phonyl-hydrazide. Formed by the action of phenylhydrazine hydrochlorido upon (β) -naphthoquinone. Long red needles; sol. lot alcohol and hot acetic acid. It does not combine with acids or bases. By SnCl_ it is reduced to (β) -amido- (α) -naphthol. By treatment in hot acetic acid solution with IINO₃ it gives di-nitro- (α) -naphthol. By the action of alkaline reducing agents it is at once converted into (β) -amido- (β) -naphthol; no intermediate product could be detected (Zincke a. Bindewald, B. 17, 3030; Zincke a. Rathgen, B. 19, 2482).

Bromo-derivative $C_{\rm lo}H_{\rm 10}N_{\rm s}{\rm Br}_{\rm s}O$; [215°-219°]: elember red needles.

Benzene-azo- (β) -naphthol $C_{ic}\Pi_{12}N_2O$ i.e. $C_{il}\Pi_{12}N_2O$ i.e. $C_{il}\Pi_{1}N_2C_{10}\Pi_{0}O\Pi$ or $C_{10}\Pi_{0}$ $N_2H.C_{0}\Pi_{12}$ [1:2].

Formation.—By the action of a diazobenzone salt on an alkaline solution of \$B-naphthol. It is not formed by the action of diazobenzene hydrate on thedy divided (\$B-naphthol (difference from (a)-naphthol) (Liebermann, \$B, 16, 2858).

Properties.—Golden plates or long metallic glistening needles. Solublo in ether, benzene, ligroine and CS.—Dissolves in H.80, with a magenta-red colour. Is insoluble in caustic alkalis. It has a slightly basic character, dissolving in HCl with a red colour, and forming an unstable hydrochloride crystallising in needles. By SnCl. it is reduced to (a)-amido-(β)-naplithel and aniline. The eame reduction products are also formed at once by treatment with alkaline reducing agents (e.g. zinc-dust and LOH, ammonium sulphide, &c.); no international account account acid solution with HNO, it gives

di-nitro-(6)-naphthol (Zincke a. Rathgen, B. 19, 2482).

Reamondering time C. H. Ren O. (1682)

Brown-derivative C_{1e}H₁₁BrN₂O: [168°]. Brownish-red needles (Zincke a. Bindewald, B. 17, 3031).

Benzene-azo-(α)-naphthol sulphonic acid C_aH₃-N₂-O_aH₃(SO_aH)(OH). From sodium (α)-naphthol sulphonate, aniline nitrate, and KNO_a (Hofmann, B. 10, 1378). Slender brown needles (from alcohol mixed with HClAq). Dyes orange.

-ΔνΔ'.—Rna'.

—AgA'.—BaA'.

Benzene-azo-(β)-naphthol sulphonic acid C₀H₂.—N₂-C₀H₄(IISO₂)OH. Brown neodles with yellowish-green reflex. Difficultly soluble in water and alcohol. Prepared by the action of an alkaline solution of (β)-naphthol-sulphonic acid on diazobenzene nitrate or chloride. —A',Ba. Yellowish red microscopic leaflets. Slightly soluble in water (Griess, B. 11, 2197). Its absorption-spectrum has been examined by Hartley (C. J. 51, 196).

Benzène-azo (β) -naphthol disulphonic acld $C_s\Pi_s - N_2 - C_{10}H_1(SO_2H)_2(OH)$. From sodium (β) -naphthol disulphonate and diazo-benzene nitrate. Sol. water. Barium salt is sl. eol. water (Stebbins, jun., A. C. J. 2, 236).

Benzenc-azo-(a)-naphthylamine

C₄H₅—N₂—C₁₀H₆NH₂. Prepared by the action of diazobenzene sulphate on (a)-naphthylaurino (Griess, T. 1861, iii. 679; Weselsky a. Benedikt, B. 12, 228).—B'.Hi.SO,4aq: microscopic needles; difficultly soluble in water.

Benzene-azo- (β) -naphthylamine $C_0\Pi_3-N_2-C_{10}H_0N\Pi_2$ or $C_{10}H_0$ $N_2\Pi_1C_0H_2$

Benzene-hydrazimido-naphthalene. [101]. Ited trimetric tables or long fine red needles. Easily soluble in alcohol and acctic acid, insoluble in water. Dissolves in strong II, SO₄ with a blue colour. Its salts exist only in presence of a large excess of acid. Formed by combination of diazobenzene with (β)-naphthylamine. By boiling with 20 p.o. aqueous II, SO₄ it is slowly decomposed into (β)-naphthylamine, phenol, and nitrogen. On reduction it gives antiline and (1, 2)-naphthylene diamine. On oxidation it gives benzene-aximido-naphthalene (q. v.).

Acetyl derivative C₁₆H₁₂AcN₃. [153°]. Small red needles, easily soluble in alcohol, insoluble in water.

Benzoyl derivative C₁₆H₁₂BzN₂: [163°], rod crystals (Lawson, B. 18, 796).

Benzene-o-azo-nitro-benzene

C_bH_z-N_z-C_cH_z(NO_z) [1.2]. Nitro-azobenzene.

[123°]. Formed by nitration of benzene-azo-benzene in acetic acid at 100°. Orange-yellow minute needles. V. sol. alcohol. Alcoholio NaOH gives a beautiful emerald-green colouration; by long boiling or by treatment with sodium-amalgam it is further reduced to a compound C_z(H₁₈N_zO. Tin and HCl or animonium sulphide first reduce it to bonzene-o-azo-aniline and thon to aniline, and o-phenylene-diamine. By bromination in acetic acid it yields brome-nitro-benzene [123°] or [132°] (Janovsky a. Erb, B. 19 _2157; 20, 360; M. 8, 56).

Benzene-p-azo-nitro-benzene C₄H_{*}, N₂, C,H_{*}, NO₂ (1:4), [137°]. Small yellow needlos (from alcohol). Formed by nitration of benzene-azo-benzene. By NH_{*}HB it is reduced

ordinary benzone-azo-aniline. Alcoholic | NH, HS produces red crystals of an intermediate C, H, N, C, H, NOH

aitrolio soid C.H.N.C.H.NOH ?) [134°],

which forms a blue solution in NaO11Aq. K, FeCy, reoxidises it to benzene-azo-p-nitrobenzene. By complete reduction it yields anilino and p-phenylene-diamine (Laurent a. Gerhardt, 4. 75, 73; Janovsky, B. 18, 1133; M. 6, 164, 455). Benzene-azo-tri-nitro-benzeno

C₄H₅-N₂ -C₄H₄(NO₂)₃. [112°]. By action of HgO on an alcoholic solution of the corresponding hydrazo-compound (E. Fischer, A. 190, 133). Slender red prisms (from alcohol).

Benzene-azo-nitro-iso-butane

C₆H₅-N₂-CH(NO₂)Pr. From diazobenzene nitrate and potassium nitro-iso-butane (Züblin. B. 10, 2088). Oil. Forms an orange solution in alkalis.

Beazene-azo-nitro-ethane C, II, N,O., i.e. C₃H₃-N₂-CH(NO₂).CH₃. [137°]. From diazobenzene nitrate and sodium nitrocthune (V. Meyer a. Ambühl, B. 8, 751, 1073). Rectangular orange crystals; v. sol. alcohol and ether, insol. cold water, sol. aqueous alkalis forming a bloodred solution. Dyes silk yellow. Conc. H.SO, forms a violet solution. It crystallises unaltered from aqueous NII, (Barbieri, B. 9, 386) but such a solution gives pps. with metallic salts.
Salts.—C,H,K,N,O,4aq.—C,H,Na,N,O,7aq.

- C, H. ZnN, O, 3aq. - C, H. PbN, O, PbO 2 jaq.

Benzene-az :-nitro-methane

C₄H₅—N₂ · CH₂NO₂ [153°]. From diazobenzene nitrate and sodinin nitromethane in very dilute solution (Friese, B. 8, 1078). Slender red needles (from alcohol). Conc. II,SO, gives a purple solution. Decomposed by HClAq only after long boiling.

Benzene-azo-tri-nitro-phenol

C₈H₅-N₂-C₈H(NO₂)₃O11. Long brown prisms. Insol. cold, sl. sol. hot, water; v. sol. alcohol. Explodes at 70°. Dyes silk and wool orange-yollow. Prepared by the action of a salt of diazobenzene on an alcoholic solution of pierie acid (Stebbins, jun., A. C. J. 1, 465; 2, 236; C. N. 41, 117; B. I3, 43).

Benzene-azo-nitro-propane

C.H. -N. - Cl1(NO.).ClL..Cll. [99°]. From potassium nitropropane and diazobenzene nitrate (V. Meyer, B. 9, 386). Orange needles; sol. alkalis.

Benzene-azo-iso-nitro-propsne

C, II, -N, C(NO)(CII3), From aqueous diazobeuzene nitrate and potassium nitro-isopropane. Oil; insol. alkalis (V. Meyer a. Ambühl, B. 8, 1076).

Benzene-azo-o-oxy-benzoic-acid

CaHa-Nz-CaHa(OH).CO.H. From diazobenzone nitrate and an alkaline solution of salicylic acid (Stebbins, jun., A. C. J. 1, 465; B. 13, 715; C. N. 41, 117). Orange-red needles; insol. water, v. sol. alcohol or ether. Dycs wool orange,

Benzene-azo-di-oxy-naphthaleae

 $\mathbf{C}_{0}\mathbf{H}_{3}$ — \mathbf{N}_{2} — $\mathbf{C}_{10}\mathbf{H}_{3}(\mathrm{OH})_{2}$ or $\mathbf{C}_{10}\mathbf{H}_{3}(\mathrm{OH})$ $\left\{ egin{array}{l} \mathbf{O} \\ \mathbf{N}_{m}\mathbf{H}\mathbf{C}_{n}\mathbf{H}_{s} \end{array} \right.$ Oxy (a) naphthoquinone phenyl hydrazide. [230°]. Formed by the action of phenylhydrazine on oxy-naphthoquinone in aqueousalcoholio solution. Yellowish-red glistening needles. V. sol. ether, hot alcohol, and but acetio acid.

Salts.-The alkali salts form slender orange needles, sol. aloohol.—A',Ba 10aq: yellowish-brown platos or long red needlos.—A',Ca 4aq: slender orange ncodles .- A'Ag: rcddish-brown amorphous pp.

Acetyl derivative: [179°]; red needles. Methyl other A'Mo: [175°]; red needles. Ethyl ether A'Et: [173°]; yellowish-red needles, v. sol. hot alcohol and hot acetic acid.

Bromo-derivative C, H, BrO, N2: [198°]; large red needles, sl. sol. alcohol (Zinoko a. Thelen, B. 17, 180)).

Bcnzene-azo-phenol $C_0H_3-N_2-C_0H_4.OH[1:4]$.

Oxy-azo-benzene. [154°].
Formation.—1. A product of the action of BaCO, on diazobenzene nitrate in the cold (Griess, A. 137, 81), -2. From benzene-azo-benzene sulphonic acid by potash fusion (Grioss, A. 154, 211). 3. From diazo-benzene nitrate and C.H.OK (Kekulé a. Heidegh, Z. [2] 6, 384).-4. By the action of p-nitroso-phenol (20 pts.) on aniline acetate (60 pts.) (Kiruich, B. 8, 1499) .-5. By gently warning benzene-azoxy-benzene, C_aH_s , $-N_aO-C_aH_s$, with 11_aSO_4 (Wallach, B. 13, 525; 14, 2617).

Properties. -Purple pyramidal plates. Insol. cold water, sol. alkalis; v. sol. alcohol. Bromine in acetic acid gives a compound [139°].

Salt, $-\Lambda g \Lambda'$.

Acetyl derivative. - Ph.N. C. H.OAo. [85°]. (above 360°). Golden scales.

Methyl ether. — Ph.N₂,C₀H,OMe [54°]. Yellow scales.

Benzene-azo-phenol snlphonic acid

CaH3-N2-C6113(OH)(SO3H)[1:4:3]. From diazobenzene nitrate and an alkaline solution of phenol o-sulphonic acid (Griess, B. 11, 2194). Red needles or tables; sol. water and alcohol. -KA'.

Benzene-azo-phenol sulphonio acid

C₁₂H_aN₂(OH)(SO₂H). From benzene azo phenol and fuming 11₂SO₄ (Tschirvinsky, B. 6, 560).— BaA',2aq.—CuA',6aq.—MgA',6aq.—KA'. Benzene-azo-phenol di-sulphonio acid

C₁₂H.N₂(011)(SO₃II)... From azoxybonzene (1 pt.) and fuming IL₂SO₄ (10 pts.) by heating for 3 hours at 150° (Limpricht a. Wilsing, A. 215, 232; B. 15, I297). Orango needles, v. e. sol. water, v. sol. dilute acids.
Salts.-K₂A" 2aq: red needles, sl. sol. cold

water; not attacked by Br. -BaA"aq: orange crystalline pp. -Ag₂A": red crystalline pp.

Benzene-azo-phenol tetra-sulphonic acid *C_{0.1}I, M_.(OH)(SO₃11)₄. From axoxybenzone and luming H₂SO₄ (L. a. W.). Will not crystallise. Salts. - C₁₂H_.N_.(OK)(SO₃K), 7¹/₂ aq; long yellow needles. Gives with bromine-water app. of tri-bromo-phenol [92°] .- Ba A 7aq: orange crystalline pp.

Other benzene-azo-phenol snlphonic acids are described as Oxy-benzene-Azo-benzene sulphonic acids, and sulpho-benzene-azo-phenol sulphonic acids.

Benzene-azo-diphenylamine

C₆H₅-N₂-C₆H₄,NH.C₆H₅. [82] (O. N. Witt, C. J. 35, 185; B. 12, 259). A solution of diphenylamine (17 g.) in alcohol (100 o.c.) is added to a solution of diazo-benzene chloride (14 g.) in alcohol (50 g.). The brown mixture is cooled with ice, and alcoholio NMe₃ (used instead of NII, because NMc, HCl is soluble in aloohol) is

added from time to time to neutralise the liquid. The oil which separatee is purified by a lengthy procees, and finally crystallised from benzoline.

Properties .- Golden leaflets or needlee. Sol. benzene, alcohol, and ether. HCl turns its alcoholic solution violet; steel-grey erystals of the hydrochloride separating. The base forms a green solution with H.SO, turned, by adding water, into indigo, violet, and finally red. With amyl nitrite and acetic acid it forms a nitrosamine, Ph.N(NO).C,11,N,Ph, [120°], forming orange needles, sparingly soluble in alcohol and acetic acid, readily in benzene.

Benzeus-azo-m-phenylsne diamine

 $\mathbf{C_4H_4}$ — $\mathbf{N_2}$ — $\mathbf{C_4H_4}$ (NH.) $_2$ [1:2:4]. Chrysordine. [117·5 $^\circ$] (W.); [110 $^\circ$] (H.). From diazobenzens ealte and m-phenylene diamine (Witt, B. 10, 850, 654; Hofmann, B. 10, 213, 388; Griess, B. 10, 390). Slender yellow needles (from water); sl. sol. water, v. sol. alcohol. Reduced by ammonium sulphide at 150° to aniline and (1, 2, 4)tri-amido-benzene. The absorption-spectrum has been examined by Hartley (C. J. 51, 178) .-B"HCl: black octahedra or small red sitky cryetale; its aqueous solution is orange, but turned red by HCl-B",H,PtCl,.-B"HNO3.

Di-acctyt derivative Ph-N₂-C, H, (NHAc), [251°].

Benzsne-azo-m-phsnylsne-diamine salphonic acid C₄H₃-N₂-C₅H₄(NII₂)₂(SO₂H). Chrysoïdine sulphonic acid. Glistening spangles or needles. Sl. sol. water and alcohol. Prspared by ths action of dinzobenzene-chloride on a sult of m-phenylene-diamino sulphonic acid. A'Na: eolubls golden needles—A',Ba: orangs needlss (Ruhemann, B. 14, 2655). By the sulphonation of chrysoidino an isomeric acid is formed (v. Di-amido-benzene-azo-benzene sulphonic acid).

Benzsns-azc-di-phenyl-methyl-pyrrol-carboxylic acid C24H19O2N, i.e.

HO,C.C.CMe

N.C.H. N. C. IIs. [195°]. Obtained HC:CPh/

by saponification of its ethyl ethor, which is formed from an acctic acid solution of acctophsnono-aceto-acetie ether (1 mol.) and benzenep-azo-aniline (1 mol.) on standing for two days. Large red erystals. V. col. alcohol, ether, and benzene; sl. sol. ligroin and acctic acid; insol. water, and conc. acids. Gives Laubenheimer's reaction.

Ethyl ether Eth': [123°]; splendid red crystals; v. sol. benzene and ligroin, sl. sol. alcohol, other, and acetic acid (Paul a. Schneider, B. 19, 3162).

Benzsne-azo-phonyl-(8)-naphthyl-amine C₂₂H₁,N, i.e. C₆H₅-N₂-C₁₆H₆NHC₆H₅ or N.C₆H₅

/y TIC"H

Benzene - phenyl - hydrazimido - naphthalene. [1420]. Obtained by combining diazo-benzene chloride with phanyl-(B)-naphthyl-amine in alcoholic colution. It is also formed by the action of an acetic acid solution of aniline upon phenyl-(3)-naplithyl-nitrosamine. Small red glietening needles.

Reactions .- 1. By heating with conc. HCl it

yielde naphthophenazins C.H.

anillne .- 2. Bromine acting on the hot acetio acid solution forms tetra-bromo-phenyl-(8)naphthylamine with evolution of nitrogen.-8. It is reduced by SnCl, to aniline and phenylonaphthylsne diamine C10H3(NH2)NHC,H4.-4. By oxidation with K2Cr2O, in acetic acid solntion the chromats of a powerful ammoniumbass C22H16N,OH is formed; the latter possibly N(OH).C,H,

has the constitution $C_{10}H_e$ N.N.C.H.

Naphthalene di-phenyl-AZAMMONIUM-hydrate (Henriques, B. 17, 2671: Zincke a. Lawson, B. 20, 1167).

Benzene-azo-di-phenyl-thic-urea

C, H, -N, -C, H, NH.CS.NHPh. [179°]. Plates. Formed by combination of phenyl-mustard-oil with benzene azo-aniline (Berju, B. 17, 1105). Benzene-azo-pyrogallol

C_sH₃—N₂—C_sH₁(OH)₃. Prepared by adding an aqueous solution of diazobenzene nitrate to an

alkaline solution of pyrogallol (Stebbins, jun., A. C. J. 1, 465; 2, 236; B. 13, 44; C. N. 41, 117). Rad needles (from acetic aeid). Insol. water, sol. alcohol. Its alcoholic solution dyee eilk and wool orange.

Benzene-p.azo-resorcin

 $C_6H_5-N_2-C_6H_3(Oll)$, [1:2:1]. [161°] or [170°]. From diazobenzens nitrate and resorcin (Typke, B. 10, 1577; Wallach, B. 15, 2819; R. Meyer, B. 16, 1329). Formed also by gently warming diazobenzene anilide with resorcin, aniline being eliminated (Heumann a. Oceonomides, B. 20, 905). Slendsr orange needles [170°] or short red needles [161°], insol. water, sol. aqueoue alkalis, v. c. sol. alcohol.

Acetyl derivative [102°]. Mono-ethyl ether [87°]. Soarlet needlee. Diethyl ether [70°]. Yellowish-red needles. Banzene-o-azo-resorcin C.H., N., C.H. (OH) [1:2:6]. Formed in small quantity (about 5 p.e.)

in the preparation of the p-isomeride.

Mono-ethylether CaH, N2.CaH3(OH)(OEt) [150°]. Long fins scarlet needles; v. s. sol. alcohol and sther, insol. water; dissolves in aqueous

alkalis with a brownish-red colour. Di-ethyl-ether C.H., Nz.C.H. (OEt)2: [90°].

Large red glistening tables; v. sol. ether, hot alcohol and acetic acid, incol. water (Pukall, B. 20, 1145).

Banzene-azo-thymol

 $C_8H_8-N_2-C_8H_2MePr(OI')$ [1:2:5:4] [85°-90°]. From diazobenzane chloride, and an alkaline solution of thymol (Mazzara, G. 15, 52, 228). Reddish-yellow needles. Reduction followed by exidation gives thymoquinone.

Benzsne-aze-thymel-sulphonic acid

C₅H₅-N₂-C₅H(CH₃)(C₅H₁)(HSO₃)OH. [216°]. Small yollow prisms. Yellow colouring matter. Prepared by the action of diazobenzens chlorido on a ealt of thymol-sulphonic acid. Salts .-A'Na: small yellow crystals.- A',Ba: fine hair-

like nesdles (Stebbins, B. 14, 2793).

Benzens, p-azo-tolnene C₆H₃.—N₂—C₆H₄(CH₃) [63° corr.]. Formsd by diazotising amido-bancorr.]. Formed by discouring announces receive p-azo-toluens $C_4H_4(CH_2)-N_2-C_4H_4(NH_2)$ dissolved in alcohol and boiling the solution (Schultz, B. 17, 466). Oranga-red platss; v. sol. alcohol. Volatils with steam. By treatment with alcoholic SnCl₂ and H₂SO₄ it is converted into a bass melting at [116°].

Bensens-axo-m-tolylens diamine C.H. -N. -- C.H.Me(NH.) .- Yellow needles. Easily soluble in alcohol, sparingly in water. Prepared by the action of diazobenzene chloride on (1, 2, 4) tolylene-diamine, [99°].—B'HCl: orango red needlee (Stebbins, jun., A. C. J. 1, 465; B. 13, 717; C. N. 41, 117).

Benzene - azo - xylenol C.H.N.C.H.Mo.OH [1:3:5:2]. [175°]. Formed by combining diazobenzene chlorido with m-xylenol CoH, Me. (OH) [1:3:4] (Grevingk, B. 19, 148). Slender brownishred needles. V. sol. alcohol, ether, and benzene, incol. water. On reduction it yields aniline and o-amido-m-xylenol $C_0H_2Me_2(NH_2)(OH)$ [5:3:2:1]. Bromo-amido-benzene-azo-p-bromo-aniline

Diacetyl derivative [2:5:1] NHAc.C,H3Br -N2-C,H3Br.NHAc[1:5:2] [282°]. Formed by warming acetyl-bromo-nitroaniline C. H. (NHAe) Br(NO.) [2:5:1] with zine and conc. NH, Aq (C. H. Matthiessen a. Mixter, Am. 8, 847). Pale red substance.

p-Bromo-benzene-azo-benzene-p-snlphonic acid [4:1] $C_aH_aBr - N_a - C_aH_a(SO_aH)$ [1:4]. Formed by sulphonation of benzene-p-azo-bromobenzene or by bromination of benzene-azo-benzene-p-sulphonic acid. Flat needlee (containing 3aq).

Salte. - KA': rhombio tablee. - NaA': yellow eilky needlee, sl. sol. water (Janoveky, M.

5, 162; B. 20, 358; M. 8, 53).

m - Bromo - benzene-azo-benzene - p-enlphonio acid [3:1] C₀H₄Br—N₂—C₆H₄(SO₃H) [1:4]. Formed by sulphonation of benzene-m-azo-bromobenzene with furning sulphuric acid. Glietening golden plates (containing 1, aq).

Salts.—KA': yellow pearly pp. of mioroscopio needles.—NaA': pp. v. el. sol. water (Janovsky a. Erb, B. 20, 359).

o-Bremo-benzene-o-azo-bremo-benzene [2:1] C₆H₁Br-N₂-C₆H₄Br [1:2]. [185°]. A product of the bromination of benzene-azobenzene in HOAc (Janovsky, M. 8, 50; B. 20, 337). Golden platee, el. sol. alcohol. On nitration it gives a tri-nitro- derivative [135°].

p-Bromo-benzene-p-azo-bromo-benzene [4:1] $C_eH_aBr-N_2-C_eH_aBr$ [1:4]. [205°]. Formed by bromination of benzeno-azo-benzene (Werigo, A. 135, 178; 165, 189). Formed also by reduction of p-bromo-nitro-benzene with zino-dust and alcoholic KOH (Schultz, B. 17, 465). Yellow needles. By alcoholic SnCl₂ and H₂SO, it is converted into a di-bromo-di-amidodiphenyl. Funing H.SO, forme a sulphonic acid C₁₂H,Br,N.SO₃H 3aq (W.).

m.Brome-beazene-m-azo-bromo-benzene [3:1] C_eH_eBr-N₂-C_eH_eBr [1:3]. [126°]. From the corresponding hydrazo-compound by Fe₂Cl_e

(Gabriel, B. 9, 1407).

Tribromo-benzene-azo-dimethyl-aniline C.H.Br. N. C.H. (NMe.). [161°]. Formed by adding an alcoholic colution of dimethylaniline (2 mol.) to C. H. Br. N. NO. (1 mol.). Crystalline pp.

Red plates (from glacial acetic acid). Insol. water, hardly soluble in alcohol. It combines with cono. HCl (Silberstein, J. pr. [2] 27, 124).

Tribromo - benzene - azo - methyl - diphenylamine C₂H₂Br₂-N₂-C₆H₄NPhMe. [138°]. From C₆H₂Br₂N₂NO₃ and NPh₂Me in alcohol. Small brownieh-red plates (from glacial acetio scid). Ineol. water, sl. sol. alcohol. Does not combine with HCl (Silberetein, J. pr. [2] 27, 125).

Brome-bensene-aso-(a)-naphthol [4:1] $C_2H_1Br-N_2-C_{10}H_2OH$ [1:4]. [196°]. From p-diazo-brome-benzene and (a)-naphthol, or by bromination of benzeno-azo-(a)-naphthol (Mazzara, G. 14, 271).

p-Bromo - benzene - azo - (3) - naphthol [4:1] C_aH₄Br—N₂—C₁₀H_aOH [1:2]. [161°]. From p-diazo-bromo-benzene and (8)-naphthol, or from benzeno-azo-(\$)-naphthol and bromine (Mazzara, G. 13, 438). Orango needles.

p-Bromo-benzone-p-azo-nitro-benzene [4:1] C₆H₄Br - N₂ ··· C₆H₄(NO₂) [1:4]. [108°]. Formed by nitration of benzene-p-azo-bromo benzene. Yellow needles, V. eol. aloohol V. eol. aloohol

(Janovsky a. Erb, B. 20, 358).

Bromo-bonzene-azo-di-nitro-benzene (?) [4:1] $C_0H_1Br - N_2 - C_0H_2(NO_2)_2$ [1:2:4] (?). [190°]. Formed by nitrating benzene-p-azo-bromo-benzeno (Janovsky, M. 8, 52).

p-Bromo-benzene-azo-nitro-ethane [4:1] C₆H₁Br -N₂--CH(NO₂).CH₃. [o. 187°]. From p-diazo-bromo-benzene nitrate and potaseium nitro-ethane (Wald, B. 9, 393). Brick-red crystals (from diluto alcohol); sol. ether, glacial HOAc, and chloroform. Salt.—Call.KNaOaBr.

Di-bremo - di - imido-di-hydro-anthracene-azodi bremo-di-imido-di-hydro-anthracene

 $C_{14}H_7Br_2N_3$ or NII

$$\begin{array}{ccc} NII & NH \\ C_{\bullet}H_{\bullet} \overset{C}{\underset{C}{\bigcirc}} C_{\circ}Br_{2}H - N_{2} - Br_{2}HC_{\bullet} \overset{C}{\underset{C}{\bigcirc}} C_{\circ}H_{\bullet} \end{array} (?)$$

$$NII & NH \\ NH & NH \end{array}$$

[233°]. Prepared by heating dibromo - nitroantbraquinone with alcoholic NH3 (Claus a. Diernfellner, B. 14, 1335). Red needles. Sublimable. Si, sol. alcohol and ether, incol. water or aqueoue acide and alkalis.

Bromo - di - oxy - benzene - azo - bromohydroquinone. Tetra-methyl derivative C.H.Br(OMe), -N. -C.H.Br(OMe), [220°]. Formed by bromination of the tetra-methyl derivative of di-oxy-benzeno-azo-hydroquinone (Baessler, B. 17, 2125). Red orystalline solid, v. sol. benzene, chloroform, and CS2, v. sl. sol. alcohol, insol. water.

Bromo-sulpho-benzene-azo-benzene sulpho-[6:3:1]C,H,Br(SO,H)-N,-C,H,Br(SO,H)[1:6:8]. Formed by oxidation of a neutral aqueous solution of bromo-amido benzene eulphonic acid. C_all₃Br(NH₂)(SO₃H) [4:3:1] with KMnO₄. Salte.—K₂A" 2aq: glistening red tables (Limpriebt, B. 18, 1422).

Di-bromo-sulpho-benzene-azo-di-bromo-benzene salphonic acid $\begin{array}{lll} \hbox{$^{\prime}$}2:6:4:1]\mathring{C}_{s}H_{s}Br_{s}(SO_{s}H).N_{s}.C_{s}H_{s}Br_{s}SO_{s}H[1:2:6:4].\\ From & potassic & di-1 \\ romo & o & amido & benzene \end{array}$ eulphonate and KMnO, (Rodatz, A. 215, 222). Red plates (containing 2aq). V. sol. water or alcohol. Reduced by SnCl₂ to the original

BaA" 3aq. — CaA" 4aq. — PbA".

Chloride. — [258° 262°]. Brown platos.

Amide .- Violet silky needles. Di - bromo - snlpho - benzene - azo - di - bromebenzene sulphonio acid [4:6:3:1]C₆H₂Br₂(SO₃H).N₂.C₆H₂Br₂SO₃H[1:4:6:8]. From potaesie di-bromo-amido-benzene enlphonate and KMnO₄ (Rodatz, A. 215, 216). Slender red needlee, containing 1 aq (from water). V. sol. water, v. c. sol. alcohol. Reduced by SnCh to the original CaH2(NH2)Brg. SO3H. Saltz .-K.A"3aq.—BaA" aq.—CaA" 4aq.—PbA" 2}aq. Chloride.—[233°]. Brick-red needles.

Amide.-Microscopic orange needles (from

alcohol). Does not melt.

An acid isomeric with the above may be got by the action of KMnO, on the tetra-bromo-Lydrazo-benzene di-sulphonic acid of Jordan, A. 202, 361.

Tri - bromo - snlpho - benzene - azo-tri - bromobenzeae snlphonic acid [2:4:6:3:1] $C_aHBr_a(SO_aH) - N_a - C_aHBr_a(SO_aH)$ [1:2:4:6:3]. From potassic tri-bromo-m-amido-bonzoate by KMnO, (Rodatz, A. 215, 225). Flat orange necdles. Reduced by SnCl, to the original C.H(NH2)Br3.SO3H.

K₂A" 3aq.—BaA" 2aq.—CaA" 7aq.—PbA" 4aq. Chloride. - [222'-224°]. Dark violet tables. Amide .- Brown crystals that do not melt. Di-bromo-snipho-bonzene-azo-(β)-naphthol

C₆H₂Br₂(SO₃H) · N₂ - C₁₀H₂OH (Stebbins, C. N. 42, 44; A. C. J. 2, 236). From diazo-dibromobenzene sulphonio acid and an alkalino solution

of (β)-naphthol,
Di-bromo-snlpho-benzene-azo-di-oxy-naphthalene $C_0H_2Br_2(SO_3H)-N_2-O_{10}H_3(OH)_2$. From diazo-dibromo-benzene sulphonic acid and an alkaline solution of dioxynaphthalenc. Needles (Griess, B. 11, 2199).

Bromo - snlpho - tolnene - azo - bromo - tolnene snlpbonic acid N₂(C₆H₂BrMe.SO₃H)₂ [1:?:4:5] From potassic bronio-p-toluidine sulphonate (of Jenssen) and KMnO (Kornatzki, A. 221, 186). Red tables with pointed ends. Salts. -"KA" 4aq.—BAN" 5aq.—CaA" 4½aq.—PhA" 5aq.
Chloride.—[226°]. Groups of red prisms.
Amide.—[above 260°]: red powder.

Di - bromo - snlpbo - toluene - azo - di - bromotolneno sniphonio aoid N₂(C₆Hhr₂Me.SO₂H)₂[1:2:7:2:5]. From potassic dibromo-o-toluidine eulphonate (of Hayduck) and KMnO₄ (Kornatzki, A. 221, 188). Blood-red plates. Salts.—

K_A"2aq.—BaA"9aq.—CaA"8aq.—PbA"9aq.

Chloride. [243°]. Red swallow-tailed plates.

Amide.—[218°]. Powder.

 $\mathbf{D}\mathbf{i}$ - $\mathbf{b}\mathbf{n}\mathbf{t}\mathbf{y}\mathbf{l}$ - p - $\mathbf{a}\mathbf{m}\mathbf{i}\mathbf{d}\mathbf{o}$ - $\mathbf{b}\mathbf{e}\mathbf{n}\mathbf{z}\mathbf{e}\mathbf{n}\mathbf{e}$ - $\mathbf{a}\mathbf{z}\mathbf{o}$ - $\mathbf{d}\mathbf{i}$ - $\mathbf{b}\mathbf{n}\mathbf{t}\mathbf{y}\mathbf{l}$ anillne (C₄H₉) N.C₈H₄ - N₂ - C_bH₄N(C₄H₉)₂. Dibutyl-aniline-azyline. [158°]. Red needles. butyl-antline-azyline. [158°]. Red needles. Formed by passing NO through an alcoholic solution of di-butyl-auiline.

Periodide. -B" Is: dark crystals with blue reflection (Lippmann a. Fleissner, B. 15, 2142

and B. 16, 1421; M. 3, 713).

m-Carboxy benzene, azo aceto acetic acid $C_{e}H_{4}(CO_{2}H)$ N₂ - CHAc.CO₂H. Formed by the action of the sulphate of m-diazo-benzoic acid upon acetoacetio ether in alkaline aqueous solution (Gricss, B. 18, 962). Small yellow plates or needles. Soluble in alcohol, nearly insoluble in water. Bitter taste.

o - Carboxy - benzene - o - azo - benzoic aold [2:1] C₈H₄(CO₂H)—N₂—C₈H₄.CO₂H [1:2]. o-Azo-benzoic acid. [238]. Mol. w. 270. o-nitro-benzoic acid and sodium-amalgam (Griess, B. 10, 1868). Dark yellow hair-like needles.

Insol. water; m. sol. alcohol. Salts.—BaA"7aq.—BaA"9aq.—Ag.A". Ethyl ether Et.A". [139°]. From o-nitrobenzoic ether by sodium-amalgam. Scarlet needles (Fittica, J. pr. [2] 17, 216).

m-Carboxy-benzene-m-aso-benzoic acid N₂(O,H₄,CO₂H₃, [1:8]. *m-Asobensoic acid.* From *m*-nitro-benzoio acid by sodium-amalgam (Streeker, A. 129, 134). Amorphous powder, decomposed by heat, v. sl. sol. water, alcohol, Gives phenazine when distilled and ethor. with lime. Its copper salt gives azobenzene on distillation. HgO and iodine gives an amorphous di-iodc derivative, C₁,H₈L₂N₂O₄ (Benedikt B. 8, 386). Salts.—BaA" 5aq.—Ag₂A".

Ethyl ether Et.A". [99°]. By the action of

EtI on Ag2A" Golubeff (B. 7, 1651) obtained two bodies isomeric with this ether, one melting at 76°, the other being a monobasic acid.

p-Carboxy benzene-p-azo-benzoio acid

 $N_2(C_6H_1.CO_2H)_2$ [1:4]. From p-nitro-benzoic acid and sodium-amalganı (Beilstein a. Reichenbach, A. 129, 144; Billinger, A. 135, 154). Formed, together with azoxybenzoic acid, by boiling nitro-benzil with alcoholic KOH (Zinin, Z. 1868, 563). Flesh-coloured amorphous powder, insol. water, alcohol, and ether. Gives phenazine when distilled with lime. Salts.—(NH₄)₂A" aq.—
Na₂A".—CaA" 3aq.—BaA".—Ag₂A".

Ethyl ether Et₂A". [88°] (Fittica,

J. pr. [2] 17, 216). m-Carboxy-benzene-azo-malonio acid

C.H.(CO.H)-N.-CH(CO.H). Formed by the action of the nitrate of m-diazo-benzoic acid on malonio ether in alkaline aqueous solution (Griess, B. 18, 962). Microscopic orange needles or plates. Easily soluble in alcohol.

Carboxy - benzene - azo - di-methyl-amido-bensoio aoid C₆H₄(CO₂H). N₂. C₆H₃NMe₂. CO₂H. From m-diazo-benzoic acid and dimethyl-m-amidobenzoic acid (Griess, B. 10, 525). Brown pp.

m-Carboxy-benzene-azo-nitro-methane C.H. (CO.H).N. CH. (NO.). Formed by adding the nitrate of m-diazobenzoic acid to a dilute aqueous alkaline selution of nitromethane (Griess, B. 18, 961). Yellowish-red plates. Sol. hot alcohol and ether, v. sl. sol. water.

m-Carboxy-benzene-azo- (β) -naphtbol $C_aH_4(CO_2H)$ - N_2 - $C_{10}H_4(OH)$. [235°]. Prepared by the action of m-diazo-benzoic acid on an alkalino solution of (B)-naphthol (Griess, B. 14, 2035). Reddish-yellow needles or plates. Sol. hot alcohol, sl. sol. cold alcohol and ether, insol.

Salts: A'2Ba31 aq: red microscopic needles. A'K 2aq: easily soluble yellow needles or plates. Ethyl ether A'Et. [104°]. Yellowish-red needles or plates. Sol. ethor, insol. water.

Amide.—Slender orange needles. Sl. sol. alcohol and ether, insol. water.

m-Carboxy-benzene-azo- (β) -naphtbol enlphonio acid C₆H₁(CO_.H) - N₂ - C₁₆H₂(OH)(HSO₃). Prepared by the action of m-diazo-henzoic acid on an alkalino solution of (β) naphthol sulphonic aoid (Griess, B. 14, 2036). Brown needles or plates. 'Sol, hot water, sl. sol, cold water and alcohol, insol. ether. Dyes wool and silk a splendid orange.—A"2H_Ba4aq: orange pp. of slender necdles.

m-Carboxy-benzene-azo-(3) naphthol-di-sulphonio acid C₀H₄(CO₂H).N₂·C₁₀H₄(OII)(HSO₂)₂. Prepared by the action of m-diazobenzoic acid on (β)-naphthol-(α)-disulphonic acid in alkaline solution (Griess, B. 14, 2037). Yellowish-red microscopic needles. Dyes silk and wool an

orange scarlet.

Salts.-BaHA" 6aq: red crystalline pp.-A.M., 12aq : red microscopic needlee. Carboxy-benzene-azo-phenol v. Oxy-benzenesso-benzoic acid.

Di-carboxy-benzene-azo-phthalio acid

C₆H₃(CO₂H)₂-N₂-C₆H₃(CO₂H)₂. Azo-phthalic acid. [about 250°]. Prepared by reduction of nitro-pbthalio acid with sodium-amalgam (H. Müller; Claue a. May, B. 14, 1330). Small yellow needles: sl. col. water, alcohol, and ether.

Salta.—A'Na,10aq: yellow monoclinic prisms, v. sol. water.—A'K,6aq: long yellow needles.—A''Mg,18aq: large orango orystals.— A''Ag,: yellow insolublo pp.-A''Ba,: yellow incoluble pp.

Di-carboxy-benzene-azo-tere-phthalic acid [5:2:I] $C_6 ll_3 (CO_2 H)_2 - N_2 - C_6 ll_3 (CO_2 H)_2$ [1:5:2].

4 so-terephthalic acid.

Formation.-1. By oxidation of hydrazoerephthalic acid with nitrons acid .- 2. By eduction of nitro-terephthalio acid with codinmımalgaın (Homolka a. Löw, B. 19, 1092).

Yellow needles. Sol. alcohol and ether, sl. iol. water. Decomposes above 200°.

Carboxy-naphthaleno-azo-(8)-naphthoic acid $C_{10}H_6(CO_2H) - N_2 - C_{10}H_6(CO_2H)$. From a nitro- β)-naphthoic acid by ammonium sulphide Rakowsky, B. 5, 1022).

exo-Carboxy-tolucne-azo-phenyl-acetic acid 30,H.CH., Call, -N., -C. II., CH., CO., II. S. (cold sleehol) 0.375; (hot alcohol) 1.57. Insol. hot vater, ether, and benzene. Does not melt below 100°. Formed by the action of sodium amalgam m nitro-phenyl-acetic acid [I51] (Wittenberg, 3l. [2] 43, 111).

Carboxy - tolnene - azo - toluic acid

5:2:1] C_all_aMe(CO_aH)—N₂—C_all_aMeCO_all[I:5:2]. 4zo-p-toluic acid. [184]. From nitro-p-toluic icid by sodium amalgam (Fittica, B. 7, 1358). linute yellow needles; m. sol. boiling water, ', sol, alcohol,

p.Chloro-benzene-azo-benzene p-sulphonic $.cid [4;1]C_sH_1Cl-N_2-C_sH_1(SO_3H) [1;4] [148^{\circ}].$ formed by warming p-chloro-azo-henzene with uming sulpburie acid (10 p.c. SO₃) at 60°-70°. Brown needles; v. e. sol. water and alcohol. It s reduced by SnCl2 to p-chloro-aniline and p-

ulphanilic acid.

Salts.-A'Na: largo orange-yellow pearly plates or small needles; sl. sol. cold water. Y2Ba: glistening flesh-coloured needles .- Tho K, Mg, Ca, Ag, Cr, and Fe, salts are white to lark-yellow needles; the Cu salt forms green plates; all are sparingly soluble.

Chloride C₆H₄Cl.N_c,C₆H₄,SO₂Cl: [130°]; distening red prisms, easily soluble in alcohol

ind ether.

 $C_611_4Cl.N_2.C_611_4.SO_2NH_2: [211°];$ Amide rellowish - brown prisms; sol. hot alcohol, paringly in other and cold alcohol, insol. water Mentha a. Heumann, B. 19, 2972).

m-Chloro - benzene - m-azo - chloro - benzene 1:3] C₀H₁Cl-N₂-C₀H₁Cl [1:3]. Azo-chloro-benzene [101°]. Obtained by acting with Fe₂Cl₆ on m-di-chloro-hydrazo benzene in alcoholic solution (Laubenheimer, B. 8, 1025). Orange needles (from alcohol).

p.Chloro-benzene-p.azo-chloro-benzene
4:1] C_sH_sCl-N₂-C_sI_sCl [I.4], [181°]. From pshloro-benzene - p - azo - nitro-benzeno and
alcoholic potaeh (Willgerodt, B. 15, 1002); or

| Cuminic-azo-cuminio acid | PrC_sH_sCO₂H_sPr.CO₂H_s

from di-chloro-azoxybenzene and fuming H,SO,

(Heumann, B. 5, 913, 918). Yellow needlee, p-Chloro-benzene-azo-ohloro-benzene sulphonic acid [4:1] C₆H₄Cl.N₂.C₆H₃Cl.SO₂H [1:4:?]. Prepared by sulphonation of the preceding body (Calm a. Heumann, B. 13, 1183; 15, 2558). Slender reddish-yellow needles. Sol. water and alcohol. Salts .- A'Na: golden plates, sl. sol. cold water. - A'K: reddish-yollow glistening plates, sol. hot water, and alcohol. - A'Ag.-A'Ba: yellow crystalline pp.-A'Ca: golden yellow plates.—A'₂Pb: orange glistening plates. Chloride C₁₂H,Cl₂N₂SO₂Cl: [161°]. Long

orange-red needles.

Chloro - benzene - azo - chloro - phenol

[3:1] Cl.C, H₄ - N₂ - C, ll₃Cl(Ol1) [1:3; ?]. [115°] Formed by the action of fuming H₂SO₄ on m-di-chloro-azoxybenzeno (Schultz, B. 17, 465). Brown plates.

Chloro-benzene-azo-ohloro-nitro-benzene

[4:1] C₀11,Cl-N₂-C₀11₃Cl(NO₂) [1:4:?]. [210°]. Prepared by reducing di-chloro-nitro-azoxyhenzene with cold alcoholic NH,HS (Calm a. Henmann, B. 13, 1184). Yellow needles. Sl. sol, alcohol.

m - Chloro - benzene - azo - di - methyl - aniline [3:1] C₀H₄Cl-N₂-C₆H₄NMe₂[1:4]. [98°]. Yellow plates. Tolerably soluble in alcohol. Prepared by adding sodium nitrite (1 mol.) to a solution of m-chloranilino (1 mol.) and dimethylaniline (I mol.) in diluto H.SO, (Staedel a. Bauer, B. 19, 1955).

p-Chloro-benzene-p-azo-nltro-benzene

[I:4] C₆H₄Cl.N₂.C₆H₄(NO₂) [1:4]. [193°]. Formed by nitration of p-chloro-azo-benzene with funing HNO3. Yellow needles; v. sol. acctic acid and hot alcohol, sl. sol, cold alcohol, insol, water. It is reduced by SuCl2 to p-chloro-aniline and pphenylene diamine (Mentha a. lleumann, B. I9, 2971).

p-Chloro-benzene-azo-phenol C_c 11 $Cl-N_2-C_c$ H $_1(OH)$. [1522]. Formed by gently warming p-chloro-diazo-benzene-ohloroanilido with phenol, p-chloranilino being elimi-

nated. Requisity of the committee of the Azo-cumene. [108°]. From nitro cinnene, [-35°], by sodium-amalgam (Pospekhoff, J. R. 1886, 49). Thin yellow leastlets, sl. sol. cold alcohol.

4-Cumene-azo-4-cumenol

[2:4:5:1] C₆H₂(CH₃)₃.N₂.C₆H(CH₃)₃.OH [I:3:5:6:2]. [148°]. Formed by combining di-azo-oumene ehloride (from ψ-cumidine) with ψ-cumenol [70°] (Liebermann a. Kostanoki, B. 17,885). Orange needles. Insol alkalis. Dissolves in H.SO, with an orange colour. No nitrogen is evolved on boiling with HCl.

ψ-Cumene-azo-ψ-cumidlne

[2:4:5:1] C_aH₂Me₃,N_a,C_aHMe₃,NH₂ [1:2:4:5:6], [139°]. Prepared by the action of cuid of hydrochloride upon diazo-cumeno-cumide (diazo-amido-cumene) dissolved in eumidino (Nölting a. Baumann, B. 18, 1147; Bl. [2] 42, 335). Orango plates (from alcohol). V. sol. ethor. On reduction with SnCl, it yields oumidine and eumylenc-o-diamine. Salt .- B'HCl: yellow

nitro - cuminic acid and sodium - amalgam (Alexéeff, Bl. [2] 88, 552; 42, 821; J. R. 1882, 198; Alexéeff a. Kissel, Bl. [2] 40, 72).

Ethers.—Me₃A".[166²].—Et₃A".[104²-108²].

Cymene-azo-cymene

C₆H₃MePr-N₂-C₆H₃MePr. [86°]. From nitro-cymeno by sodium amalgam (Werigo, Z. 1864, 721). Cherry-coloured plates.

Ethyl - amido - benzene - azo - bsnzene - p entlybonic acid [4:1] C_all₄(SO₃II),N_xC_aH_yNI|Et [1:4]. p. Sulpho. beazene azo ethyl-aniline. [c. 244°]. Obtained by combining p-diazobenzene-sulphonic acid with ethyl-aniline in combination. Standblan profiles. Nearly used acid solution. Steel-blue needles. Nearly insol. alcohol and cold water, sl. sol. hot water. Tho sodium salt (NaA') forms orange-red plates. On reduction with (NH1)2S it yields mono-ethyl-pphonylene-diamine and p-sulphanilic acid (Bornthson a. Goske, B. 20, 929).

Di-ethyl-p-amide-benzene-azo-di-ethylaniline C₆H₄(NFt₂) - N₂ - C₆H₄(NEt₂). Diethylaniline azyline. [170°]. Prepared by passing NO through an alcoholic solution of di-ethylanilino; yield 50 p.c. of the theoretical. Red monoclinic crystals, a:b:c=1:.7108:.9493. $\beta=$ 90° 30'. V. sel. chloroform, sl. sol. cold alcohol.

Reactions .- 1. Nitrous acid acting on the acotic solution gives p-nitro-di-ethyl-aniline.-2. On reduction it yields u-di-ethyl-p-phenylene-diamine.—3. Heated with ethyl iodide at 100° it gives tetra-ethyl-p-phenylene-diamine.-4. Mcl at 100° gives di-methyl-di-ethyl-p-phenylenediamine.

Salts .- B"H2Cl2PtCl4: small brownish-red trimetric tables.—I'errocyanide B"2H4FeCy8: brown plates.—Pierato B"(C8H4(NO2)30H)2: yellow sparingly solublo needles (Lippmann a. Fleissner, B. 15, 2136; 16, 1415; M. 3, 286, 788).

Di-ethyl-amido-benzene-azo-tolnidine Acetyl derivative

[4:1] C₆H₄(NEt₂)-N₂-C₆H₂Me(NHAc) [1:6:3]. [159]. From diazotieed acetyl-tolylene-diamine C₄H₄(NH₄)Me(NHAo) [1:6:3] and di-othyl-aniline (Wallach, A. 234, 359).

o-Ethyl-benzsne-o-azc-ethyl-benzene $[2:1]C_bH_4(C_2H_5)-N_2-C_bH_4(C_2H_5)[1:2].[47^{\circ}corr.].$ Formed by reduction of o-nitro-ethyl-benzene with zinc dust and alcoholic NaOH (Schultz, B. 17, 473). Long red dimetric prisms, a:c = 1: 3455. V. sol. atcohol. By treatment with SuCl. and HCl in alcoholic solution it yields a di-amido-diethyl-diphenyl.

p. Ethyl-benzene-p-azo-ethyl-benzene [4:1]C₆H₄(C₂H₃)=N₂-C₆H₄(C₂H₃)[1:4].[63°corr.]. (above 340°). Formed by reduction of p-nitro-ethyl-benzene with zinc-dust and alcoholic No.01 (Schultz B. 17.475°). NaOH (Schultz, B. 17, 475). Orange-red plates or thick prisms. V. sol. alcohol. By treatment with SnCl₂ and H.SO₄ in alcoholic solution it yields a di-amide-di-ethyl-diphenyl.

p-Ethyl phenyl amide-benzene-p-azc ethyldl-phenylamine

[4:1] C.H.NEtl'h-N.-C.H.NEtPh [1:4]. [178°]. From ethyl-di-phenyl-amino and NO (Lippmann

a. Fleissner, M. 4, 706). Monoelinic red crystals.
Ethyl-pyrrol-azo-(β)-naphthalene
C₁₈H₄N.C.(H₃NEt. [74°]. Obtained by adding (B)-diazo-naphthalene chloride (1 mol.) to ethylpyrrol (1 mol.) dissolved in alcohol containing godium acetate. Thick red tables. Sparingly

soluble in dilute HCl. Dissolves in conc. H.SO. with a dark reddish-yellow colour. The platino-chloride forms small sparingly soluble red needles (O. Fischer a. Hepp, B. 19, 2258).

Ethyl-pyrrol-azc-p-tolnene C.H.Me.N.C-CH C.H.Me.N.C.H.NEt probably NEtl

HC = CH Formed by adding p-diazo-tolnene ehloride (1 mol.) to an alcoholic solution of cthyl pyrrol (1 mol.) containing sodium acetate. Thick red prisms. Dissolves in conc. H.SO, with a yollow colour, in diluto HCl with a reddish yellow colour. The platino-chloride forms sparingly coluble red needles (O. Fischer a. Hepp, B. 19, 2257).

Iodo-carboxy-benzone-azo-iodo-benzcic acid C.113I(CO211) - N2 - C.113I(CO2H). Azo-iodo-benzoic acid. From m-amido-benzoie acid, iodine, and HgO (Benedikt, B. 8, 386).

Mesitylene-azo-mesitylene C.H. (CH.) - N. - C.H. (CH.). Aso mesitylene. [75° corr.]. Prepared by oxidising an aqueous solution of mesidino hydrochloride (5 pts.) with a solution of 45 pts. of potassium ferricyanide and 10 pts. of KOH (Schultz, B. 17, 476). Thin red needles. Sol. hot alcohol. It does not appear to yield a hexa-methyl-benzidine by treatment with SnCl₂ and HCl in alcoholic

Methyl - amide - benzene - azc - benzeae - snl $phonio\,acid[4:1]\,C_{_{6}}H_{_{4}}(SO_{_{3}}H).N_{_{2}}.C_{_{6}}H_{_{4}}.NHMe[1:4].$ p-Sulpho-benzene-azo-methyl-aniline. Obtained by combining p-diazo-benzene sulphonic acid with methyl-aniline in acid solution; yield 30 p.e. of theoretical. Also formed by the action of cold dilute acids upon p-sulpho-diazo-benzenomethyl-anilide C₆H₄(SO₂H).N_{...}NMcC₆H₅. Steol blue needles, al. sol. water. The sodium-salt (A'Na) forms largo orange-red plates, v. sol. hot water. On reduction with (NII,) S it yields psulphanilie acid and mono-methyl-p-phenylene diamine (Bernthsen a. Goskc, B. 20, 925).

Di-metbyl-amidc-benzene-azo-benzene-snlphonic acid [4:1] CoH, (NMc.) - N2-CoH, SO2H [1:4] Helianthin, or Orange III.

Preparation .- 1. From dimethylaniline and p-diazobenzeno sulphonic acid (Griess, B. 10, 525). -2. Dimethylamido-azobenzene (1 pt.) ie dissolved in 20 pts. cold sulphuric acid of 30 p.c. anhydrido value and allowed to stand for 24 hours (Möhlau, B. 17, 1491). The absorption spectrum has been mapped by Hartloy (C. J. 51, 192).

Di-met hyl-amido-benzene-azo-benzoic acid [4:1] C₆H₄NMo₂-N₂-C₆H₄,CO₂H [1:3]. From di-methyl-aniline and m-diazo-benzoic acid (Griess, B. 10, 525).

Di - methyl - amido - benzene - azc-di-methylaniline [1:4] C,H,(NMe2)-N2-C,H,NMc2 [1:4] [266]. Dimethylaniline-azylinc. Rcd necdles.

Formation.-1. By diazotising p-amido-dimethyl-aniline and combining the diazo-compound C_sH₄(NMe_s),N_s,Cl with dimethylaniline (Nölting, B: 18, 1143),—2. By passing NO through an alcoholic solution of di-methylanilino for several days.

Reactions .- 1. Nitrous acid acting on the aeotic seid solution gives p-nitro-di-methylaniline.—2. On reduction it gives u-di-methyl-p. phenylene-diamine.

Balts.—B"H,Cl.PtCl.: dichroic crystalline powder.—Picrate B"C,H,(NO,),OH+C,H,OH: glistening green needles (Lippmann a. Fleissner, B. 15, 2136; 16, 1415; M. 3, 708).

Di-methyl-amido-benzene-azo-p-tolaene C,H,—N₂—C_eH₄.NMe₂. [168°]. Golden platoe. Easily soluble in alcohol and ether.

Preparation. -- A solution of 65 pts. of NaNO2 (100 p.c.) and 35 pts. NaOII in 465 pts. of water is slowly added to a cooled solution of 100 pts. of p-toluidine, II3 pts. of dimethylaniline and 200 pts. of HCl in 300 pts. of water

On reduction it yields p-toluiding and di-methyl-p-phenylene-diamine. The hydrochloride and sulphate form violet prisms, giving a red

solution in alcohol (Möhlau, B. 17, 1492).

Di-methyl-amido-bsazene-azo-p-tolnene-sul-phonic acid C,H_e(HSO₃).N₂.C_eH_e,NMe₂. Formed by the combination of p-diazo-toluene sulphouic acid (Me:IISO₃:N₂=1:3:4) with dimethylaniline (Möhlau, B. 17, 1493). Dark-violet prisms. Soluble in water and alcohol with an orange oolour, insoluble in ether. The sodium salt forms orange glistoning plates.

Di-methyl-amido-benzene-azo-telnidine

[4:1] $C_sH_4(NMo_s)-N_2-C_eH_4Me(NH_2)$ [1:6:3] [145°]. From its acetyl derivative [200°] which is formed by the action of diazotised-acetyltolylene-diamine C_sH₃(NH₂)Me(NHAo) [1:6:3] upon di-methyl-anilino (Wallach, A. 234, 355).

Di-methyl-amido-benzene-azo-telaidine [4:1] $C_eH_1(NMc_s)-N_2-C_eH_2Me(NH_s)$ [1:4:3] [215°]. From its acetyl derivative, [192°] which is formed by the action of diazotised acctyl-; tolylene-diamine C_aH_a(NH_a)Me(NHAc) [1:4:3] upon di-methyl-aniline (Wallach, A. 234, 359).

p-Methyl-phenyl-amido-benzene-azo-methyl-

di-phenyl-amine

[4:1] C.II.,NMoPh.N.,.C.H.,NMoPh [1:4] [150°]. Yellow crystals, got by the action of NO on mothyl-diphenyl-amine (Lippmann a. Fleissner, M. 4, 798).

(a)-Naphthalene-(a)-azo-naphthalene

[a] $C_{10}H_1$, N_2 , $C_{10}H_7$, [a], [190°]. Prepared by boiling diazo-naphthalene azo-naphthalene with alcobol: 1 pt. of naphthalcno-azo-naphthylamino is dissolved in 100 pts. of 95 p.c. alcohol and 5 pts. of II SO, are added; the still warm solution is then treated with a saturated solution of (1 mol. of) NaNO2; the fluid is heated and finally boiled for a few hours, and the azonaphthalene precipitated by water. It is recrystallised by dissolving it in hot aniline and adding alcohol (Nietzki a. Goll, B. 18, 298, 3252). Steel-blue orystals. Sublimes in thin yellow plates. Soluble in aniline, sparingly in alcohol. It dissolves in cold H2SO, with a blue colour, but on heating the solution to about 180° it becomes violet and exhibits a rod fluorescence. By alcoholic NH, HS, or zinc-dust and alcoholic KOH, it is reduced to hydrazonaphthalene.

(α)-naphthalene-(β)-azo-naphthalene

[a]C₁₀H₁N₂C₁₀H₁[β]. [186°]. Obtained by diazetisation of (β)-naphthalene-p-azo-(α)-naphthylamine and beiling with alcohel. Dark-brown platos with steel-blue reflex. Sol. alcohol, acetic acid, &c. Dissolves in conc. H.SO, with a violet colour (Nietzki a. Göttig, B. 20, 612). Laurent's naphthase [275°], got by heating nitro-naphthalene with zine dust, or (a) naphthylamine with PhO, has been regarded as naphtha-

lene-azo-naphthalene, but Witt has lately (B. 19, 2794) shown it to be an azine U, H, N, v. (aB)-NAPHTHAZINE (Laurent, A. 109, 384; Doer, B. 3, 291; Alexeoff, B. 3, 868; Schichuzky, B. 5, 365; Klobulowsky, B. 10, 570, 772, 873).

Naphthalene-p-azo-(α)-naphthol

C10H2-N2-C10H0OH. Formed by adding diazo-naphthalene chloride to a solution of (a)naphthol in NaOlIAq; it is ppd. by HCl. Crimson powder. Soluble in alkalis forming crimsou solutions (P. F. Frankland, C. J. 37,

Naphthalene-o-azo-(β)-naphthol

C10H7-N2-C10H6OH. [176°]. Formed by diszotising (β)-amido-azo-naphthalene and heating the diazo- compound with water (Nietzki a. Goll, B. 19, 1282). Sublimos in glistening golden needles

(a)-Naphthalene-(a)-aze-(a)-naphthylamine $C_{10}H_1$ - N_2 - $C_{10}H_0$ NH₂. [180°]. Prepared by adding KNO₂ (1 mol.) to a dilute solution of (a)naphthylamine hydrochloride (2 mel.) and making the mixture slightly alkaline with Na.CO, (Perkin a. Church, A. 129, 108; Nietzki a. Goll, B. 18, 298). Hartley (C. J. 51, 190) hae mapped the absorption-spectrum. -B'4HCl.-B'HCl.-B'2HCl. -B',H,SO,

(β)-Naphthaleno-p-azo-(α)-naphthylamine C₁₀H₁,N₂,C₁₀H₂,NH₂ [152°]. Formed by mixing aqueous solutions of equal mols. of (β)-diazonaplithalene chlorido (from (\$)-naplithylamine) and (a)-naphthylamine hydrochloride. Yellowishbrown needles with green reflex (from alcohol). The base and its salts are far more soluble than the (a)-azo-(a)-naphthalene. H.SO, dissolves it with a violet colour (Nietzki a. Göttig, B.

20, 612).
(β)-Naphthalene-aze-(β)-naphthylamine

ŅÑ N₂H.C₁₆H, or C,0II,.Nz.C,0II,.NII, [1497].

Reddish-yellow needles. Easily soluble in benzone and acetic acid, insoluble in water. Formed by the action of amyl nitrito upon (3) naphthyl-

amine. The absorption-spectrum has been mapped by Hartley (C. J. 51, 191).

Reactions .- 1. By hoating with dilute H2SO (20 p.c.) it is decomposed with evolution of nitrogen. -2. On reduction with SuCl, it yields (B)-naphthylamino and (1:2)-naphthylenodiamine. - 3. Treated with bromine in alcoholic or acetic acid solution it is converted into di-bromo-(β)-naplithylamino and a brominated β -naphthol. 4. May be diazotised in the following manner: 15 grins, of the amidoazonaphthalene are finally suspended in a mixture of 90 grms. H2SO, and 90 grms. of water, cooled with ico and slowly treated with a concentrated solution of 5 grme. sodium nitrite. The diazo- compound is very unstable (Nictzki a. Goll, B. 19, 1281).

Acetyl derivative C₂₀II₁₁N₃Ao

prisms; easily soluble in benzenc, sparingly in alcohol and petroleum-spirit.

Benzoyl derivative C20 II N3Bz [177°]; silky red needlos; casily soluble in benzene. sparingly in alcohol and petroleum-spirit (Lawson, B. 18, 2422).

(a)-Naphthalene-azo-peri-naphthylene-diamiae C10H2-N2-C10H5(NH2)2. Prepared by the action of (a)-diazonaphthalene chloride on perinaphthylene-diamine (Stebbins, jun., B. 18, 717; C. N. 41, 117; A. C. J. 1, 445). Sol. alcohol, insol. water.—B'HCl: el. sol. water, m. eol. alcobol with a brown colour; dissolves in etrong H2SO, with a blue colour.

(8) Naphthol-azo-naphthalene sulphonio acid C10Ho(OII) N. C10HoSO3II. From (8)-naphthol and (a) diazo naphthalene sulphome acid

(W. v. Miller, B. 13, 268).
(a) Naphthol-p-azo-diphenyl snlphonio acid C10H6(OH) - N2-C1.H5.SOJII. From (a) naphthol and p-diazo-diplicnyl sulphonic acid (Carnelley a. Sohlevolmann, C. J. 49, 383). - NaA': dyes wool brown - BaA'

(B)-Naphthol-p-azo-diphenyl sulphonio acid C16H16(OH)-N2-C12H16SO3H. From (8)-naph. thol and p-diazo-diphenyl sulphonic neid. -NnA': bright red pp., sl. sol. cold, v. sol. hot, water; dyes wool red. - BaA', (Carnelley n. Schlevelmann, C. J. 49, 383).

o-Nitro-benzene-azo-acoto-acetio acid

 $\mathbf{C}_{\mathbf{g}}\mathbf{H}_{\mathbf{1}}(\mathbf{NO}_{\mathbf{r}}) = \mathbf{N}_{\mathbf{2}} = \mathbf{CH}(\mathbf{CO}_{\mathbf{r}}\mathbf{CH}_{\mathbf{J}}).\mathbf{CO}_{\mathbf{2}}\mathbf{H}.$ [185°]. Obtained by saponification of the ethyl-ether which is prepared by the action of o-nitro-diazobenzeno chlorido on an nikuline solution of ncoto-acetic ether (Bamberger, B. 17, 2415). Glistening brown plates. V. sol. acetic acid and hot alcohol, v. sl. sol. ether and cold alcohol. On heating it evolves CO, and yields o-nitro-henzene - azo - acetono CaH1(NO2). N2. CH2. CO. CH3. The same decomposition is produced by heating with nlkalis.

Salts. - A'NH, *: yellow needles. - A'Ag: erystallino pp. --A', Cu*: green pp., sol. hot wator. - A', Ba*: yellow needles. - A', 2Hg: glistening plates.

Ethyl ether A'Et [93°], glistening yellow plates or fine needles, sol. aloohol, ether, acctic acid, and liet water.

o-Nitro-benzene-azo-acetone

 $C_aH_a(NO_z) - N_z - Cll_xCO.CH_x$. [124°]. Formed by heating o-nitro-benzene-azo-aceto-acetic acid to its melting-point, or by boiling the acid or its ether with alkalis.

Preparation. -– o∙Nitranilino dissolved in absolute alcohol is diazetised by passing into the well-cooled solution a stream of N2O3, tho product is poured into iced water, the solution filtered and mixed without cooling with a dilute solution of acetaectic ether (1 mol.), and KOH (I niol.), after mixing the finid must be acid, it is digested at about 40° for 40 hours, the red pp. is then filtered off and purified (Bamberger, B. 17, 2418). Long silky yollow needles. Easily coluble in hot water, alcohol, ethor, &c.

o-Nitro-beozone-azo-acotophenone [2:1] C_sH_s(NO_s) -- N₂ -- Cl_sCO.C_sH_s. Formed together with o-nitro-benzene-azo-benzoyl-acetic other, by adding a solution of o-nitrodiazo-benzene chlorido to an iced alkaliue solution of benzoyl-acetic ether (Bamberger a. Calman, B. 18, 2565). (distening yellow needles. Easily soluble in ordinary solvents.

m-Nitro-benzene-azo-aniline

[3:1] NO, C, II, N, C, II, NII, [1:4]. [c. 210°]. From diazotised m-nitruniline hydrochloride and aniline hydrochloride (Meldola, C. J. 45, 112).

a range fern-like leaflets (from alcohol). Call armid dilute acids; forms yellow solutions in C₁₀H₁····ol, acetone, and beuzeno (Meldola, C. J. 45, (8)-diaz On reduction it gives m- and p-phenyleuepyrrol be. Salt .- (B'HCl), PtCl,

Nitro-benzene-aso-benzene sulphonic acid [8:1] C_sH₄(NO₂)-N₂-C_sH₄SO₂H [1:4]. From benzene azo-benzene eulphonicacid by nitration (Jnnovsky, M. 3, 505; 8, 60).-KA:-BaA',-

PbA'...
Nitro-benzene-azo-benzene eulphonic acid [4:1] C₆II₄(NO₂)-N₂-C₆H₄:SO₃H [1:4]. Formed, together with the less soluble isomeride just deecribed, by heating benzene-nzo-bonzene p-sulphonic acid with nitric acid (S.G. 1.41). Leaflets.-KA' (Janovsky, M. 3, 506; 5, 157; B. 16, 1486). Ammonium sulphido reduces it to an amido-benzeno azo benzeno sulphonie acid which is different from that formed from diazotised sulphanilio acid and p-pbenyleno-diamine, although on complete reduction it gives oulphanilic noid and p-phenyleno-diamine.

Di-nitro-benzene-azo-benzene-snlphonic acid [3:4:1] C₆H₃(NO₂)₂—N₂—C₆H₄·SO₃H [1:4]. From either of the two preceding acids or from benzeneazo-benzene sulphonic acid nud nitrie acid (S.G. 1.45). Ornnge leaflets. - KA'. -BaA'2: S. 7 at 68° (Janovsky, M. 3, 507; 5, 157).

o. Nitro. benzene-azo-benzoyl-acetic acid [2:1]C₆H₄(NO₂)—N₂—CII(CO.C₆H₄).CO.H.[177°] Its ethyl-ether is formed, together with o-nitrobenzeuc-azo-acotophenone, by adding n solution of o-nitro-diazo-benzene eliloride to an iced alkaline solution of benzoyl-neetic ether (Bani-berger a. Calman, B. 18, 2565). Long, yellow silky needles. Sl. sol. alcohol and ether. By long hoating at its melting-point it loses CO. giving o-nitro-benzene-azo-acetophenone.

Oxim C11II10N2(NO2)(CO2II)NOH

orange-yellow needles.

m - Nitro - bonzene - azo - m - chloro - di-methylaniline [3:1] C_cH₄(NO₂).N₂·C_cH₃Cl.NMe₂ [1:2:4] [156°]. Prepared by adding sodium nitrito (1 mol.) to a solution of m-nitranilino (1 mol.) and m-chloro-di-methyl-anilino (1 mol.) in dilute 11.SO, (Staedel a Bauer, B. 19, 1956). Reddish. yellow plates (from alcohol).

p-Nitro benzene azo di-methylaniline [4:1] NO, C, II, N., C, H, NMc, [1:4]. [230°]. hydrochlorido is deposited as crystals with steelbluo reflex when aqueous diazo-p-nitro-benzene chloride is added to aqueous dimethylaniline hydrochloride (Meldola, C. J. 45, 107).

Properties - Chocolate-brown powder, slightly solublo in alcohol whence it separates as brown needles. Solutions are orange in benzene and in glacial acotic acid, orange in conc. H.SO. red on dilution. Salt .- (B'HCl), PtCl,

m. Nitro. benzene. p. azo-dimetrylaniline [3:1] NO., C., II., N., C., H., NMe., [1:4]. [159°]. From NO., C., II., N., C., H., NMe., IICl (Meldola, C. J. 45, 120; Staedel a. Bauer, B. 19, 1954). Orange crystalline powder. Solutions in alcohol, ben-zeno, acetone, and glacial acetic acid, aro yellowish-orango; in oono. H2SO, palo orange, turned red by dilution. After reduction by HCl and zine dust, Fe₂Cl₆ forms a blue dye.

m. Nitro-bonzene-azo-(a)-naphthol [3:1] C₀H₄(NO₂)-N₂-C₁₀H₆OH [1:4]. From mdiazo-nitro-benzene and (a)-naphthol (Stebbine, jun., A. C. J. 2, 446). Brown pp., sol. water.

p-Nitro-benzene-azo-(a)-naphthol [4:1] $C_0H_4(NO_2)$ — N_2 — $\hat{C}_{10}H_4\hat{O}H$ [1:4]. From p-diazo-nitro-benzene obloride and an alkaline solution of (a)-naphthol (Meldola, C. J. 47, 661). Dull red powder, melts above 360°. V. al. sol.

biling alcohol. H₂SO, forms a violet solution; | acid (O₂H₄(NO₂)-N₂-O₂H₄:NOH)₂(?) [\$18°] of ot NaOHAq gives a blue colour. | which the sodium ealtie blue. It is re-oxidized by ot NaOHAq gives a blue colour.

m-Nitro-bensene-azo- (β) -naphthol 1:1] $C_0H_4(NO_2)-N_2-C_{10}H_4OH$ [1:2] or

[194°]. From m-diazo-N.H.C.H.(NO.) itro-benzene oblorido and an alkaline solution f (3) naphthol (Meldola, C. J. 47, 668). Lustrons range sealos (from toluenc). Insol. aqueous lkalis; sol. aleoholic KOH. H.SO, gives a asgenta red solution. It is not reduced by

ammonium sulphide.

p-Nitro-benzene-azo-(β)-naphthol [4:1] $C_0H_1(NO_2) - N_2 - C_{10}H_2(OH_1(?))$ [1:2]. [219°]. From p-diazo nitro benzene chloride and sodium (β)-naphthol (Meldola, C. J. 47, 663). Orange necdles. Insol. hot NaOHAq. Conc. H.SO.

givee a magenta-red solution.

m-Nitro-benzeno-azo-(B) -naphthol-disulphoio acid [3:1]C₁H₁(NO₂).N₁,C₁₀H₁(SO₂H)₂(OH) (?).

Prepared by acting on di-azo-m-nitro-benzene

with (β)-naphthol disulphonic acid in alkalino Stebbins, jun., A. C. J. 2, 446).

m-Nitro-bonzenc-(a)-azo. (a)-naphthylamine [B:I] NO_C_0H_i-N_2-C_0H_i_NII__[I:4]. [203°]. From NO_C_0H_iN_CI and $C_{10}H_iNII_LIICI$ (Meldola, C. J. 45, 114).

Properties.-Red needles. Solutions in alcohol, acctone, and benzene, are orange; in acetic acid, red; in cone. ILSO, violet-red turned red by Jilution. Completely decomposed by

ammonie sulphide.

p-Nitro-benzene-azo-(a)-naphthylamine [4:1] $NO_{2}C_{6}\Pi_{4}N_{2}C_{10}\Pi_{6}NH_{2}$ [1:4]. [252°]. From aqueous p-nitro-diazo-benzene chloride and alcoholic (a)-naphthylamine hydrochloride (Meldola, C. J. 43, 430). Brown needles (from Forms a crimson alcoholio solution. (B'HCl)₂PtCl₄. Salts hardly soluble in alcohol.

Reactions.—Reduces to p-phenylene-diamine

and (a,a) naplithylene-diamine.

m-Nitro benzene-azo-(\$)-naphthylamine [3:1] $C_6H_4(NO_2)-N_2-C_{10}H_8(NH_2)$ or NH_2 ⟩C₁₀Π₆. [177°]. From

O.H.(NO.)-N.H NO2, C. H., N2Cl and (8) naphthylamine. Splendid orange needles. Solutions in tolucne, chloroform, and glacial acetic acid, are orange; in alcohol and in acctone, orange but turned red by HCl; in couc. H.SO, violet (Meldola, C. J.

p-Nitro benzene-azo (β)-naphthylamino [4:1] NO₂.C₆H₄-N₂-C₁₀H₆NII₂ (\mathfrak{f}) [1:2]. [180°]. From nqueous p-nitro-dinzo-benzeno chlorido

and aqueous (8)-naplithylamino hydrochloride (Meldola, C. J. 43, 420). Needles, with golden lustre (from alcohol). Its solutions in alcohol, acetone, and chloroform are red, in benzene and toluene, orange, in cone. IL SO, violet. salts are readily soluble in alcohol. -

(B'HCl) PtCl

p-Nitro-benzene-azo-p-nitro-benzeze [4:1] C₀H₄(NO₂)-N₂-C₂H₄(NO₂) [1:4]. Di-nitro-azo-benzone. [2013]. Formed by nitration of benzene-azo-benzene (Laurent a. Gerhardt, A. 75, 73; Janovsky, M. 6, 159; 7, 135; B. 18, 1134). Red crystals (from glacial HOAc). Gives, when reduced by ammonium sulphide, a nitrolic K_xFeCy_e to p-nitro-benzene-azo-p-nitro-benzene.
m-Nitro-benzene-azo-m-nitro-benzene

[8:1] C₆H₄(NO₂)-N₂-C₆H₄(NO₂) [1:3]. A red oil, formed in the preparation of the preceding (Janovsky, M. 6, 455). Ammonium culphide and NaOH give a violet colonr.

o-Nitro-bonzene-p-azo-uitro-benzene [2:1] $C_8H_4(NO_2) - N_2 + C_8H_4(NO_2)$ [1:4], [208°], From nitro - benzene - d - azo - nitro - benzene (Janovsky, M. 7, 131). Orange lamina. Alcoholio ammonium sulphide mixed with NaOll Aq gives a permanent blue.

Nitro-benzene-azo-nitro-benzene

 $C_uH_1(NO_2)-N_2+C_uH_1(NO_2)$ [1:4]. [205°], **A** by-product in the nitration of benzene-azo-benzene-p-sulphonic acid (Janovsky, M. 7, 132), Orange lamina. Ammonium sulphide and NaOH gives a permanent blue nitrolate.

Nitro-benzone-azo-nitro-benzeno

C, II, (NO.) - N2 -- C, II, (NO.). [180]. A product of nitration of benzene-azo-benzeno (Janovsky, M. 7, 134). Pale, ashestos-like, needles. Ammonium sulphide and NaOH give a blue nitrolate changing to brown.

Nitro-benzenc-azo-di-nitro-bonzene

 $[4:1] C_b H_4(NO_a) - N_2 - C_b H_2(NO_a)_a [1:2:3 \text{ or } 5 \text{ or } 6]$ or [1:3:5]. $[112^a]$. Formed by nitration of benzeueazo-benzene (Petrieff, Z. [2] 6, 564) or benzeneazo - p - nitro - benzene (Janovsky, M. 7, 125). Yellow needles. Boiling with a mixture of alcoholic NaOH and aqueous ammonime sulphide gives a green colour, changing to brown.

Nitro-benzeno-azo-di-nitro-benzenc $[3:1] \; C_{n} H_{1}(NO_{2}) + N_{2} - \; C_{n} H_{3}(NO_{2})_{2} \; [1:3:1], \; \; \textbf{[170°]}.$ Formed by mitration of benzene - p-azo - nitrobenzene or m - mitro - henzene - m - azo - mitrobenzeno (Janovsky, M. 7, 126). Yellow tables. Alcoholie ammonium sulphide and aqueous NaOH give an olive-green colour, turning brown. Nitro - benzeno - azo - di - nitro - benzene

 $[3:1]C_0H_4(NO_1) - N_2 - C_0H_3(NO_2)_2[1:3:2 \text{ or } 5 \text{ or } 6]_4$ [121]. Formed by nitrating m-nitro-benzene-m-azo-nitro-benzene (J.). Yellow prisme. Nitrolio reaction: emerald-green changing to

Nitro-benzene-azo-di-nitro-bonzone [4:1] $C_0H_1(NO_2)-N_2-C_0H_3(NO_2)$, [1:4:3or2]. [185°]. Formed by nitrating p-nitro-benzenep-azo-nitro benzene or benzene-azo benzene-(Janovsky, M. 6, 461; B. 18, 1135). Needlee.

Nitro - benzene - azo - di - nitro - banzene [4:1] $C_{\rm e}H_1(NO_2)-N_2-C_{\rm e}H_1(NO_2)_2$ [1:4:2 or 3]. [160°]. Formed in the preparation of the proceding substance (Janovsky, M. 6, 462; 7, 125; B. 18, 1134). Yellow needles (from alcohol). Nitrolic reaction: green, turning blue. This body and the preceding, both give (1, 2, 4)-tri-amidobenzene and p-phenyl ac-diamine on reduction.

Nitre benzone azo-uitro ethano $[3:1]C_*H_1(NO_*) - N_* - CH(NO_*).CH_3$. From potaesium nitro-ethane and m-diazo-nitro-bonzene nitrate (Halimann, B. 9, 391). Yellow powder. Reduced by tin and HCl to the tin salt B"H, SnCl. of an unstable base di-amido-phenyl-ethyl-hydrazine Caffa(NfL) -- NaHa-CH(NHa).CHa.

Mitro-beuzene-aze-nitre-phenol [3:4]C.H.(NO.E.N.,C.H.(NO.)(OH)(I:3:4?].[1739]. Toward by heating the isomeric di.m.nitroazovy · benzene C.II.(NO2)-N2O--C.II.(NO2)

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with strong H2SO, for some time to about 140° (Klinger a. Pitechke, B. 18, 2552). Yellowiehbrown crystale. Discolves in alkalis with an orange colour .- A'Ag : red cryetalline pp.

p-Nitro-benzene-azo-o-oxy-benzoic acid [4:1] C₆H₄(NO₂)-N₂-C₆H₃(CO₂H)(OH) [1:3:4]. From diazotisod p-nitro aniline and a cooled alkaline solution of salicylic acid (Moldola, C. J. 47, 666). Brown needles (from dilute acetic aoid); sol. alkalis. H.SO, gives an orange solu-Blackens at 225°.

p. Nitro-benzene-azo-phenol [4:1] $C_b\Pi_1(NO_2)$ $-N_2-C_bH_0\Pi$ [1:4]. [184°]. From diazotised p-nitro-aniline and sodium phenol (Meldola, C.J. 47, 658). Golden scales; v. sl. sol. water; sol. boiling dilute alkalis. H2SO, gives an orange solution.

m-Nitro-benzeae-p-azo-diphenylamine [3:I] (NO₂)C₄H₄· N₂-C₅H₄(NHC₆H₅) [I:4]. [137°]. From NO₂C₆H₄N₂Cl and NH(C₆H₅)₂ (Meldola, C. J. 45, 118). Reddish-brown scales dola, C. J. 45, 118). Reddish-brown scales (from diluto alcohol). Solutione in alcohol, acctone, glacial acetic acid, and benzene are orange. On adding HCl to the alcoholic solution the liquid turns crimson, and, if concentrated, a brown gelatinous hydrochloride is ppd. Couc. H₂SO, forms a violet solution. After reduction by Zn and HCl, Fo₂Cl₆ forms a blue dye. Its salts are unstable.

Nitroso derivative [128].

p-Nitro-benzene-azo-di-phenylamine $[4:1](NO_z)C_0H_4-N_2-C_0H_4(NH.C_0H_5)[1:4]$ [151°]. From aqueous diazotised p-nitranilino and alcoholic diphonylamine. The pp. is treated with ammonium carbonate, and the base crystallised from dilute alcohol (Moldola, C. J. 43, 440). Brown loaflets. Solutions are orange in alcohol, turned violet by HCl; violet in conc. H.SO. The hydrochloride forms needles, with violot roflex, but is very unstable.

p-Nitro benzene-azo resorcin [4:1] $C_6H_4(NO_2) - N_2 - C_6H_3(OH)_2$ [1:2:4]. From p-diazo-nitro-benzene nitrato and resoroin in alkalino solution (Moldola, C. J. 47, 660). Brickred crystalline powder; KOHAq forms a violet, H.SO, an orange, solution.

p-Nitro-benzeae-azo-m-xylidine [4:1] (NO.), C_cH_4 — N_2 -- C_cH_2 Me. (NH.) [1:3:5:2]. [141°]. From aqueous p-nitro-diazo-benzeno ohlorido and alcoholic m-xylidino hydrochloride (Meldola, C. J. 43, 428). There results a bulky reddish pp. of NO₂-C₆H₄N₂-NHC₆H₃Me₂ which, on standing, changes to the scarlet hydrochlorido of the azo-compound.

Properties .- Brick-red needles (from dilute alcohol). Forms orange solutions in alcohol, acetone, benzeno, ohloroform, and cono. H.SO..

Salts.-The chloride, sulphate, and nitrato form red needles with violet reflex, insoluble in alcohol.-(B'HCl),PtCl.

Nitro-oarboxy-benzene-azo-nitro-benzoic acid N₂(C_aH₃(NO_a).CO_aH)₂(?). Formed by nitrating oarboxy-bonzene-azo-bonzoic acid (Golubeff, J. R. 6, 197). - Na₂A". - K₂A" 3aq. - BaA". - Et₂A": [104°].

Di-nitro-oxy-amido-benzene-azo-xylene C_sH(NO₂)₂(NH₂)(OH) -N₂ -C_sH₃Me₂. From diazo-xyleno chlorido and di-nitro-amido phenol in alkalino solution (Stebbins, jun., A. C. J. 2, 236). Brown powder, sl. col. cold water.

Nitro-oxy-bensene-aso-bensene enlphonic ${\rm acid}[3:4:1]C_sH_3({\rm NO}_2)({\rm OH}) - {\rm N}_2 - C_sH_4.{\rm SO}_sH[1:4],$ From diazoticed culphanilic acid and o-nitrophonol (Griese, B. 11, 2195; R. Meyer a. Krels, B. 16, 1331).

Nitro - oxy - benzene - azo - naphthalene sulphonicacid C₆H₃(NO₂)(OH) -N₂-C₁₀H₆·SO₃H. From diazotised (a)-naplithylamino sulphonic acid and o-nitro-phenol (Stebbins, jun., A. C. J. 2, 236). Red ncedles, v. eol. water.

Di.nitro-oxy-benzene-azo-napthylamine anlphonicacid C,H,(NO.),(OH).N,C,OH,(NH.).SO,H. From diazo-di-nitro-phenol and (a)-naphthylamino sulphonic acid (Stebbins, jun., A. C. J. 2, 446). Reddish-brown dyo; sol. water.

Nitro oxy-honzene-azo-nitro-phenol.

Ethyl ether [2:x:1]C₆H₃(OEt)(NO₂).N₂.C₄H₃(OEt)(NO₂)[1:2:x]. [190°]. Formed by nitrating o-oxy-benzene-oazo-phenol ethyl ether, and separated from the isomeric compound by alcohol, in which it diesolves (Andreae, J. pr. [2] 21, 322). Neodlee (from alcohol).

Nitro-oxy-benzene azo-nitro-phenol.

hydrazine.

Ethyl ether $[2:x:I]C_aH_a(OEt)(NO_2).N_2.C_aH_a(OEt)(NO_2)[1:2:x].$ [285°]. Formed together with the preceding (q, v). Brownish-red crystals (from ohloroform). Insol. alcohol. Dissolves without change in cono. H.SO,. Reduced by alcoholic ammonium sulphide to the di-cthyl other of dinitro-dioxy di-phonyl-

 $\begin{array}{lll} \textbf{Di-nitro-oxy-benzene-azo-phenol} & \textbf{eulphonio} \\ \textbf{id} & \textbf{C_oH_2(NO_2)_2(OH)-N_2-C_oII_3(OH).SO_3H.} \end{array}$ From diazotised di-nitro-amido-phenol and an alkaline solution of phenol o-sulphonic acid (Stebbins, jun., A. C. J. 2, 236; C. N. 42, 44). Brown lustrous needles, sl. sol. hot water.

Nitro-diphenyl-azo-nitro-diphenyl (?) $[4:I] C_0H_1(NO_2).C_0H_1-N_2 C_0H_4.C_0H_4(NO_2)[1:4].$ [187°]. From p-dinitro-diphenyl and sodium amalgam (Wald, B. 10, 137). Yellow powder (from alcohol).

m-Nitro-tolucne-azo-aceto-acetic acid $\{4:2:1\}$ C_a H_a (CH_a) (NO_a) $-N_a$ -CH (CO,CH_a) CO_a H[176°]. Obtained by saponification of the ethylcther formed by the action of nitro-diazo-toluene chlorido (from nitro-p-toluidine [114°]) on an alkalino solution of aceto-acetic ether (Bamberger, B. 17, 2121). Long yellow eilky needles. V. sol. hot alcohol and HOAo.—A',Ba.

m-Nitro-toluene-azo-acetone [4:2:1]C₆H₃(CH₃)(NO₂)-N₂-CH₂·CO.CH₃·[134°]. Formed by the action of a dilute solution of aceto-acetic ether (1 mol.) and KOII (I mol.) on a solution of nitro-diazo-toluene nitrate (from nitro-p-toluidinc[114°]) (Bamberger, B. 17, 2421). Orange-red prisms. V. sol. alcohol and ether.

m-Nitro-tolnene-azo-acetophenoae [4:2:I]C,H₂(CH₃)(NO₂)-N₂-CH₂.CO,C_aH₃, [168°]. Glistening yellow needles. Formed, together with nitro-toluenc-azo-benzoyl-acetic ether, by adding a solution of nitro-diazo-toluene chlorido (from-m-nitro-p-toluidine [II4°]) to an iccd alkaline solution of benzoylacetic ether.

Ketoxim C₁₃H₁₃N₂(NO.);NOH:[174°]; orange needles (Bamborgor a. Calman, B. 18, 2566).

m-Nitro-p-toluene-azo-benzoyl-acetic acid [4:2:1] $C_aH_a(CH_a)(NO_a).N_a.CH(CO.C_aH_a).CO_aH_a$ [194°]. Its ethyl ether is formed, together with m-nuro-p-toluene-azo-acetophenone, by adding a solution of m-nitro-p-diazo-toluene chloride to an iced alkaline solution of benzoyl-acetic ether (Bamberger a. Calman, B. 18, 2566). yellow needles. V. sl. sol. cold alcohol and acetic acid, more easily at the boiling-point.

p-Oxy-benzene-azo-benzene-m-sulphonic acid [4:1] $C_{n}\Pi_{n}(OH) - N_{2} - C_{n}\Pi_{n}SO_{3}H$ [1:3]. From diazotised amido-benzene m-sulphonic acid and an alkaline solution of phenol. Leaflets, with violet reflex; insol. ether, v. sol. water and alcohol. - KA': long needles (Griess, B, 11, 2194).

p-Oxy-benzene-azo-benzene-p-sulphonic acid [4:1] $C_6 H_1(118O_3) - N_2 - C_6 H_4(OH)$ [1:4]. paoline Y. Prepared by the action of an aqueous alkaline solution of phenol on p-diazobenzene snlphonic acid (Griess, B. 11, 2192). Yellowish red prisms. V. sol. water and alcohol.

Salts. - BaA"; orange pp.-BaA', 2aq. -BaA', 5aq: minute orange tables, sl. sol. water. -KA': yellow rhombic leaflets, S. 26 at 15? (Wilsing, A. 215, 232).

p-Oxy-bonzeno-azo-benzene sulpbonio acid [4:1] C₆H₄(OH) N₂ - C₆H₄, SO₂H. From azoxybenzene (1 pt.) and fuming ILSO, (5 pts.) at 110° (Limpricht, B. 15, 1295; Wilsing, A. 215, 229); Tschirwinsky (J. R. 5, 217) considers this neid to be identical with the preceding. Small lustrous reddish plates, v. sol. water, m. sol. dilute acids or alcohol. Br does not act on the potassium salt. SuCl2 forms no aniline by reduction. Salts.—KA' aq: S. 85 at 15°. - BaA', AgA. - MgA', 6aq. - CuA', 6aq. Chloride, [122°]. Orange 6- or 8-sided

plates.

Amide. [212°]. Plates.

Di-oxy-benzane-azo-benzene sulphonic acid [4:2:1] C₆H₃(OH)₃ -N₂-C₆H₄.SO₃H [1:4]. Tropactin O. Chrysoine. Formed by sulphonating benzene-azo-resorcin at 100° (Witt, C. J. 35, 183) or from diazotised amido-benzeno p-sulphonic acid and resorein dissolved in KOHAq (Griess, B. 11, 2195). Red leaflets with steelblue reflex; v. sl. sol. alcohol and cold water. - KA'.—BaA'. 4aq.

The absorption-spectrum has been examined

by Hartley ($\dot{C}.~J.~51,~182$).

Di-oxy-benzane azo-benzone m-sulphonic acid [4:2:1] $C_6H_4(OH)_2-N_2-C_6H_4$. SO₃H [1:3]. From resorein and diazotised amido-beazene m-sulphonic acid. Orange needles. - KA': hygroscopic needles.

Tri-cxy benzono azc-benzeno sulpbonio acid $[2:4:6:1] C_6H_2(OH)_3-N_2-C_6H_4.SO_3H[1:4]$. From diazotised amido-benzene p-sulphonic acid and an alkaline colution of phloroglucin (Stebbins, C. N. 42, 44; A. C. J. 1, 465; 2, 236; B. 13, 716). Yellow leaflets with green lustre. -NaA': yollow leaflets, easily soluble in water.

 $p ext{-}\mathsf{Oxy-benzene-}m ext{-}\mathsf{azo-benzoic}$ acid

[4:1] C.II.(OH) -N. -C.II.,CO.H [1:3]. [220°]. Formation.—1. From m-diazo-benzoic acid and phenol (Griess, B. 14, 2032), -2. By gently warming m-carboxy-diazo-benzene m-carboxyanilide C₆H₄(CO₂H).N₂.NH.C₆H₄(CO₂H) phonol, m-amido benzoic acid being eliminated (Heumann n. Oeconomides, B. 20, 906). Red needles or plates; sol. alcobol and ether, sl. sol. water. Dyes wool and silk yellow.-BaA's 31aq. Di-cxy-benzene.m-azo-benzoic acid

[4:2:1] C₆H₂(OH)₂-N₂-C₆H₄·CO₂H [1:3]. Pre-

pared by the action of m-diazobenzoic acid on an alkaline solution of resorcin (Grieee, B. 14, 2034). Brownish rod needles or brownish-yellow plates. Sol. alcohol. Dyos wool and eilk yellow.

Oxy-benzene-azo-p-cresol. Ethyl ether

[4:1] $C_0H_4(OEt)-N_2-C_0H_4(OH_3)(OH)$ [1:5:2]

[104°]. Formed N,H.C,H,Me(OEt)

by combining p-diazo, phenetol with p-cresol (Liebermann a. Kostanecki, B. 17, 883). Golden plates. Sol. alkalis. Dissolves in H.SO, with a brown colour. On reduction it gives p-amidophenetol and amido-p-cresol.

o-Oxy-benzene-aze-o-cresel. Methyl cther $C_aH_a(OMe) - N_a - C_aH_aMe(OH)$. [68°]. From diazotised o-anisidine and o-cresol (Kunonnikoff, J. R. 1885, 369). Di-methyl ether [103°]. o-Oxy-benzeno-azo-m-crosol. Methyl

ether. [161°]. Prepared like the preceding (K.).

Di.oxy-benzene-azo-\psi-cumsns

[1:2:1]C.H. (OII)..-N., -C.11, Me. [199°]. Formed, together with the disazo-compound, by combining diazo - cumeno chlorido (from amidopseudo-cumene [62]) with resorcin (Liebermann a. Kostanecki, B. 17, 131, 882). Small red needles. Dissolves in alkalis with a brownish yellow colour.

Di - oxy - bonzene - azc - hydroquinone [5:2:1] $C_6H_3(OH)_2-N_2-C_6H_3(OH)_2$, [1:2:5]. Azohydroquinone. Tetra-methyl derivative C_nH₃(OMe), N. C_oH₃(OMe), [140°]. Formed by reduction of nitro-di-methyl-hydroquinone in alkaliue solution (Baessler, B. 17, 2124; C. C. 1886, 671). Red needles. V. sol. alcohol, bonzene, chloroform, and CS., v. sl. sol. water. Dissolves in strong HCl with a blue colour.

Tetra-ethyl derivative [128]. From nitro-di-ethyl-hydroquinone, powdered zino, and alcoholie potash (Nietzki, B. 12, 39).

Di-oxy-benzanc-azo-naphtbalene snlphonic acid [4:2:1] C₄H₃(OH)₂-N₂-C₁₆H₄-SO₃H [1:4]. From diazotised (a) naphthylamine sulphonio acid and an alkaline solution of resorcin (Stebbins, jun., A. C. J. 2, 36; C. N. 42, 41). Darkbrown needles, sol. water.

p-Oxy-benzene-azo-(a)-naphtbylamine [4:1] $C_0H_1(OH)-N_2-C_{10}H_0NH_2$ [1:4]. [170°]. Prepared by the action of p-diazophenol nitrate on (a)-naphthylamino (Weselsky a. Benedikt, B. Orange needles (containing 3aq) .-B' H.SO, 6aq: green needles, insol. water.

o-Oxy-benzene-azc-(8)-naphthylamine

[2:1] $C_sH_s(OII).N_sC_{10}H_s$ or $C_{10}H_s < N_sH.C_sH_sOH_s$ o-Oxy-benzene-hydrazimido-naphthalene. [193°]. Formed by combining o-diazo-phenol with (8)naphthylainine. Slendor red needles (from alcohol or acetic soid), or dark red plates (from benzene). It dissolves in aqueous or alcoholic NaOII, but is insoluble in water. By heating at 150° with HCl it is split up into (\$\beta\$)-naph-thylamine, pyrocatechin, and nitrogen. Bromine in cold acetic acid solution gives di-bromo-(8)naplithylamine, a brominated pyrocatechin, and nitrogen. On reduction with zinc dust and acctic acid it yields (1:2)-naphthylene-diamine.

Acetyl derivative [198°]. Benzoyl derivative [183°].

0 G 2

Methyl ether C.H. (OMe).N.C., H. [1380]; dark-red monoclinio prisms; insoluble in water (Sachs, B. 18, 3125).

p-Oxy-benzene-azo-(β)-naphthylamine [4:1] C₀H₄(OH).N₃C₁₀H₅ or C₁₀H₆; N.H.C.H.OH.

p-Oxy-benzene-hµlrazimido-naphthalene. [193°]. Formed by combining p-diazo-phenol with (β)naphthylamine. Flatred prisms. V. sol. benzene and acetic acid, insol. water. By heating at 150° with HCl it is split up into (B)-naphthylamine, hydroquinoue, and nitrogen. Bromine in cold acetic acid solution gives di-bromo (B)naptlitlylamine, a brominated hydroquinone, and nitrogen. On reduction with zinc dust and acctic acid it yields (1:2)-naphthylene-diamine (Sachs, B. 18, 3125).

Mono-acctyl derivative [218°] Mono-benzaylderivative [241°].

Oxy-benzene-azo-orein. Methyl ether C.H.(OMe).N.,C.H.(OH).Me. Hair-like needles (Stebbins, A. C. J. 5, 32).

o-Oxy-benzene o-azo phenol

[2:1] C.H.(OH) N. C.H.(OH) [1:2]. o-Azo-phenol. [171°]. S. (alcohol) 3 at 20°. Prepared by fusing o-nitro-phenol with KOH (Weselsky a. Benedikt, B. 11, 398; A. 196, 344). Golden leaflets; may be sublimed. Insol. water.

Reactions. -1. Bromine added to an ethereal solution forms a tetra-bromo-derivative. Chlorine passed into an acctic acid solution forms a tri-chloro derivativo C₁₂H₁Cl₃(O(I),N₂ [235°] (Bohn a. Honnaum, B. 17, 275).—3. Nitric acid forms (1, 2, 4)-dt-nitro-phenol Ethyl other {C,H,(OEt)},N, [131°], c-Nitro-phenyl-ethyl ether C,H,(OEt)(NO.)

is dissolved in alcohol and reduced with sodiumamalgam (5 per cent.), the liquid being allowed to become hot. The product is poured into water, and the pp. extracted by strong HCl which dissolves the azer-compound. On pouring the HCl solution into water, a ano-phenetol is thrown down (R. Schmilt a. Möhlan, J. pr. 126, 202). Properties .- Long red prisms (from alcohol). Melts under water, but insoluble therein and not volatile with steam. It begins to boil at 240°, but suffers decomposition at the same time. Reactions .- 1. Reduced by alcoholic ammoninm sulphide to the corresponding hydrazo-compound (q. v.). 2. Gold funning 11NO, forms a nitro-, and a disnitro-, derivative.

m-Oxy-benzene-m-azo-phenol Ethylether [B:1] C.H. (OEt.) = N. - C., H. (OEt.) [1:3]. m-Azo-phenetol. [91°]. Formed by reducing m-nitrophenetol in alcoholic solution with sodium amalgam (M. Buchstab, J. pr. [2] 29, 299).

Properties. - Orange prisms (from alcohol). Sol. ether. Insol. water and (difference from ocompound) in conc. HCl. Reduced by H.S and alcoholic ammonia to m-hydrazo-phenetol.

p-0xv-benzenc-p-azo-phenol [4:1] C, U, (O11) N, C, H, (OH) [1:4], phenol. [2042].

Formation. -1. By potash-fusion from p-nitroso-phenol (Jacger, B. 8, 1499), p-nitrophenol (Weselsky a. Benedikt, A. 196, 339), poxy-benzene-azo-benzene p-sulphonic acid, or psulpho benzene azo benzene p sulphonic acid (Bohn a. H. maann, B. 15, 8037).—2. From p diazo-phenol nitrate and phenol potassium.

Properties .- Slender brown needles with blue reflex (containing aq); sl. sol. water, v. sol. alcohol. Bromine gives a tetra-bromo derivative. Nitrio acid (1, 2, 4)-di-nitro-phenol. Chlorine givee tri-chloro phenol.

Salt.—BaC₁₂H₂N₂O₂4aq. Ethyl ether C₂H₄(OEt)—N₂—C₄H₄(OEt). p. Azo-phenetol [160°] (S. a. M.); [158°] (A.). Formed by adding sodium-amalgam (5 p.c.) to an alcoholic solution of p-nitro-phenyl-ethyl ether, precipitating the product with water, boiling it with dilute HCl to remove amido-phenylethyl-ether, and crystallising from alcohol (Schmitt a. Möhlau, J. pr. 126, 199; Hepp, B. 10, 1652; Andreac, J. pr. 129, 333). Properties.— Glittering golden plates. M. sol. cold alcohol, v. sol, ether and chloroform. Distils with difficulty. Reactions. - 1. Finning HNO, forms the ethyl ether of di-miro-phenol (q. v.) and two isomeric tri-nitro-azoxy-phonol ethyl ethers .- 2. HClAq at 130° gives EtCl and p-axo-phenol N. (C.H.OH)2. but at 150° chloro-p-amido-phenol ie got (Schmitt, J. pr. [2] 19, 313).

p-Oxy-benzenc-azo-diphenyl sulphonio aoid [4:1] $C_a\Pi_a(OH) - N_2 - C_b\Pi_aC_a\Pi_a$, SO_aH . From diazotised p-amido-diphenyl sulphonic acid and phenol (Carnelley a. Schlevehnann, C. J. 49, 380). Yellow dye.—BaA'2: insol. cold water.

m . Di - oxy - henzene-azo-diphenyl eulphonio acid [4:2:1] C₀H₃(OH)₂ -N₂ -C₀H₁,C₀H₂,S₀H₄.

From diazotised p-amido-diphenyl sulphonic acid and resorein (Carnelley a. Schlevelmann, C. J. 49, 382).—NaA'.—BaA'.

p-Di-oxy-benzene-azo-diphenyl sulphonic acid [5:2:1] C,H,(OH), N,,C,H,C,H,SO,H. diazotised p-amido-diphenyl sulphonic acid and hydroquinone (Carnelley a. Schlevelmann, C. J. 49, 382).

p-Oxy-bonzeue-azo-phlorogluciu

Call₁(OH) -N₂-C₃H₂(OH)₃. Two modificatione appear to be formed by the action of p-diazophenol nitrate on phloroglucin (Weselsky a. Benedikt, B. 12, 227). (a) Red crystalline powder (containing 3aq), sol. alcohol. (b) Green amorphous mass, insol. alcohol.

p-Oxy-benzene-azo-reservin. Ethyl-ether [4:1] C_aH_b(OFt)—N_a—C_aH_a(OH)_c [1:2:4]. [167°]. Formed by the action of p-diazo-phenetol on resorcin (Liebermann a, Kostanecki, B. 17, 883). Red plates. Sol. alkalis. Dissolves in H.SO. with a brownish-red colour.

Dioxy-benzenc-azo-recorciu

[2:6:1] C₈H₃(OH)₂-N₂-C₆H₈(OH)₂[1:2:4]. Tetraory-aro-benzene.

Diethyl ether CaHa(OEt), Nu. CaHa(OH), [182']. Formed by combination of the di-ethyl ether of o-diazo-resorein C, H, (OEt), N, OH [6:2:1] with resorcin. Light reddish-yellow needles. V. sol. alcohol and other, insol. water. Diecolves in conc. alkali with a reddish yellow colour, which becomes a splendid carmine red on dilution (Pukall, B. 20, 1151).

Di-oxy-benzene-azo-resorcin

Di-cthyl other

C, H3(OFt), -N2 - C, H3(OH), [193]. Formed by combination of the di-cthyl other of p-diazoresorein C₈H₃(OEt), N₂OH [4:2:1] with resorein. Small brown needles or chort prisms with green reflex. V. sol. alcohol, ether, and aqueous alkalis, insol. water. (Pukall, B. 20, 1144). Carmine-rod dyestuff

Oxy-benzene-azo-resoroin-di-eulphonic acid Mono methyl ether

C₆H₄(OMe).N₂.C₆H(OH)₂(SO₃H)₂. Yellow plat (Stebbins, A. C. J. 5, 55).—BaA"aq: needles. Yellow plates

p. Oxy.benzene-p. azo-tolneno
[4:1] C,H,Mo-N,-C,H,(OH) [1:4]. [151].

Formed by gently warming p-diazo-toluene-ptoluide with phenol, p-toluiding being split off (Heumann a. O conomides, B. 20, 905), or by warming p-nitroso-toluene with p-toluidine acctate (Kimich, B. 8, 1030). Orange prisms, with blue reflex.

m-Di-oxy-benzene-o azo-toluene

[2:1] $C_6H_4(CH_4) = X_4 + C_6H_3(OH)$, [1:2:4]. [195°], (W.); [178°] (F.). Formed by the action of σ -diazotoluone on resorcin. Brownish-red felted needles. Acetyl derivative [75°]. Orange-yellow

plates (Wallach, B. 15, 2825; cf. Fischer, B. 20, 1579).

m-Di-oxy-henzone-p-azo-toluene

[4:1] $C_a H_a (CH_a) = N_a = C_a H_a (OH)_a$ [1:2:4] {184°}. Prepared by the action of p-diazo-toluene on resorcin, or by gently warming p-diazo toluenep-toluide with resorcin, p-toluidine being split off (Heumann a. Occonomides, B. 20, 906). Reddish-yellow needles: sol. alcohol, ether, and aqueous alkalis. Acetyl derivative [98].

Oxy-benzene-azo-toluidine [5:2:1] $C_aH_a(GH_a)(NH_a)-N_a-C_aH_aOH$ [1:1]. [172] Formed by saponitication of the acetyl derivative. Slender brown needles. Sol. alcohol and ether, v. sl. sol. cold water. Dissolves in aqueous acids and alkalis.

Acetyl derivative

C₆II₁(CH₃)(NIIAc) - N₂-C₆II₁OH. [253°]. Prepared by diazotising the mono-acetyl derivative of (1:2:4)-tolylene-diamine and combining it with phenol (Wallach, B. 15, 2826). Yellow plates. Sol. acetic acid and in aqueous alkalis, sl. sol. alcohol.

Di.oxy.benzeuo.azo.xylene (Wallach, B. 15, 25). Formed by adding diazo-xyleno chlorido to an alkaline solution of resorcin.

Di-oxy-benzeno-azo-xylene sulphonio acid [4:2:1] $C_6H_3(OII)_2 - N_2 - C_6H_2Ne_2SO_3H$. Prepared by the action of an alkaline solution of resorein upon diazo-xylene sulphonic acid. Slender orange needles, sl. sol. hot water; m. sol. alcohol (Griess, B. 11, 2197).

Oxy-carboxy-benzene-azo-naphthalenc

O₁₀H.,—N₂—C₄H₃(OH)(CO₂H) [1:4:3]. From naphthylamine hydrochloride (9 g.), HCl, water (500 g.), and NaNO₂ (3:45 g.) at 0°, the filtrate being ponred into a solution of salicylic acid (6.9 g.) and NaOH (6 g.) in water (500 g.). The liquid is filtered and the sodium sait ppd. by NaCl (P. F. Frankland, C. J. 37, 747). Salt.— NaA'. S. 07 (cold). Dyes silk pale yellow. Reduced by Sn and HCl to naphthylamino and amido-salicylic acid, CaH, (NH,)(OH)COaH.

Oxy.carboxy.benzene-azo-(\beta)-naphthol

Mono-methyl derivative C_sH₃(OMe)(CO₂H)—N₂-C₁₀H₆(OH). Prepared by the action of diazo-anisic acid on an alkalino solution of (β)-naphthol (Griess, B. 14, 2039). Small red needles or plates (containing 11 aq). Si. eol. alcohol. A', Ba 4aq : red minute needles.

Oxy-carboxy-benzene-azo-(β) naphtbol sulphonic acid. Methyl derivative

C_zH_z(OMe)(CO_zH).—N_z—C_{te}H_z(OH)(SO_zH).From diazo anieto acid and (β) naphthol enliphonic acid. Brown needles. Dyos wool scarlet. BaA", 8aq (Griess, B. 14, 2039).

Oxy-carboxy.benzene.azo.(3).naphthol-(a).disulphonic acid Methyl derivative

 $C_bH_a(OMc)(CO_aH) - N_a - C_{10}H_4(OH)(SO_aH)_a$. Propared by the action of diazo anisic acid on an adkuline solution of (B), naylithol (a)-di-sulphonio acid (Griess, B. 14, 2010). Small dark-red needles (containing 3aq). Sol. water, and alcohol, insol. ether. Dyes a blueish shado of scarlet. A"HK, baq: red crystals, col. hot water.

Oxy-carboxy-benzene-azo-oxy-benzoic acid

Dimethul derivative

 $C_nH_n(OMe)(CO_nH) = N_2 - C_nH_n(OMe)(CO_nH)$ Formed by the action of sodium numbrane on an alkaline solution of nitro-anisic acid. water. BaA" aq (Alexejeff, C. R. 55, 172).

Dioxy. carboxy . methyl-phthalide-azo-dioxyphthalide-acetic acid

Tetra-methyl derivative Call No. On i.e.

N. {C, H(OMo), CO - O }

Azo-meconic-acetic acid. [c. 257°]. Obtained reduction of nitro-di-methoxy-phthalideacetic acid CaH(OMe) (NOz) CO CO CH CHarCOaH

with zine-dust and aqueous NH2. Yellow crystale. lusol, water. Dissolves in conc. Il. SO, with a deep blneish-violet colour (Kleeman, B. 20, 880).

Tri - oxy - carboxy - tolucno - azo-tri oxy-toluio acid. Anhydride of the tetra methyl derivative.

[c. 245]. Formed by reduction of nitroso-opianio acid C₆H(OMe)₂(NO)(CHO)(CO₂H) with zincdust and aqueous NII, Dissolves in alkalis with a yellow colour, in cone. It SO, with an intense purple colour. — AgA': microscopio needles (from hot water) .- EtA': [101°], yellow needles, v. sol. alcohol, ether, and benzone (Kleemann, B. 20, 878).

(β)-0xy-naphthalone-azo-hippurio acid $CO_{\alpha}H^{\prime}CH_{\alpha}NH_{\alpha}CO_{\alpha}G_{\alpha}H_{1} + N_{2} - C_{1\alpha}H_{\alpha}(OH)$. pared by the action of diazo-hipparic acid on an alkaline solution of (B)-numbthol (Griess, B. 14, 2010). Reddish-yellow needles. alcohol, v. sl. sol, water and ether.

(a)-0xy-(a)-naphthaleno-(a)-azo-naphthaleno-(a)-sulphonic acid

[4:1] $C_{10}H_{a}(OH) - N_{4} - C_{1a}H_{a} \cdot SO_{3}H$ [1:4]. From diazotised (a)-naphthylamine sulphonic acid and (a)-naphthol. The all orntion-spectrum has been examined by Hartley (C. J. 51, 198).

(β) - 0xy naphtbal ne azo naphthalene sulphonic acid C10H1 (OH) - N2 - C10H SO3H. From diazotised (a) naplithylamine sulphonic acid and (β)-naphthol (Caro; Griess, B. 11, 2199). Reddish-brown needles (from alcohol). Red dye .-The absorption-spectrum has been examined by Hartley (C. J. 51, 197).

Oxy - propyl - carboxy - benzeno - azo - cxypropyl-benzoic acid

 $CO_sH_sC_sH_s(C(OH)Me_s)-N_s-C_s\Pi_s(C(OH)Me_s)CO_sH_s$ Formed by reduction of nitro-oxy-propyl-benzoic acid with sodium - amalgam and water (Widmann, B. 15, 2550). Yellow plates. V. sl. sol.

most ordinary solvents, sl. sol. acetic acid .-- | Na.A" IOaq: thin red rectangular tables.

Oxy-snlpho-benzene-azo-henzoic acid [3:1] $C_6H_4(CO_2H) - N_2 - C_6H_3(OH)(HSO_3)$ [1:4:3]. Prepared by the action of m-diazobenzoic acid on an alkaline solution of phenol-o-sulphonic acid (Griess, B. I4, 2033). Brownish-red crystals (containing ½aq). Sol. water, alcohol and other. Ysllow dyc. Salts.—A"HK aq: yellow plates or needles, sl. sol. cold water.—A", II.Ba: small yellow needles or plates.—A"Ba aq: yellow crystalline pp.

Oxy-snlphc-benzene-azc-naphthalsne sulphonic acid C_aII₃(OII)(SO₃H)—N₂—C₁₀II₃.SO₃H. From diazotised (a) naphthylamins sulphonic acid and phenol sulphonic acid (Stebbins, A. C. J.

2, 446).

Oxy-tolusne-azo-toluene snlphenic acid [4:2:1] $C_4H_4(CH_4)(SO_4H)-N_4-C_4H_4(CH_4)(OH)$ [1:5:2]. Formed by the action of p-diazo-tolusus sulphonio acid (by diazotising p-tolnidine-sulphonic acid) en an alkaline solution of p-cresol (Nölting a. Kohn, B. 17, 358). Reddish-brown crystals with violet reflection. V. sol. water, sl. sol. alcohol. Salts .- A'Na: soluble yellow plates. --A'2Ba 4aq: small reddish brown needles, el. sol. hot water.

Diphenyl-azc-diphenyl C.H.C.H.-N2--C.H.C.H. [250°]. Orangered plates. Sol. ether, insol. water, alcohol, and aestio acid. Formed by reduction of p-nitrodiphenyl with sodium amalgam, and by the dry distillation of hydrazo-diplicnyl. Prepared by oxidising an alcoholic solution of hydrazo-diphenyl with Fe Cl. (Zimmermann, B. 13, 1962).

Phenyl - acstic - aze - phenyl - aestic acid v. exo-Carboxy-toluene-azo-phenyl-acetic acid. Phsnyl - amide - bsnzsne - aze - benzene - snl-

phonic acid [4:1] C₈H₄(SO₃H) -N₂-C₆H₄NHPh [1:4]. Troparoline O.O. Prepared by the action of p-diazobenzana sulphonic acid on an alcoholio solution of diphenylamine (Witt, C. J. 35, 187; B. 13, 262). Steel-blue hair-like needles. Sl. sol. water. Salts .- A'K: flat yellow needles; sl. sol. cold water.—A'Na.—A'NH₄.—A'NHMe₃: large yellow leaflsts.—A'₂Ba and A'₂Ca: insoluble yellow pps.

Phanyl-glycellic-o-aze-phenyl glycellic acid

N₂(C₆H₄,O.CH₂,CO.H). [162°].

Preparation. — o Nitrophenylglycollio acid (18.6 g.) water (140 g.) and Na CO3 (5 g.) are treated at 60° with sodium-amalgam (215 g. of 4 p.c. amalgam). The crystals which separate on cooling are disselved in very little water and the acid is ppd. by acetic acid. The product is recrystallised several times from alcohol (A. Thate, J. pr. [2] 29, 161).

Properties.-Orange silky needles, containing 2aq (from water or diluto alcohol). When dry (at 110°) it is brick-rod. Sol. other, alkalis and strong acids. Its solutions are yellow or red.

Reactions .- I. Aqueous colution is acid to litmus and gives with AgNO, a red gelatinous pp., and with Pb(OAc), a flocculent yellow pp.—2. Reduced by alcoholic NH, and ILS to the corresponding hydrazo compound, the potassium salt of which, N2H2(C6H4O.CH.CO2K), 3aq, eryetallises from alcohol in rhomhohedra.

Salts.-K, A" Saq. Orangs plates. solution gives with BaCl, a red orystalline pp.; with Pb(OAc), an orange flocculent pp.; with

AgNO, a red flocculent pp.; with Fe,Cl, an orange pp.; with CuSO4, a brownish-yellow pp.; with HgCl2, on boiling, a red pp.; with MgSO4 Na, A' 3aq.—Ag, A' 3aq.—BaA' 2aq.—CaA' 8aq.

Ethyl ether. Et, A''. [III']. Red oryetals.

Phenyl-glyoxylic-azc-phsnyl-glyoxylic acid CO.H.CO.C.H. - N. - C. H. CO.CO.H. Azo-ben-zoyl-formic acid. Orangs needles (containing 2aq and melting at [135°]. When dry it melts at about 151°. Prepared by reduction of mnitro-benzoyl-formic acid with FeSO, and KOH: yield, 50 p.c. (Thompson, B. 16, 1308). Sl. sol. ether and cold water, insol. acidulated water, chloroform, or benzene. Very etable body. A cold saturated aqueous solution of the acid gives with BaCl, or CaCl, a micro-crystalline pp., with AgNO, a yellow flocculent precipitats.— A"Ba: orange crystalline pp., insol. water.— A"Ag2: orange-yellow slightly soluble pp.

Phenyl-pyrrel-aze-hsnzsne

 $C_6H_5.N_2.C = CH$ $NPh \mid$ C.H.N.C.H.NPh probably HC = CH

[II7°]. Prepared by adding diazobenzene chloride (1 mol.) to plienyl-pyrrol (I mol.) dissolved in alcohol containing AcONa. Brown prisms or long reddish-yellow needles with blueish reflection. V. sol. alcohol. Dissolves in conc. H, SO. with a reddish-violet colour, in alcoholic HCl with a blood-red colour. Reduced by zinc dust and NH3 or NaOH to aniline and (probably) amidophonyl-pyrrol (O. Fischer a. Hepp, B. 19, 2256).

Di-propyl-amide - henzene - azc - di - propylanilins $C_aH_1N(C_3H_1)_2-N_2-C_aH_4N(C_3H_1)_2$. Dipropyl-aniline-azyline. [90°]. Formed by passing NO through an alcoholic solution of di-propylanilino (Lippmann a. Fleissner, B. 15, 2140; 16, I417). Large trimetric crystals, a:b:c:= I: 629: 913. Picrato B"(C, IL (NO.), OH); orange-red insoluble crystals. — Psriodide B", Ia: violet glistening needles.

Pyrrel - azc - benzsne C.H.-N2-C.H3NH C_all_a.N₂.C - CH NH | NH | . [62°]. Prepared by probably

adding the calculated quantity of a moderately conc. solution of diazo-benzenc chloride to a wellcooled solution of pyrrol (2 pts.) in alcohol (100 pts.) with addition of sodium acetate (5 pts.) (Fischer a. Hepp, B. 19, 2251). Yellow needles. V. sol. alcohol, ether, and petroleum spirit, sl. sol. water. It has basic properties. Easily reduced (e.g., hy zine-dust and alkalis) to aniline and (prohably) amido-pyrrol. Readily combinss in alkaline or neutral colution with a further quantity of diazo-compound, giving disazo-hodies. Dissolves easily in dilute HCl with a reddish-yellow colour. Dissolves in conc. H.SO, with a yellow colour. The platino-chlorido forms small red cparingly solubls needles.

Pyrrol-p-azc-di-methyl-aniline

 $C_{8}H_{1}(NMe_{2})-N_{2}-C_{4}H_{8}NH.$ [159°]. by combining p-diazo-di-msthyl-amido-benzsne with pyrrol in dilute alkaline solution. Glittering green plates. Dissolves in very dilute HCl with a grass-green colour, in cone. HCl with a greenish-yellow colour. PtCl, gives a dark green amorphous pp. (Fischer a. Hspp, B. 19, Fyrrol-(a)-and-naphthalone C. H. N. C. H. NH

Ö₁₀H₄.N₂.G = CH NH | HC = CH probably . [103°]. Formed by

adding (a)-diazo-naphthalene ohlorido (1 mol.) to pyrrol (1 mol.) dissolved in nlcohol containing sodium acetate. Reddish-yellow plates. sol. alcohol (O. Fischer a. Hepp, B. 19, 2255).

Pyrrol-(\$)-azo-naphthaleno

 $C_{10}H_7.N_2.C = CH$ C10H7-N2-C4H2NH probably ŇΗ HC - CH

[101°]. Prepared by adding (\$\beta\$)-diazo-naphthalene chloride to an alcoholic solution of pyrrol containing sodium acetate. Gold-bronzy plates (O. Fischer a. Hepp, B. 19, 2255).

Pyrrol-p-azo-toluene C.H.Me--N2--C.H3NH C,H,N,C=CH

probably ŇIII [82°]. Prepared by HC = CH

adding p-diazo-toluene chloride (1 mol.) to pyrrol (1 mol.) dissolved in alcohol containing sodium acetate (Fischer a. Hepp, B. 19, 2254).

m-Sulphi - bsnzene - azo - bonzene-m-sulphinic acid C₆II₄(SO₂H) -N₂--C₆II₄(SO₂H).

Azo-benezue di-sulphinic acid. Obtained from C_cH₁(SO.SII).N_zC_cH₁(SO.SII) by treatment with sodium amalgam (Limpricht, B. 18, 1473; Bauer, A. 229, 363). Yellowish amorphous mass, sl. sol. cold water, insol. ether.

Salts. -Na, A" raq. - CaA" 1, aq. -PbA". These salts are readily exidised (c.g. by KMnO, or I in K1) to the corresponding disulphonates. Conc. ammonic sulphido converts them into the di-thio-di-sulphonates. They are not reduced by sodium amalgam. Boiled with HCl the acid clots together, but conc. HCl. at 110° forms (2 p.c. of) an isomeric base.

p-Sulphi-benzene - azo - benzene - p-sulphinic

acid SO₂II.C₆H₄—N₂—C₆H₄SO₂H.

Azo benzene p-di-sulphinic acid. Prepared from $\begin{array}{c} SO_2Cl.C_cH_1-N_2-C_cH_1.SO_2Cl \ and \ Ba(SH)_2, \ or \\ from \ NaS.SO_2.C_cH_1.N_2.C_cH_1.SO_2.SNa \ by \ sodium \end{array}$ amalgam (Limpricht, B. 18, 1475; Bauer, A. 229, 369). The fres acid is ppd by HCl from its salts as a bulky yellow mass, sparingly solublo in water or alcohol. Clots together when heated with neids. Salts.—Na, A" 4aq.—BaA".

Sulpho-benzsne-azo-amido-ethane v. Sulpho-BENZENE-AZO-ETHYLAMINE.

p-Sulpho-bsnzene-azo-di-amido-henzoio acid $[4:1]C_6H_4(SO_3H)-N_2-C_6H_2(NH_2)_2CO_2H[1:4:2:6],$ Formod by the notion of p-diazo-benzene-sulphonio acid on s-di-amido-benzoic neid (Griess, B. 15, 2199). Needles or plates. Sl. sol. water, cold nicohol, and ether. Very unstable. Decomposes by boiling with water. On reduction it gives sulphanilic acid and (5:3:2:1)-tri-amidobenzoic acid.

Sulpho-benzene-azo-aniline-sulphonio *acid [4:1] $\vec{C_e}H_1(HSO_1) - N_2 - C_eH_1(HSO_1)NH_2$ [1:7:4]. Amido-azo-benzene disulphonic acid. Formed by sulphonation of p-amido-benzeno-azo-benzene-p-sulphonio neid (Griess, B. 15, 2187). Violst glistening noedles. Sol. hot water. Dyes silk and wool yollow. On reduction with tin and HCl it gives sulphanilio acid and p-phenylenediamino-sulphonic acid. BaA"7 aq: ornnge needles, sol. hot water.

m-Sulpho-henzene-azo-benzene-m-sulphonic acid [3:1] SO.H.C.H.-N.-C.H.SO.H [1:3].

Formation.- 1. From nitro-bensene m-sulphonic acid by treatment with sodium-amalgam (Claus a. Moser, B. 11, 762) or, better, with powdered zino and KOH (Mahrenholtz a. Gilbert, A. 202, 332).-2. One of the acids got by sulphonnting benzene - azo - benzeno at 150° (Janovsky, M. 3, 244).—3. From potassium mamido-benzene sulphonate and KMnO,

Monoclinie prisms, sl. sol. water and alcohol,

insol. ether.

Salts.—Na,A" 3½sq.: monoelinie erystals.— (NH.),A" 2aq. — CaA" 4aq. — BaA" 5aq. — PbA" ¼aq.

Amide [290°]. Prisms, sl. sol, water.

Ethyl ether Et.A". [100°]. Chloride C.H. (SO.CI).N., C.H. (SO.CI) [166°]. Acts upon cold conc. aqueous Ba(SH). thus: $C_a H_a(SO_a Cl) \cdot N_a \cdot C_a H_a(SO_a Cl) + 2BaH_aS_a = N_a(C_aH_a SO_a S)_aBa + BaCl_a + 2H_aS$ forming thiosulpho-benzene-azo-benzene-thio-sulphonioacid, part of which then decomposes according to the following equation: $N_2(C_0H_1,SO_2,S)_2Ba + H_2S = II_2N_2(C_0II_1,SO_2,S)_2Ba + S$ forming the barium salt of hydrazobenzsno di-thio-di-sulphonio acid (Bauer, A. 229, 353).

m-Snlpho-honzene-azo-benzene-p-sulphonio acid [4:1] $C_{\rm e}H_4(11{\rm SO}_3)$ - N_2 - $C_{\rm e}H_4(11{\rm SO}_3)$ [1:3]. Formed, together with the p-neid, by heating benzeus-azo-benzeus with 11,80, at 160° (Limpricht, B. 14, 1356; Rodatz, A. 215, 216), and by the exidation of n mixture of m- and ppotassium audio benzono sulphonate with KMnO. Uncrystallised syrup. On heating with dilute HCl to 150° it gives p- and mamido-benzene-sulphonio aoids (Limpricht, B. 15, 1155).

Salt's. $-K_2A'' 2_2^1$ aq; yellow needles, v. solwater. $-A_{\mathbb{S}_2}A''$.

Chloride [125°]: red needles.

 $A \, mide \, [258^{\circ}]$: slender yellow needles.

p-snlpho-bsnzene-azo-benzene - p - sulphonio acid. [4:1] SO₃H.C₆H₄-N₂-C₆H₄SO₃H [1:4].

Formation. -1. By oxidising muido-benzene-Permation.—I. by continuing minar-centeral-p-sulphonic acid with KMnO₄ (Laar, B. 14, 1928; Limpricht, B. 18, 1414).—2. Among the products of the sulphonation of benzene-azo-benzene at 160° (Limpricht, B. 14, 1856; 15,

1155; Janovsky, M. 3, 242).

Properties.—Ruby-red needles containing 2 or 3 aq (J.), or aq (L.). Melts at nbout 50° or, when dry, at about 150°. HClAq at 150° gives sulphanilio acid and other products.

Salts: K_2A'' $2\frac{1}{2}aq$: sl. sol. water. $-Na_2A''$. $-(NH_4)_2A''$. $-\Lambda g_2A''$. -CaA''. -PbA''aq. -CuA''6aq.

Chloride [222°]. Red needles. Amide [above 300°]; orangs plates or needles, sl. sol, hot water.

Di-snlpho-henzeae - azo-henzene - disnlphonis acid [5:3:1] C_aH₃(SO₃H)₂·N₂·C_aH₃(SO₃H)₂ [1:8:5]. From nitro-henzene-di-sulphonic acid, zino dust. nnd baryta (Reicho, A. 203, 64). Very deliques-cent crystals.—K,A'* 3aq.—Ba,A'* 5aq.

Di-snlpho-henzene azo-benzene - di-sulphonie acid [4:3:1] C₆H₃(SO₃H)₂·N₂·C₆H₃(SO₃H)₂ [1:3:4], From the corresponding nitro-benzone di-sulphonic neid, zinc dust, and baryta-water (Reiche, A. 203, 70). Salts. -K, A' 3aq. -- Ba, A' 4aq.

Chloride [58°]; radiating needles. Amide [222°]; white neodles.

p-Snlpho-henzene-azo-o-oresol [4:1] $C_0H_4(SO_0H) - N_2 - C_0H_4(CH_2)(OH)$ [1:8:4]. Formed by the action of diazo-benzene-p-sulphonio acid (by diazotising sulphanilio acid) on an alkaline solution of o-cresol (Nolting a. Kohn, B. 17, 364). Small reddish-brown needles. Sol. hot water. V. sl. sol. alcohol. On reduction with tin and HCl it gives sulphanilic acid and amido-o-cresol C₆II₃(CII₅)(NH₂)(OH) [I:5:2].

Salts. A'Na 2aq: yellow soluble plates.

A'2Ba 3aq : yellow tables, sl. sol. hot water.

p-Sulpho-benzenc-azo-m-crosol [4:1] $O_c \Pi_d(SO_3\Pi) - N_2 - C_d \Pi_d(G\Pi_d)(O\Pi)$ [1:2:4]. Formed by the action of diazobenzenc-p-sulphonic acid on an alkaline solution of m-cresol (Nölting a. Kolm, B. 17, 366). Small reddish-brown crystals with violet reflex. V. sol. V. sol. water and hot alcohol. Orango-yellow dyo stuff. On reduction it gives sulphanilic acid and amido-m-cresol C, II, (CII,) (NH2) (OII) [I:2:5].

Salts: A'Na: small yellow soluble needles. A'2Ba: yellow plates, v. sl. sol. cold water.

Sulpho-benzeno-azo-p-cresol

[4:1] $G_8H_1(SO_3H) = N_2 - G_8H_2(CH_3)(OH)$ [1:5:2]. C,H,CH,

C.H.(SO,H)-N.11 Formation .- 1. By the action of p-diazobenzeue-sulphonic acid on an alkaline solution of p-crosol .- 2. By sulphonation of benzeneazo-p-cresol (Nölting a. Kohn, B. 17, 355). Yellowish-brown plates with violet rotlex. Y. sol. water and hot alcohol. Dyes silk and wool

orange-yellow. On reduction with tin and HCl it yiolds sulphanilio acid and amido-p-cresol C_eH_s(CII_s)(NII_s)(OII) [1:3:4].

Salts: A'Na: soluble yellow plates. A'K 3aq.— A'_Mg 5aq.—A'_Ba: yellowish-brown tables, sl. sol. hot water.

p-Sulpho-benzeue-azo-\psi-cumenol

 $\{4:\hat{\mathbf{I}}\}\mathbf{C}_{a}\mathbf{H}_{a}^{\dagger}(\mathbf{HSO}_{a}) = \mathbf{N}_{2} + \mathbf{C}_{a}^{\dagger}\mathbf{H}(\mathbf{CH}_{a})_{3}\mathbf{OH}\{1:3:5:6:2\}.$ Formed by combining diago-benzene-p-sulphonic acid with \psi-cumenol [70]. -KA'2aq; orange needlos (Liebermann a. Kostanecki, B. 17, 887).

Sulpho-benzene - azo - ethylamine. Potassium salt. C.114(SO3K)-N2-CH(NH2).Cl13. From the potassium salt of the corresponding nitro-compound by reducing with ammonium sulpbide (Kappeler, B. 12, 2285). Silvery plates (from water); sl. sol. water, insol. Na CO.Aq. NaOHAq dissolves it with crimson colour.

m-Snlpho-bonzene-azo-(a)-naphthol

[8:1] $C_6H_4(HSO_3) - N_2 - C_{10}H_6 - OII$ [1:4]. pared by the action of an alkaline solution of (a)-naphthol on m-diazobenzono sulphonic acid (Griess, B. 11, 2197). Small greenish leaflets. Sl. vol. cold water and cold alcohol.

m-Sulpho-benzone-azo-(β)-naphthol [3:1] $O_0\hat{H}_1(HSO_2) - N_2 - \hat{C}_{10}\hat{H}_0(\hat{O}H)$ [1:2]

C10He. Prepared by the C.H.(HSO,)-HN. action of an alkaline solution of (3)-naphthol on m-diazobeuzene sulphouio acid (Griess, B. 11, 2197). Slender rod needles. V. sol. alcohol and water. BaA'25aq: yellowisb-rod scales. 81. sol. water.

p-Sulpho-benzene-azo-(a)-naphthol [4:1] C₆H₄(SO₃H) -N. -C₁₀H₆(OH) [1:4]. Tropæoline OOO, No. 1. From p-diazobenzeno sulphonio soid and an alkaline solution of (a)-naphtbol [Liebermann a. Jacobsen, A. 211, 61). Orange-

Its absorption spectrum is given by Hartley (C. J. 51, 184).

p-Sulpho-benzene-azo-(B)-naphthol

[4:I] C₆H₄(SO₅H)-N₂-C₁₆H₆(OH) [1:2] or

C10 He. Trop woline OOO No.2 C.H.(SO.H)-HN. From p-diazo-benzeno sulphonic acid and (β-) naphthol (W. v. Miller, B. 13, 268; Holmann, B. 10, 1378; Griess, B. 11, 2198). The absorption spectrum has been examined by Hartley (C. J. 51, 185).

p-Sulpho-bonzene-azo- (β) -naphthol sulphonlo acid [4:1] C₀H₁(HSO₃)—N₂—C₁₀H₃(HSO₃)OH. Prepared by the action of p-diazobenizene sulphonic acid on an alkaline solution of (9)naphthol sulphonic acid (Griess, B. 11, 2198; Stebbins, A. C. J. 2, 236). Yellowish red crystals. Excessively soluble in water. BaA"71aq: difficultly soluble orange microscopic needles.

p-Sulpho-benzenc-azo-(a)-naphthylamine [4:1] $C_0 \Pi_1 (\Pi SO_3) - N_2 - C_{10} \Pi_2 N \Pi_2 [1:1]$. From diazotised sulphanilic acid and (a)-naplithylamine (Griess, B. 12, 427). Brownish-violet needles, v. sl. sol. boiling water. Its acid solutions have a deep magenta colour (Griess's test for nitrous acid); its alkaline solutions are orange. On reduction with tin and HCl it gives sulphanilio

reduction wist on all and a grand and (1, 4)-naphthylone-diamine.

Salts: KA'3aq: brownish-yellow plates, sol. hot water.—BaA', 3aq: sparingly soluble brown needles (Griess, B. 15, 2190).

p-Salpho-benzene-azo- (β) -naphthylamine [4:1] $C_6 II_1(IISO_3) - N_2 - C_{16} II_6 NH_2$ [1:2] HN C.H. Formed by the

C.H.(HSO.)-HN. action of p-diazo-benzene-sulphonic acid on (β)-naphthylamine hydrochloride (Griess, B. 15, 2191). Small yellowish-red needles. St. sol. water, v. sol. hot alcohol, insol. ether. On reduction with tin and HCl it gives sulphanilio acid and (1, 2)-naphthylene-diamine. KA'7 aq: orange plates, sol. bot water.

Sulpho-benzene-azo-(a)-naphthylamine sulphonic acid G₆H₄(SO₂H).N. G₁₀H₃(SO₂H).NH₂(1:4:2]. Formed by the action of p-diazo-benzene-sulphonic acid on (a)-naphthylamine-sulphonic acid (Griess, B. 15, 2194). Needles or plates. Sol. water and alcohol, insol. ether, dyes silk and wool orange.—BaA" 71aq: red needles or plates, sol. hot water.—BaH2A'28aq: sparingly soluble violet-brown needles.

p-Sulpho-benzene-azo-(β)-naphthyl-phenyl- $C_0H_1(SO_3H) - N_2 - C_{10}H_0 \cdot NHC_0H_3$ amino C₆H₄(SO₃H).HN₂.C₁₀H₄ . Prepared by slowly

N.C.H. adding dry p-diazobenzene-sulphonio aoid (18 g.) to a solution of phenyl-(8)-naplithylamine (22 g. in glacial acetic acid (100 c.c.) at c. 50°, followed by finely powdered dry K.CO, (7 g.); the compound separates out in glistening red needles of the potassium-salt. It is a splendid scarlet dyestuff, but is very fugitive in light. potassium salt is easily solublo in water; when cold its solution solidifies to a transparent red jelly. IICl precipitates the free acid. By SnOl2 it is reduced to phenyl-o-naphthylone-diamine and sulphanilio acid. By boiling with dilnte mineral solds it is converted into naphthophenazine and sulphanilio acid:

O.H.(SO.H).N.C.,H.NHO.H. $C_{10}H_6 < \frac{N}{N} > C_4H_4 + C_6H_1(NH_2)SO_3H$. The Ba and Ca salts are crystalline insoluble pps. (Witt, B.

20, 572).

p-Sulpho-benzeue-azo-nitro-isobutaue $C_aH_1(HSO_s)-N_s$ - $C_sH_s(NO_s)$. Prepared by the action of p-diazobenzene-salphonic acid on an alkaline solution of nitro-iso-butane. -- KA'aq: orange-yellow needles. Soluble in alkalis to a red solution. Dyes silk orange (Kappeler, B. 12, 2288).

p-Sulpho-bonzene-azo-nitro-ethane

C.H. (HSO.) - N. - C.H. (NO.). Prepared by the action of p-diazobenzene-sulphonic acid on an alkaline solution of nitre-ethane. A'K: golden yellow leaflets, sparingly soluble in cold water, seluble in alkalis to a blood-red solution (Kappeler, B. 12, 2286).

n-Sulpho-henzene-azo-nitro-methano

C_aH₁(HSO₃) - N₂ - CH₂(NO₂). Prepared by the action of p-diazonenzene-sulphonic acid on an alkaline solution of nitro-methane, "KA' Baq: orange needles. Dyes silk erange (Kappeler, B. 12, 2286).

p-Sulpho-henzene-azo-nitro-propaue C₆H₄(HSO₃)-N₂-C(NO₂)(CH₃)₂. Prepared by the action of p-diazobenzene-sulphonic acid on an alkaline solution of nitro-isopropane. A'K: light-yellow leaflets. Has no dycing power. Insoluble in alkalis (Kappeler, B. 12, 2287).

p-Sulphobenzeue-azo-orciu

[1:1] C_aH₄(HSO₃)—N₂—C_aH₂(CH₃)(OH)₂. Small yellowish-red needles. Difficultly soluble in water. Prepared by the action of an alkaline solution of orein upon p-diazobenzene-sulphonic acid.-KA' 2aq (Griess, B. 11, 2196).

p-Sulpho-benzene-azo-o-oxy-beuzoic acid $[4:\hat{1}] C_a H_a(SO_a H) - N_a - C_a H_a(OH)(CO_2 H) [1:4:5].$ I'ron diazotised sulphanilic acid and an alkalino solution of salicylic acid. Golden needles; sl. sol, hot water (Griess, B. 11, 2196; Stebbins, B. 13, 716). BaH.A"2.

p-Sulpho-benzeno-azo-oxy-quinoline

 $\begin{array}{c} \textbf{p-Sulphotoshstate} \\ \textbf{(B,1)} & \textbf{(B,1)} \\ \textbf{C}_{6}\textbf{H}_{1}(\textbf{HSO}_{2}) - \textbf{N}_{2} - \textbf{C}_{6}\textbf{H}_{2}(\textbf{OH}) & \vdots \\ \textbf{N} : \textbf{CH} \end{array}$ Formed

by the combination of p-diazo-beuzene-sulphonic acid with (B. 4)-oxy-quinoline (Fischer a. Renouf, B. 17, 1642). Small needles. Orange dye.

p.Sulpho-henzene acc. phenol disulphonic acid C₂H₄(SO₃H). N.—C₄H₂(SO₄H). (OH). Fermed by heating azoxybenzene with fuming H.SO. Small soluble flat red needles with green lustre. On reduction it gives p amido-benzene-sulphonic acid and amide-phenol-di-sulphonic acid. Salts.—A"K3 3aq: yellow micr

Salts.—A"K₃ 3aq: yellow microscopic needles, easily soluble. Bromine-water gives tribromo-phenol .-- A"Ag : unstable red pp. --A"2Ba37uq: brewn crystalline pp. -A"2Pb3 13aq. Chloride: red crystalline powder [220].

Amide: yellew plates [260], sparingly soluble in alcohol (Limpricht, E. 15, 1297; Wilsing, A. 215, 234).

p.Sulpho-beuzeue-azo-xyleuol 4:1] C_sH_s(SO₃H) -N₂-C_sH_sMe_sOH [1:3:5.2]. Formed by combining diazobenzene-p-sulphonic

(Grevingk, B. 19, 148,. Dyes wool and silk a brownish yellow from au acid bath. On reduction it yields sulphauilio acid aud o-amido-mxylenol $C_0H_2Me_2(NH_2)(OH)$ [5:3:1:2]. Sulpho - carhoxy - heuzeue - azo - (β) -naphthol-

(a)-di-sulphonic acid

 $C_aH_a(CO_aH)(SO_aH) - N_2 - C_{10}H_1(OH)(SO_aH)_{20}$ Prepared by the action of m-diazo-sulpho-benzoic acid on an alkaling solution of (B)naphthol-(a)-di-sulphonic acid (Griess, 14, 2038). Orange needles or prisms. V. sol. water and alcohol, insol. other. Salts .-A" H.Ba. 3au: slightly soluble yellow needles. A Ba, 5aq: nearly insoluble red crystalline pp.

Sulpho-earboxy-benzene - azo - oxy - naplithoic acid $C_n\Pi_n(SO_nH)(CO_n\Pi) \cdots N_n \cdots C_{1n}\Pi_n(O\Pi)(CO_nH)$. Prepared by the action of diazosulphobenzoio acid on an alkaline solution of (a)-oxy-naphthoio acid (Griess, B. 11, 2199). Brown microscopic needles or leallets. Sl. sol. water.

Sulpho-naphthalene-azo- (β) -naphthol-disulphonic acid C₁₀H₁(SO₂H), N₀, O₁₀H₁(OH)(SO₂H)₂. Crimson dye (Stebbins, A. C. J. 2, 416).

Sulph :- to uene azo-tolnene-sulphonic acid (2:1:1) C.H.Mc(SO.H).N. C.H.Mc(SO.H) [1:2:4]. From potassium o-toluidine sulphonate (of Gerver) and KMuO, (Kornatzki, A. 221, 183). Small red prisms, grouped in tables, very soluble in water and in alcohol. K.A"; red plates grouped in clumps. BaA"aq. - CaA"3aq. BbA"aq.

Chloride. [218]. Red needles (from CaHa). Amide. [250]. Tables (from aqueous NH2). Sulpho-toluenc-azo-tolaene-sulphonic acid

[2:5:1] C_aH_aMe (SO_aH), $N_a.C_aH_aMe$ (SO_aH) [1:2:5]. Azo-toluene-disulphonic acid. From o-nitrotoluene sulphonic acid, zine dust, and KOHAq (Nunle, A. 203, 74); or from a toluidino sulphonic acid of Haydnek and KMnO, (Kornatzki, A. 221, 181). Salts. -BaA" 1aq. -K₂A" 2\aq. - CaA" 5aq. -PbA" 4aq.

Chloride. [220°]. Red prisms. Amide. [300°]. Red pewder.

Snlpho-toluene-azo-toluene-sulphonic acid [4:6:1] $C_nH_3Me(SO_3H).N_2.C_nH_3Me(SO_3H)$ [1:4:6]. From potassium p-toluidine sulphonate and KMnO, (Koruatzki, A. 221, 182). Salt.—BaA" 3aq.

Sulpho-toluene-azo-toluene-sulphonic acid 4:5:11 C.H.Mc(SO.H).N. C.H.Mc(SO.H) [1:4:5]. Frem p-nitro-toluene o-sulphonic acid, KOHAq, and zinc dust (Neale, A. 203, 80); or from potassic p-teluidine sulphonate and KMnO.

K.A" 3aq. - CaA" 3aq. - BaA" aq. - PbA" 2aq. Chloride. [194°]. Red Amide. [270°]. Wellow. Red crystals.

Fire - Sulpho - toluene - azo - tolueno - exo - sul phonic acid SO₃H.Cl) "C₄H₄N., C₄H₄CH₄SO₃H.
Formation.—1. From C₄H₄(NO₂).CH₄SO₃H

by boiling with zine dust and KOH or Ba(OH)2.-2. From C. H. (NIL.). CH .. SO , K and KMuO, (Mohr, 1. 221, 223). Salts.-K2A" 2aq: orange plates.-BaA" 1 aq. -Ag₂A" aq. Chloride. [149'].

Snlpho-xyleue-azo-di-bromo-uaphthol C_H_Me_(SO_H)-N_2-C_0H_Br_2(OH). From pdiazo-xylene sulphonic acid and di-bromo-(a)naphthol (Stebbins, jun., A. C. J. 2, 446). Sol. hot water, forming a searlet solution.

Sulpho-xylene-azo-(a)-uaphthol said with m-xyleuol C.H.Me.(OH) [1:8:4] C.H.Me.(SO.H)-N.-C.oH.OH. From p-diszoxylene sulphonic acid and (a)-naphthol (Stebbins, jun., A. C. J. 2, 446). Brown dye; sol. water.

Sulpho-m-xylene-aso-(3)-naphthol

[1:8:6:4] $C_aH_2Me_a(SO_3H)-N_2-C_{10}H_a(OH)$.

Formed by the action of diazo-m-xylene sulphonic acid (from m-xylidine eulphonio acid) upon an alkaline solution of (β)-naphthol (Nölting a. Kohn, B. 19, 139). Metallie green crystals. Sl. eol. cold water. Dyes wool and eilk from an acid bath a yellowish shade of searlet.

Salts.—A'Na×: red coluble platee.—A',Ba: sl. sol. hot water.

Sulpho-xylene-azo-(\$)-phenanthrol. From pdiazo-xyleno eulphonic acid and (\$)-phenanthrol (Stebbins, A. C. J. 2, 416). Reddish-brown dye.

Sulpho-xylene-azo-resorcin v. DI-OXY-BEN-MENE-AZO-XYLENE SULPHONIO ACID.

Snipho-xylene-azo-xylene sulphonio acid [2:4:5:1] C₀H₁Mc₂(SO₃H) - N₂ - C₀H₁Mo₂(SO₃H) [1:2:4:5]. Formed by exidising (1, 3, 6, 4)-xylidine eulphonio acid with dilute KMnO₄ (Jacobsen a. Ledderboge, B. 16, 194); er by reducing (6, 1, 3, 4)-nitro-xylene sulphonic acid with zine-dust and NaOH (Limpricht, B. 18, 2191). Orange plates; v. sol. water; el. eol. acids.

Salts. - K2A" 4aq. - KHA" 4aq. Chloride. [86]; red crystals. Amide. [174].

m-Thio-enlpho-benzene - azo-benzene - enlphinio acid [3:1] (IIS.SO.,)C.H., N., C.H. (SO.H) [1:3]. [below 100]. A solution of the barium thioeulpho-benzene-azo. (or hydrazo-) benzene-thieeulphonate gives, on evaporation, S and the salt of the present acid. This salt forms red crusts which are sparingly soluble in water, but are converted by boiling Na₂CO₃ into the soluble Na salt, whence HCl separates the free acid as a bulky flocculent pp. hardly seluble in water, but resinited by beiling with it. It is scluble in alcohel. Oxidised by KMnO, to N. (C.H., SO, K), Salts.—BaA" (dried at 140°).—K₂A".—Na₂A" xaq.—PbA" (dried at 130°).

Isomer.—Ammonia converts the acid into a brown amorphous base, isomeric with it (Limpricht, B. 18, 1472; Bauer, A. 229, 360).

m-Thio-eulpho-benzene-azo-benzene-m-thiosulphonic acid

[3:1] HS.SO, C, H, N, C, H, SO, SH [1:3].

f91°-93°]. From its salts by adding glacial acctio acid. A veluminoue yellow pp. insel. water or alcohol, and resinified wher boiled with them (Limpricht, B. 18, 1471; Bauer, A. 229, 358).

Barium salt .- BaA" 5aq. One of the products of the action of paric sulphydrato upon the chloride of sulpho-benzene-azo-benzene-sulphenic acid (q. v.). V. sol. hot water, el. sol. sulphenic acid (q. v.). V. sol. hot water, el. sol. cold water, nearly inseluble in alcohol. Yellow ammenic sulphide slowly converte it into the corresponding hydrazo-compound.

Na A" raq. 1te eolutions give amorphous ppe. with salts of Cu, Pb, Ag and Fe".

p-Thio-sulpho-benzene-azo-benzene thiosulphonio acid

[4:1] C_eH₄(SO₂SH)—N₂—C_eH₄(SO₂SH) [1:4]. Yellow amorphous colid. Sl. col. water and alcehel. Formed by the action of a saturated aqueous colution of Ba(SH), npon the chloride of eulpho-benzene-aze-benzene-eulphonie acid. Na.A" xaq: very coluble yellow warty oryetale. B. 19, 2488).

-BaA": yellow warty crystals, sol. hot water (Limpricht, B. 18, 1474; Baner, A. 229, 368).

p-Teluene-azo-aceto-acetio acid

[4:1] C₆H₄(CH₃)-N₂-CH(CO.CH₃).CO₂H. [188⁶]. Ethyl cther A'Et: [70²]; yellow needles. Formed by the action of p-diazo-toluene ohloride on an alcoholic colution of codio-acet acetio ether (Züblin, B. 11, 1419; Richter a. Münzer, B. 17, 1929).

p-Toluene-azo-acetone

[4:1] C₀H₁(CH₂)-N₂-CH₂.CO.CH₃. [115°].

Formation.-1. By heating p-toluene-azoaceto-acetic ether with a dilute alcoholic solution of NaOH .- 2. By heating p-toluene-azo-acetoacetio acid above its melting-point, CO2 being evolved (Richter a. Münzer, B. 17, 1929). Yellow needles. Sl. eol. water.

Toluene-azo-bromo-toluene

C₆H₄Me-N₂--C₆H₃BrMe. Bromo-azo-toluene. [136°] (P.); [138.5°] (J. a. E.). Formed by brominating p-tolucne-p-azo-tolucne (Petrieff, B. 6, 557; Janovsky a. Erb, B. 20, 363). Golden plates or needles. Reducee to a hydrazo- compound [119°].

Toluene-azo-chloro-toluene

[4:1] C.H.Me.N., C.H.MeCl [1:5:2], [97°], Formed by the action of cuprous chloride upon diazotised p-toluene-p-azo-toluidine (from p-toluidine); yield, 20 p.c. of theoretical. Brown platee. V. sol. alcohol, ether, and benzene (Montha, B.19, 3026).

p-Tolnene-azo-p-cresol [4:1] C₈H₄(CH₃)-N₂-C₆H₂(CH₃)(OH) [1:5:2] [113°]. Obtained by the action of p-diazoteluene chleride on an alkaline solution of pcresol. It is also formed by diazotising p-telueneazo-p-toluidine $C_gH_4(CH_3)-N_2-C_gH_3(CH_3)(NH_2)$ and beiling the product with water (Nölting a. Kohn, B. 17, 354). Reddish needles or yellow

tables. V. sol. ether, benzene, and hot alcohol.

Acetyl derivative [91°], yellow needles.

Benzoyl derivative [95°], emall yellow

o-Toluene-azo-ethyl-(\$)-naphthyl-amine [2:1] $C_8H_1(CH_3)$ -N₂- $C_{10}H_1(NHEt)$ [1:2]. [132°]. Formed by heating ethyl-(β)-naphthyl-nitrosamine with an acetic acid colution of o-toluidina (Henriques, B. 17, 2670).

p-Tolnene-azo-ethyl-(eta)-naphthyl-amine [4:1] C_eH₄(CH₄).N₂·C₁₀H_e(NHEt) [1:2]. [113°]. Fermed by heating ethyl-(β)-naphthyl-nitrosamine with an acetic acid colution of p-toluidine (Henriques, B. 17, 2670).

o-Toluene-p-azo-(a)-naphthol

[2:1] C₆H₄Me,N_aC_{1e}H₆(OH) [1:4]. (a)-Naphtho-quinone-o-tolyl-hydrazide. [146°]. Formation.—1. From o-diazo-toluene and

(a)-naphthel.-2. From (a)-naphthoquinone and o-tolyl-hydrazino.

Properties .- Red glietening needles. V. sol. alcohol, anetic acid, and benzene, lese readily in benzolino. HNO, cenverts it into di-nitro-(a)naphthol. With HCl and HBr it gives dark bluc metallio-glistening calte. Diesolvee in diluto NaOH ..

Methylether C. H. N. (OMe) 1930; reddish. brown glistening needles; easily soluble in ordinary solvents.

Ethyl cther C, H12N2(OEt) [94°]; red plates er dark thick noodlee (Zincke a. Rathgen, p-Toluene-p-aso-(a)-naphthol 4:1) $C_cH_cMen.N_x.C_{1c}H_c(OH)$ [1:4]. (a)-Naphtho-quinone-p-tolyl-hydrazide. [208°].

Formation.—1. From p.diazo-toluene and a)-naphthol.—2. From (a) naphthoquinone and

1-tolyî-hydrazine.

Properties.—Metallio - glistening dark - rod pangles. V. sol. acetone, aniline, and hot itrobenzene, sl. sol. alcohol, acetio acid, and enzene. Dissolvos in dilute NaOH. 11NO, converts it into di-nitro-(a)-naphthol. Not attacked by bromine in acetic acid solution. With mineral acids it forms salts which separate in bluish-green metallic-glistening plates. By heating with baryta-water it is rendered insoluble in alkalis.—B'HCk.—B'HBr.

Methyl ether C₁,H₁,N₂(OMc) [104°]. Ethyl ether C₁,H₁,N₂(OEt) [127°], large red crystals or red needles.

Acetyl derivative $C_1, H_1, N_2(OAc)$ [102°], fine yellowish needles (from benzoline) (Zincko a. Rathgen, B. 19, 2486).

o Toluene o azo-(a)-naphthol [2:1] $C_6H_1Me.N_r.C_{10}H_6(O11)$ [2:1] or

C₀H₄Me. HN₂ C₁₀H₄. (β) -Naphtho quinone-o-tolyl-hydrazide. [156°]. Formed by the action of o-tolyl-hydrazine upon (β)-naphthoquinone. Glistening red plates. Easily soluble in ordinary colvents. HNO₃ converts it into di-nitro-(α)-naphthol. Bromino gives a di-bromo-derivative [254°] (Zineke a. Rathgen, B. 19, 2492).

p-Toluene-o-azo-(a)-naphthol [4:1] $C_6H_4Me.N_2.C_{10}H_6(O11)$ [2:1] or

C.H.Me.N.H. C., bH. (β)-Naphtho-quinone-p-tolyl-hydrazide. [145°]. Formed by the action of p-tolyl-hydrazine npon (β)-naphtho-quinone. Red slender glistening needles. V. sol. alcohol, benzene, and acotio acid, sparingly in benzoline. By SnCl₂ it is reduced to (β)-amido-(a)-naphthol and p-toludino. HNO, converts it into di-nitro-(a)-naphthol. Bromino gives a di-bromo-derivative [236°] (Zincko a. Rathgen, B. 19, 2491).

p-Toluene-o-azo- (β) -naphthol [4:1] $C_0H_4Mc.N_2.C_{10}H_4(OH)$ [1:2] or

C₄H,Mo.N.H. C₁₀H₄. [135°]. Formed by combination of p-diazo-tolueno with (β)-naphthol. Thick red needles or tables. V. sol. alcolol, benzeno, acetio acid, and acetone. Insoluble in cold dilute NaOH. With acids it forms unstable salts. Bromino in acetic acid converts it into a di-bromo-derivative [190°]. HNO₃ gives di-nitro-(β)-naphthol (Zincke a. Rathgen, B. 19, 2490).

o-Toluene-o-azo-(β)-Laphthol [2:1] C_eH₄Me.N_{...}C₁₀H₆(OH) [1:2] or

Σ_eH₄Mc.N₂H C₁₀H₆. [131°]. Formed by sombination of o-diazo-toluene with(β)-naphthol. Fine red needles or plates. Insol. cold dilute NaOH. With acids it forms unstable salts. INO₂ converts it into di-nitro-(β)-naphthol. Bronine forms a mono-bromo-derivative [167°]. Zincke a. Rathgen, B. 19, 2491; Fischer, B. 20, 580).

p-Toluene-azo- (β) -naphthol disulphonic acid ${}^1_4H_4Me-N_2-C_{10}H_4(OH)(SO_2H)_2$. From sodium-3) naphthol disulphonate and p-diazo-toluene

nitrate (Stebbins, A. C. J. 2, 236; C. N. 42, 44). Red leaflets, v. sol. water. Scarlet dyo. The corresponding o compound dyes yellower, the m-compound, redder.

p-Tolnene-azo-(a)-naphthylamine

[4:1] C₆11₁(Cl1₄)—N₂—C₁₆H₆NII₂ [1:4]. [145°]. Prepared by the action of p-diazo-toluene sulphate on (a)-naphthylamine (Weselsky a. Benedikt, B. 12, 229). Red leaflets; insol. water.—B'.1I₂SO, 3aq: steel-bluo needlee.
o-Toluene-azo-nitro-ethane

[2:1] C₈H₁Me-N₂-CH(NO₂), CH₃. [88°]. From o-diazo-toluene nitrato and potassimu nitroethane (Barbieri, B. 9, 387). Unstable orange needles.—NaA': golden spangles.

p-Toluene-azo-nitro-ethane. [133°]. Propared like the preceding (B.). Orange prisms with steel-blue lustro. Its alkaline solutione are

deep red.

Toluene-azo-nitro-toluene

C_ol1,Mc-N₂-C_oH₃(NO₂)Me. Nitro-azo-toluens [114*]. Among the products of the nitration of toluene-azo-toluene dissolved in glacial acetic acid (Janowsky a. Erb, B. 20, 363). Orange monoclinic needles (from 90 p.c. alcohol).

Toluene-azo-nitro-toluene. [76°]. From tolueneazo-toluene and HNO, (S.G. 14) (Petrieff, B. 6,

o-Tolnene-azo-orcin

C₆H₄Me-N₂-C₆H₂Me(OII)₂ [203°-206°]. From o-diazo-toluene and orein (Sciellilone, G. 12, 223). Red-brown crystale.

p-Tolnene-azo-thymol sulphonic acid Cp-Tolnene-azo-thymol sulphonic acid Cp-Tolnene-azo-thymol sulphonic Dp-Tolnene-chloride on sodium thymol-sulphonic — A'Na:

chloride on sodium thymol-sulphonate.—A'Na: slender yellow needles; sol. alcohol and hot water, almost insoluble in cold water (Stebbins, B. 14, 2795).

o-Toluene-o-azo-toluene

[2:1] $Me.C_0H_4-N_2-C_0H_4$. Me [1:2]. o-Azo-toluens [55°].

Preparation.—1. By distilling o-nitro-tolueno with alcoholic potash; or by reducing it with zinc-dust and alcoholic NaOH (Schultz, B. 17, 497). Cannot be prepared by reducing o-nitro-toluene in alcoholic solution with sodium-amalgam (Perkin).—2. From o-toluidine and KMnO₄ (Hoogeworff a. van Dorp, B. 11, 1203).

Properties.—Dark red trimetrio prisms; a:b:c=2:225:1:1708, Volatile with steam. Givos a mono-nitro derivativo [c. 67°], a di-nitro-derivativo, [142°], and a tri-nitro-derivativo that

decomposes before melting (Petrieff).

p-Toluene-p-azo-toluene. p-Azo-toluene

[4:1] Mc.C.H.—N.—C.H.Mo [1:4]. [144°].
Formation.—From p-toluidine and CrO, in glacial acetic acid; or by treating a solution of p-toluidine in chloroform with bleaching-powder (R. Schmitt, J. pr. [2] 18, 198). Or by oxidising p-toluidine with H.O. (Leeds, B. 14, 1382), or benzoyl peroxide. Cannot be prepared by distilling p-nitro-toluene with alcoholic potash

(Perkin), for by such treatment a red condensation product is obtained which on further reduction gives di-ami lo-di-phonyl-ethylene [227°]. (Bender a. Schultz, B. 19, 3237).

Preparation.—p-Nitro-toluene (20g.) in alcohol is treated with sodium-anualgam added gradually, the mixture being frequently cooled. The brown solid that separates is crystallised from glacial acetic acid (Perkin, C. J. 37, 554, cf. Jaworsky, J. pr. 94, 223; Werigo, Z. 1864, 610; Alexejeff, Z. 1866, 269; Melms, B. 3, 549; Schultz, B. 17, 472).

Properties. Red trimetric needles. V. sol. alcohol. alcohol and ligroin, sl. sol. alcohol. Slowly reduced to hydrazo-tolucne by ammonium sulphide. In alcoholic solution it is reduced by SnCl., and HCl to tolidine [91°]. (S.). Nitric acid forms a mono-nitro-derivative, [76°], a dinitro-derivative [110°], and a tri-nitro-derivative [201°] (Petricff).

o-Toluene m-azo-toluene

[2:1] $C_n H_4(CH_3)$ $N_2 - C_n H_4(CH_4)$ [1:3]. Obtained by diazotising o-tolucue-azo-o-toluidine (from o-toluidine) and treating the diazo-compound with alcohol (Schultz, B. 17, 470). Red oil. Volatile with steam. V. sol. alcohol and ether. By SnCl₂ and HCl in alcoholic solution it is convorted into an unsymmetrical tolidine.

m-Toluenc.p-azo-toluene

[4:1] C_aH₄(CH₃) -N₂-C_aH₄(CH₃) [1:3]. [58°]. Formation,—1. By the action of zinc-dust and alcohol upon c-diazo-toluene-azo-toluene,—2. By the action of Ag₄O and alcohol upon the compound C₄H₁₁N₄ the reduction-product of c-diazo-toluene-azo-toluene (Zincke a. Lawson, B. 19, 1458). Brownish-red plates. V. sol. alcohol, other and benzene.

o-Toluene-azo-o-toluidine [2:1] $C_eH_1(CH_3)-N_2-C_eH_3(CH_3)(NH_2)$ [1:3:4] [100°]. Formed by passing nitrons acid gas into o-toluidine (Nietzki, B. 10, 662). Trimetrio crystals, a:b:c=10116:1:13268. Heated with aniline hydrochloride and alcohol at 160° it

forms a red dye resembling saffranin, Salts.—B'HCl: orange tables.—B',H,PtCl_s. Acctyl derivative

C₇H₇—N₂—C₇H₈(N11Ac). [185°]. Slender red needles, v. sol. alcohol (Schultz, B. 17, 469).

m-Tolueno-azo-m-tolnidino

[8:1] C_uH₁Mc—N₁—C_uH₃Me(NH₂) [1:2:4]. [80°]. Formed by treating an alcoholic solution of mt-tolnidine with nitrons acid gas (Nietzki, B. 10, 1155). Golden needles.—B'HCl.—B'₂H₂PtCl₃. Gives p-te³yleno diamino, [64°], on reduction.

p-Toluone-azo-o-toluidine

[4:1] C_oH₄Me-N₂. C_oH₃Me(NH₂) [1:3:4], [128°]. From p-diazo-toluene toluido and o-toluidine bydrochlorido (Nictzki, B. 10, 832). Gives p-tolylone-diamine, [61°], on reduction. Heated with aniline hydrochloride it forms a violet dyc. Salte,—B'HCl,—B'₂H₂PtCl₆.

p-Talneuc-azo-m-toluidine

[4:1] C_eH₁Me—N₂—C_eH₈Me(NH₂) [1:2:4]. [127°]. From p-diazo-toluene toluide and m-toluidine hydrochloride in alcoholic solution (Nietzki, B. 10, 1150). Large yellow plates. Gives p-tolylene-diamine [64°] on reduction.

Salts.-B'HCl.-B',H,PtCl.

p-Toluene.axo.p-toluidine [4:1] C₂H₁Me—N₂—C₂H₂Me(NH₂) [1:5:2] or NH | C₂H₁Me. [119°]. o.Amido.azo-

C₆H₄Me.N₂H Tolnene-hydrazimido-toluene. toluene. pared by heating p-diazo-toluene-p-toluide (diazo-amido-tolnene), dissolved in 5 or 6 times its weight of melted p-toluidine, with p-toluidine hydrochloride (1 mol.) at 65° for 12 hours. Orange-red glistening needles. V. sol, hot alcohol, acetic ether, and benzenc. On reduction it gives p-toluidine and tolylene-o-diamine. CrO: oxidises it in acetic acid solution to tolueneazimido-toluene C,H,- N,- C,H, (Zincke, B. 18, 31-12). Heated with p-tolnidine hydrochloride and p-tolnidine at 100° it gives a body C. H. N. analogous to azophenine which forms flat red needles. Heated to a higher temperature dyostuffs of the induline series are formed. It is converted into curhodino $C_{17}H_{13}N_3$ by heating with (a)-naphthylamino hydrochloride (Witt, C. J. 49, 393). The salts of o-amido-azo-p-toluene are yellow in the solid state, but dissolve to green solutions .- B'HCl: slender light-yellow needles.

Acetyl derivative. [157°]; yellow felted needles.

Benzoyl derivative. [135°]; orangoyellow needles (Witt a. Nölling, B. 17, 77). Disulphonic acid C₁₄H₁₈N₃(SO₅H)₂ Formed by sulphonating with funding H₂SO₄ (N. a. W.). Greyish white needles. Is a yellow dyestuff of redder shade than 'acid yellow.'— BaA" 4aq: brownish-red crystalline powder.

p-Tolnene-azo-tolylene-diamine
[4:1] C_aH,Mo-N_z—C,H,Mc(NH.)₂ [1:3:4:6],
[183°]. From p-diazo-toluene nitrate and tolylene-m-diamine (Hofmann, B. 10, 218). Oranga needles, v. sol. alcohol, insol. water.—B"HCl.—

B"_zH_PtCl_n.

Xylone-azo-(β)-naphthol-(β)-sulphonic acid,
Diazo-xylene does not combine with Rumpf's 'a':
sulphonic acid of (β)-naphthol in dilute alkaline
solution, although some other diazo-compounds
(such as diazo-benzene) do combine with it under
the same conditions. II, however, the solution
is very concentrated, the combination with
diazo-xylene takes place. The product forme
red needles, dissolvos in H_zSO_z with a red
colour, and dyes wool a somewhat yellower shada
than the compound from Schäfer's 'β'-acid
(Schultz, B. 17, 461).

Xyleue-azo-thymol-snlphonio acid

C₂II₂(CH₃)₂-N₂-C₃H(CII₃)(C₃H₇)(HSO₃)OH. Slender yellow needles. Prepared by the action of diazo-xylene eliloride on sodium thymolsulphonate,—A'₂Ba: small yellow needles or plates (Stebbins, B. 14, 2795).

Xylone-azo-xylene

Xylone-azo-xylone

\$C_0H_x(CH_x)_x - N_x - C_0H_x(CH_x)_x.\$ Azo-xylone [126°-corr.] Formed by roduction of nitro-m-xylone with sodium-amalgam or with zinc-dust and alcoholio NaOH; very small yield. A better yield is obtained by oxidation of xylidine with alkaline polassium ferricyanide (Werigo, Z. 1864, 723; 1865, 312; Samonoff, Bl. [2] 39, 597; J. R. 1882, 327; Schultz, B. 17, 476). Red needles. Sol. hot alcohol. It does not appear to give a dixylyl base by treatment with SnCl₂ and HCl in alcoholie solution.

xylene. [78°]

Preparation.—Diazo-m-xylene-m-xylide, prepared by adding a solution of 1 mol. of sodium nitrite to a mixture of 1 mol. of its hydrochloride, is dissolved in m-xylidino and gently warmed for a long timo with about 5 p.c. of m-xylidine hydrochloride. The inixture is then acidified with dilute HCl, the precipitated hydrochloride is filtered off, washed with water, alcohol, and ether, basified, and crystallised from alcohol or benzene; the yield is 70 p.c. to 80 p.c. of theoretical. Orange plates. V. sol. benzene, and hot alcohol, v. sl. sol. water.

Reactions.—On reduction it yields m-xylidine and m-xylylene-o-diamino $C_6H_2Me_2(NH_2)_2$ [1:3:5:6].

Salts.—B'HCl: yellow crystalline powder, dissolves sparingly in alcohol with a green colour, soluble in phenol with a splendid green colour (Nölting a. Forel, B. 18, 2682).

m-Xylene-p-aze-m-xylidine
[2:6:1] C_LL_Mc_-N_2-C_LL_Mc_(NH_-) [1:3:5:4]
[78°]. Yellow plates. Easily soluble in alcohol
and benze-re. Prepared from m-xylidino
C_bL_Mc_-NH__[1:3:2] by the same method as that
described under m-xylene-o-azo-m-xylidine

Salts. --1'_H_CLPtCl_; red crystalline powder. The hydrochloride dissolves in phenol or alcohol with a red colour (Nölting a. Forel, B. 18, 2684).

m-Xylene-p-aze-m-xylidino
[3:5:1] C_aH₃Me_z-N_z-C_aH₄Me_z(NH₂) [1:2:6:4].
[95°]. Prepared from m-xylidine C_aH₄Me_zNH₂
[1:3:5] by the same method as that described for m-xylene-o-uzo-m-xylidine. Yellow plates. On roduction it gives symmetrical m-xylylene-p-diamine C_aH₄Me_z(NH₂), [1:3:2:5]. The hydrochloride dissolves in phenol with violet-red colour (Nötting a. Forel, B. 18, 2684).

o-Xylene-p-aze-o-xylidine
[2:3:1] O,H,Mu,-N,-C,H,Me,(NH,) [1:2:3:4].
[111?]. Frepared from σ-xylidino C,H,Me,-NH,
[1:2:3] by the same method as that described under m-xylene-o-azo-m-xylidene (Nölting a. Forol, B. 18, 2684). Glistening yellow plates (from alcohol or benzene). On reduction it yields σ-xylidino and σ-xylyleno-p-diamine C,H,Me₂(NH,)₂[1:2:3:6]. The hydrochloride dissolves in phenol with a red colour.

m-Xylene-p-azo-p-Xylidine [2:4:1] $\mathbf{O}_0\mathbf{H}_2\mathbf{M}\mathbf{e}_2-\mathbf{N}_2-\mathbf{C}_0\mathbf{H}_2\mathbf{M}\mathbf{e}_2(\mathbf{N}\mathbf{H}_2)$ [1:2:5:4]. [111°]. Red plates.

Preparation: 50 c.o. of a solution of sodium nitrite containing 227 grms. NaNO, per litro are added to a mixture of 20 grms. of p.xylidino and 26 grms. of hydrochloride of m.xylidino C.H.Mc.(NII.) [1:3:4]; the diazoamide so formed is dissolved in 20 grms. of p.xylidine and gently warmed with 4 grms. of p.xylidene hydrochloride.

Reactions.—On reduction it yields rs. xylidino and p. xylylene-p. diamine C. H. Me. (Nll.), [1:4:2:5]. The hydrochloride dissolves in phenol with a red colour (Nictzki, B. 13, 470; Nölting a. Forel, B. 18, 20:80.

p-Xylene-p-azo-p-xylidine [2:5:1] O₄H₄Me₂—N₂—C₆H₄Me₂(NH₂) [1:2:5:4]. [150°]. Red plates (from alcohol). Prepared from p-xylidine $C_aH_aMe_2(NH_a)$ [1:4:5] by the same mothod as that described under m-xylene-o-azo-m-xylidine. On reduction it yields p-xylidino and p-xylyleno-p-diamine $C_aH_aMe_a(NH_a)_2$ [1:4:2:5]. The hydrochloride is red, and dissolves in phenol with a violet-red colour (Nölting a. Forel, B. 18, 2085).

o-Xylene-o-azo-o-xylidine

[3:4:1] C₀H₁Me₂—N₂—C₀H₂Me₄(NH₂) [1:3:4:6] or NH C₀H₂Mo₂—N₂H

C₀H₂Me₂ [179°]. Yellow

C₀H₃Mo₂(NH₂) [1:2:4] by the same method as that described under m-xylene-o-azo-m-xylidine. On reduction it gives rise to o-xylidine and o-xylylene-o-diamine C₀H₂Mo₂(NH₂); [1:2:4:5]. Its hydrochlorido dissolves in phenol with a green

colour (Nölting a. Forel, B. 18, 2685).

DIAZO-COMPOUNDS. A class of bodies formed by the action of nitrous acid upon primary amido-compounds: X.N.[11]+O.N.OH=X.N._OH+H.O. They contain a pair of nitrogen atoms (Fr. azate) which are united to only one hydrocarbon radicle, whilst in thoazo-compounds the N.g group is united to two hydrocarbon radicles X.N.Z. The diazo-radicles X.N.Z. ennot of course exist in the free state, but they occur as hydrates X.N._OH, chlorides X.N._CH, amides X.N._NHR, &c. For the sake of convenience reactions will usually be represented in this article as taking place with the hydrates.

The diaze-salts X.N.A may be regarded as derived from the salts of amines X.NI₄A by the displacement of H₃ by N. This may take place in two ways. According to Kekulé's view, which is that most generally adopted, both nitrogen atoms are trivalent: X.N.N.A. On the other hand, Blomstrand (Chemie der Jetstzeit, p. 272, and B. 8, 51) assumes that the nitrogen attached to the carbon is pentavalent: X.N.A.;

Strecker (B. 4, 786) and Erlenmeyer (B. 7, 1110) also concur in this view. The reduction of diazo-compounds to hydrazines, which have the undoubted constitution X.NH.NH.₂, E. Fischer (A. 190, 67) regards as a proof of the correctness of Kekulé's formula, since a body of the constitution X.N.A would, he considers, give on N

reduction X.NH₂. Crnm Brown, however, has

pointed out in a private communication that this argument is fallacious, since the product of the reduction is not a hydrazine itself but a hydrazine salt, and X.N.A, by adding II 2 to each

N, would give the hydrazine salt X.NH.A. He

considers the pentad N in the salts of hydrazines is most probably that connected to the hydrocarbon nucleus, in which case to explain their formation by adoption of Kekula's formula would necessitate a shifting of the acid from one N to the other. The strongest argument against Kekula's formula is that it represents diazo-salts, by not containing pentad nitregen, as differently constituted to the salts of all other nitrogen bases. On the other hand, the formula X.N.A would necessitate a rearrange-

ment of the molecule in the formation of azocompounds which undoubtedly have the con-

stitution X.N:N.Y.

The simplest member of the series H.N. OII should be formed by the action of nitrous acid upon NII, but it has not yet been obtained, probably by reason of its extreme instability. The best-known diazo-compounds are those derived from aromatic amines and amido-compounds, some of which are tolerably stable bodies. No diazo-compounds have at present been obtained from fatty amines, for, like the first member of the series, II.N. OII, they are so unstable that they are probably scarcely capable of existence, and at once break up into the alcohol sud N2. The only known fatty diazo-compounds are a few which have lately been prepared from fatty amido-others (e.g. glycocoll). In their constitution they differ from the alomatic hydrates by containing a moleculo of water less: (EtO_C)CHI_N_.OH -II_O = (EtO_C)CH:N_.

A Aromatic diazo-compounds. The disstitution they differ from the aromatic diazo-

A. Aromatic diazo-compounds. The discovery of these bodies, and a large portion of our knowledge concerning them, are due to P. Griess (A. 106, 123; 113, 201; 117, 1; 120, 125; 121, 257; 137, 39; &c.), who, in a series of classical researches, opened up a field of investigation which in a few years has produced more discoveries of scientific interest and practical utility than almost any other branch of

organic chemistry.

Formation.—1. By the action of nitrous acid, or any compound readily forming nitrous acid (e.g. NOCl, NOBr, SO₂(OH)(NO₂), zinc-dust and HNO₃, &c.) upon salts of primary amines.—2. By oxidation of primary hydrazines (F. Fischer,

A. 190, 97).

Preparation.—The details vary very much with individual eases and the purposes for which the diazo- compounds are required. The amine can be dissolved in water, alcohol, acetie acid, HCl, H.SO,, &c., and can then be treated with nitrous acid gas, sodium nitrite, or a nitrous ether. When required in the solid form, a common method is to mix the nitrate of the amino with a little water, cool in a freezing-mixture, and saturate with N₂O₄ gas; the diazonitrate is then ppd. by addition of alcohol and ether. Diazo-compounds can also be isolated from their aqueous solutions by ppn. as platinoehlorides, perbromides, piorates, sulphites, &c. When the diazo-compound is required for a subsequent reaction it is seldom necessary to isolate it, but the compound can be prepared under the conditions suitable to the second reaction. For instance, when the diazo-compound is to be conjugated with an amine or phenol to form an azo- compound, the amino is usually dissolved in water containing 2 mol. of HCl for each NII, group, cooled by addition of ice, and mixed with an aqueous solution of sodinninitrite (1 mol. to each NH.). The solution of the diazo-chloride thus prepared can be at once treated with a solution of the phenol or amine.

The diazetisation of simple amines, in not too dilute solutions, usually takes place quantitatively, and the reaction is tolerably rapid. For instance, the diazetisation of aniline in a 10 p.o. solution is so complete within an hour

that it forms the most accurate method of estimating nitrous said or aniline (Green a. Ridesl, C. N. 49, 173; Green a. Evershed, S. C. I. 1886, 633). The greater the molecular weight of the amine the slower and less complete is the diazotisation. The diazotisation of heavy amidobodies is facilitated by the presence of a very largo excess of mineral acid, using as little water as possible. Alcohol in many oases appears to have a contrary effect. Amido-groups cannot be diazotised unless combined with an acid: thus if the ordinary hydrochlorido of pplienylene diamine C, II, (NH.) (NII, Cl) is treated with HNO, only one NH, group is diazotised; but if a large excess of HCl is employed so that C.H.(NH,Cl), is present, both NH, groups are diazotised. The final products of the action of nitrous acid upon the mono-acid salts of diamines vary with the constitution of the latter. Thus o-phenylene diamine gives azimidobenzene:

 $\mathbf{C}_{e}\mathbf{H}_{4}(\mathbf{NII}_{2}).\mathbf{N}_{2}.\mathbf{OH} - \mathbf{\Pi}_{2}\mathbf{O} = \mathbf{C}_{e}\mathbf{H}_{1} \underbrace{\mathbf{N}}_{N}\mathbf{N}\mathbf{H}_{2}.$

m - Phenylene diamino gives tri · amido · azobenzeno, tbus: $C_vH_1(\mathrm{NH}_2)$, $N_v.OH + C_vH_1(\mathrm{NH}_2) = C_vH_1(\mathrm{NH}_2)$, $N_v.C_vH_1(\mathrm{NH}_2)$, H_1 . Whilst the diazo · coinpound $C_vH_1(\mathrm{NH}_2)$, $N_v.OH$ [1:4], from p-phenylene diamine, does not undergo any further transformation. The di-amido-benzoio acids react with nitrous acid in an exactly similar manner according as the NH $_2$ groups are 0, m, or p to each other (Griess, B, 17, 607).

Properties.—The diazo- salts are in general very unstable crystalline solids. When dry they often decompose with detonation, by heat or percussion. Their solutions slowly decompose at the ordinary temperature, mere quickly on heating, with evolution of nitrogen. The hydrates are even more unstable than the salts, and have scarcely ever been isolated. The stability is increased by substitution in the nucleus; thus diazo-benzone sulphonio acid is more stable than diazo-bonzene. The diazo-derivatives of substituted phenols and of o- and p- sulphonic acids usually occur in the form of anhydro-compounds, e.g.

C₆H₁₂Br₂
$$\stackrel{N_2.OH}{OH}$$
 - H₂O = C₆H₁₂Br₂ $\stackrel{N_2}{\downarrow}$, and C₆H₄ $\stackrel{N_2.OH}{SO_3H}$ - H₂O = C₆H₄ $\stackrel{N_2}{\downarrow}$. In a similar

manner o-amido-diazo- compounds form inner amides (e.g. azimido-benzene, v. supra).

Reactions.—The diazo-compounds are extremely prone to undergo reactions; they play a most important part in organic syntheses and the determination of the constitution of isomeric aromatic compounds, by serving as an intermediate stage by means of which NII₂ groups can be replaced by H, OH, Cl, Br, I, F, CN, SH, NO₂, &c. Their power of combining with amines and phenols to form azo-compounds renders them of great technical importance for the production of colouring-matters, for which purpose they are prepared in large quantities. The majority of their reactions consist in the evolution of N₂, and its replacement by the atom or group (Cl, OH, &c.) previously united to it.

1. By heating the aqueous solution nitrogen ! Is evolved, with formation of the corresponding phenol: X.N2.OH = X.OH + N2. The best method is to dissolve the amine in a considerable oxcess of dilute H₂SO₄, diazotise by adding NaNO₂ to the iced solution, and finally heat to boiling.

2. When heated with strong alcohol the normal reaction appears to be the replacement of the N₂ group by OEt, with formation of ethoxy-compounds (Wroblewski, Z. 6, 164; B. 17, 2703; Haller, B. 17, 1887; Hofmann, B. 17, 1917; Remsen, Am. 8, 243; B. 18, 65; Hayduck, A, 172, 212; Zander, A. 198, 25; Heffter, A. 221, 352; Paysan, A. 221, 212, 363; Mohr, A. 221, 222; Hesse, A. 230, 293).

3. Under certain circumstances, at present undetermined, the reaction with alcohol takes a different course, resulting in the substitution of H for the N2 group, with production of the corresponding hydrocarbon together with aldehyde:

 $X.N_{..}O11 + C_{..}H_{..}O = X.11 + C_{..}H_{..}O + N_{..} + H_{..}O$ (Griess). (a) The amido-compound is treated with a solution of nitrous acid in absolute alcohol, warmed till nitrogen comes off freely, allowed to cool, resaturated with N2O3, and the operation repeated until but little gas is evolved on heating (Neville a. Winther, C. J. 37, 452). (b) The amido-compound is dissolved in a considerable excess of cone. H.SO, the solution diluted with a small quantity of water is cooled in a freezing-mixture, and the necessary quantity of solid sodium nitrite added. When diazotised the solution is poured in a thin stream into two or three times its bulk of alcohol; the mixture becomes warm enough to complete the reaction without further heating (McIdola, C. J. 1885, 507).

4. Mercaptan, when heated with diazo-compounds at 170°, heliaves similarly to alcohol in reaction 3, causing the displacement of N2 by hydrogen with simultaneous formation of diethyl-di-sulphide (Schmitt a. Mittenzwey, J. pr.

5. The displacement of the N, group by H is also effected by reduction to the corresponding hydrazine (q. v.), and treatment of this with $CuSO_4$ or Fe_2Cl_4 (B. 18, 90).

6. Reduction of a diazo-chloride with excess of SnCl, also effects the displacement of N. by H: $X.N_{u}Cl + SnCl_{u} + Il_{u}O = X.II + SnOCl_{u} + HCl + N_{u}$ A diluto aqueous solution of the diazo-chloride is treated with an excess of SnCl, at 0°, and finally heated for two hours with an inverted condenser; the yield is good (Effront, B. 17, 2329; Gasiorowski a. Wayss, B. 18, 337).

7. By treatment of a cold solution of a diazocompound in cone. HCl with (2 mols. of) SnCl,, the corresponding hydrazine (\hat{q}, v_i) is produced: $X.N_m.Cl + 2SnCl_k + 3HCl = X.N.H.N.H. + 2SnCl_k$.

8. The reduction of the sulphites of diazocompounds with SO₂, or with zinc-dust and acetic acid, also gives rise to hydrazines.

9. By heating with dilute HNOs nitrated phenols are obtained (Nölting a. Wild, B. 18,

10. The platino-chloridos on distillation with dry Na.CO, yield the corresponding chloro-derivatives: (X.N₂.Cl)₂PtCl₄ = 2X.Cl + 2N₂ + PtCl₄.

11. The replacement of N2 by Cl is also effected by boiling the diazo-compound with fuming HCl in large excess: X.N₂.Cl = X.Cl + N₂ can be converted into sulphonic acids by oxida-

(Griess, B. 18, 960; Gasiorowski a. Wayss, B. 18, 1936).

12. The same replacement is most readily effected by treating the aqueous solution of the diazo-chloride with cuprous chloride, which appears to act by intermediate formation of an addition product R.N. Cl. Cu.Cl., (a) A 10 p.c. solution of Cu.Cl., is prepared by adding 100 pts. of cone. HCl and 13 pts. of copper turnings to a hot solution of 25 pts. of crystallised CuSO, and 12 pts. of NaCl, boiling till decolourised, and making up the weight to 203 pts. with conc. HCl. A dilute HCl solution of the diazo- compound is allowed to run slowly into the above solution (about 5 times the weight of the amine used) licated nearly to boiling; the product, if volatile, is distilled with steam, or it is separated and purified by crystallisation. (b) In most cases instead of separately diazotising the amine, its solution in dilute HCl can be mixed with about 5 pts. of the 10 p.c. Cu₂Cl₂ solution, and a solution of the calculated quantity of NaNO, run into the nearly boiling mixture (Sandmeyer, B.

17, 1633, 2650; Lellmann, B. 19, 810). 13. The perbromides (g. v.) of diazo-compounds, on heating by themselves, or with dry Na CO, or best by boiling with glacial acetic avid, yield bromo derivatives: X.N₂.Br₃ = X.Br + Br₂ + N₂ (Neville a. Winther, C. J. 37, 452).

14. The replacement of N, by Br is also effected by boiling the diazo compound with luming HBr in large execss: X.N.Br - X.Br + N2 (Griess, B. 18, 960; Gasiorowski a. Wayss, B. 18, 1936).

15. The same replacement is most conveniently effected by means of cuprous browide (cf. reaction 12). A solution of 125 pts. of crystallised CuSO, (\frac{1}{2} mol.), 360 pts. of KBr (3) mols.), 800 pts. of water, and 110 pts. of cone. H.SO, (1 mol.), is boiled with 200 pts. of copperturnings till decolourised. The amino (1 mol.) is then added, and into the mixture, heated nearly to boiling, is slowly run a solution of 70 pts. NaNO. (1 mol.) in 400 pts. of water (Sandmeyer, B. 17, 2650; 18, 1492).

16. By boiling diazo-compounds with aqueous HI tho N₂ group is replaced by I forming iodo-compounds: X.N₂I-XI+N₂ (Griess, B. 18,

17. By boiling with HF the N. group is replaced by F giving fluoro-compounds: X.N. F= XF + N₂ (Griess, B. 18, 960; Paterno a. Oliveri, G. 12, 85; 13, 533; Wallach, A. 235, 255).

18. By heating diazo- salts with Cu.(CN), the N. group is replaced by CN (cf. reactions 12 and 15). The nitriles so formed can be converted iuto carboxylie acids by saponification, so that by means of this reaction an NH₂ group can be replaced by CO₂H. 28 pts. of KCN (96 p.c.) are added to a hot solution of 25 pts. of crystallised CuSO, in 150 pts. of water; into this solution, heated to about 90°, is slowly run an aqueous solution of the diazo-chloride. If the nitrile is required for conversion into the acid, it is not always necessary to isolate it, but the crudo product of the reaction can be at once saponified (Sandmeyer, B. 17, 2650; 18, 1492, 1496).

19. By the action of a warm alcoholic solution of K₂S the N₂ group is replaced by SH, thus: X.N₂.SH = X.Sli + N₂. The mercaptans so formed tion with cold KMnO4; so that by means of this reaction an NH, group can be replaced by SO.H (Klasen, B. 20, 349).

20. Hydric sulphide converts diazo-benzene into phenyl sulphide (CaH3)2S (Grache a Mann, B. 15, 1683).

21. Diazo- compounds combine with ethyl mercaptan to ferm unstable bedies X.N., SEt. which when boiled with alcohel yield sulphides;

X.N. SEt = X.SEt + N. (Stadier, B. 17, 2075). 22. By heating with acctic anhydride, acetylated phonols are fermed; X.N.,OH + Ac.O = X.OAc + N2 + AcOH (Wallach, A. 235, 234).

23. SO in presence of boiling alcohol cenverte some diazo compounds into the correeponding sulphonic acids: X.N.OII+SO₂=X.SO₂OH+N. (Hubner, B. 10, 1715).

24. Alkalis give insoluble pps. of complex constitution (Frankland, C. J. 37, 750).

25. Action of cyanogen compounds (v. Griess, **B**. 9, **13**2; **12**, **21**19; Gabriel, B. 12, 1637).

26. Cuprous nitrite replaces the diaze group by NO. The amine (1 mol.) is dissolved in exactly 2 mols. of dilute IINO, (or 1 mel. of dilute H.SO4) and the iced solution treated with one half of a solution of 2 mols, of NaNO,, the other half being added when the diazetisation is complete. The solution of the diazo-nitrite is added to a paste of 1 mol. of Cu.O (obtained by reducing CuSO, with glucose and NaOH) and the decomposition allowed to proceed in the celd. The yield of nitrobenzene from ani-line is 42 p.c. of the theoretical, but with other bases it is smaller (Sandmeyer, B. 20, 1494).

27. Primary and secondary amines, react at once with diazo-compounds forming diazo-amides $(q, v_*): X.N_*.O11 + 11.N.Y = X.N_*.N11Y + 11.O.$ In these bodies the diazo-radiole replaces H united to N, and on treatment with excess of acid they readily regenerate the diazo- compound and amine. When diazo- compounds act upon salls of aromatic amines, the diazo- residue may replace H in the carbon-nucleus, with formation of amido-azo- compounds: X.N. OH + HY".NH X.N., Y", NH., + H.O. In the case of tertiary aromatic amines the latter reaction is the only one possible, but with primary and secondary aromatic amines the replacement in the nucleus may be preceded by the formation of a diazoamide, when there is ne large excess of mineral acid present. The readiness with which amidoazo-compounds are formed varies greatly with the amine: in some cases (e.g. (a)-and (b)-naphthylamme, phenylene diamine, &c.) the replacement in the nucleus appears to take place almost instantly; in others (c.g. miline) the reaction, at the ordinary temperature, takes ceveral hours for its completion, allowing the intermediate formation of the diaze-amide (Friswell a. Green. C. J. 1885, 917; Proceedings 1887, 26). In many cases the formation of an amido-azocompound will take place in presence of a large excess of acid, under which conditions the formation of a diazo-amide is precluded. In the formation of amido-azo- con-pounds of the benzeno series the diazo-residuo enters in the para- position to the NII, group; but when this place is already occupied it takes the ortho-position. The ortho-amido-ano- compounds appear to be differently constituted to the para-amido-azecompounds (v. Azo- compounds). If the amine

does not contain any free para- or ortho- position the formation from it of an amido-azo- compound doee not appear to be possible. When a diamine contains displaceable H atome para to each NH. group, it is capable of reacting with 2 mols, of ϵ diazo- compound to form a disazo- compound: 2X.N.OH + H., Y' (NH.), =

 $(X.N_2)_2Y''(NIL_2)_2 + 2IL_1O_1$

28. Diazo - compounds readily react uper phenols and their sulphonic and carboxylic acids in alkaline selution, forming oxy-azo-cempounds X.N₂.OH + H.Y".OH = X.N., Y".OH + H₂O Compounds of the form X.N.OY analogous to the diaze-amides have never been obtained. The above remarks (reaction 24) with regard to the position taken by the diazo-group in the benzenenucleus apply equally to oxy- and to amido-azo compounds. Also, the di-exy- compounds, simi larly to diamines, can give rise to disazo-compounds (X.N.,), Yi (OH), when they contain displaceable II atoms para to each OH group.

29. Diaze- compounds readily react with pyrrol, with formation of azo- and disazo-bedies X.N., C, II, NII and (X.N.,), C, II., NII, but ne com pounds analogous te diazo-amides (O. Fischer a

Hepp, B. 19, 2251).

30. Piperidine, on the other hand, gives rise to piperidides X.N₂.NC₃H₁₀ (O. Wallach, A. 235

31. Amido-thiophene, unlike aniline, does not appear to form diazo-amides when treated with diazo- compounds, but gives at once amido-aze compounds X.N.,C.SH.,NII. (Stadler, B. 18 2318). This is no doubt due to the tendency to replacement of the hydrogen being greater in the thiophene ring than in the benzene ring.

32. Diazo-compounds react with those bodies of the falty series which contain H united to C, replaceable by sodinm, e.g. nitro-methane, nitroethane, malonic ether, accto-acctic ether, other ketonic ethers, &c. The products are mixed azo- compounds: those from nitro methane, for inslance, have the constitution X.N. CH₂(NO₂) (Meyer, B. 8, 751, 1073; 9, 381; Zublin, B. 11, 1417; Kuppeler, B. 12, 2286; Richter s. Münzer, B. 17, 1926; Griess, B. 18, 961; Bamberger, B. 17, 2415; 18, 2563).

Salts.—The diazo-salts have the general formula X.N.A. They are mostly white erretalline solids, very easily soluble in water, but sparingly in alcohol and other. They are usually very unstable, and in the dry state are often very explosive, ospecially the nitratee and picrates. By the action of K.SO, upon diazo-chlorides, sulphites are formed of the constitution X.V.SO.K. Stannic and eupreus salts give double compounds of the formula (X.N_x.Ol), SuCl., (X.N.,LC))Cu,Cl., (X.N.,LE)Cu,Br., &c. (Griess, B. 18, 965; Lclinann, B. 19, 610). The platine-chloridee (X.N.Cl) l'tCl, are sparingly seluble pps. An excess of bromine produces yellow or red ppe. of the per-bromides X.N.Br., The diazo-sulphonic and carboxylic acids form salts X."(SO,M.N..OH with bases, as well as X."(SO,M.N..OH with bases, as well as X"(SO₂H).N₂A with acids.

Amides. As already mentioned (reaction 27), the action of primary and secondary amines upon diazo-compenieds gives rise to diazoarrides of the general formula X.N., NHY, or X.N. NY,Y. Thus diazo-benzene and aniline

diazo-benzene-anilide(diazoamidobenzene): II.N.OH + C.H.NH. = C.H.N.N.NH.C.H. bodies are in general yellow orystalline etable below 100° when free from acid. eatment with an excess of mineral acid are resolved, even in the cold, into the salt and amine. When the amine is stic the resolution products again slowly bine, if the conditions are favourable, to ce an amido-azo- compound X.N...Y".N11, ic with the original diazo- amide. The tion and recombination take place conatly when the diazo-amide is treated with and, of cold dilute HCl, or with unstable such as ZnCl, CaCl, aniline chloride, &c., coholic or anilino solution (Friswell a. R. J. 1885, 917; Wallach, A. 235, 233). chem a diazo-amido is treated with a salt of a different from that of which it is a comsand, the diazo- salt generated will react upon base of the two whose nucleal hydrogen is con readily replaced. Thus diazo-benzene disthyl-amide C.H., N., NMe, treated with aniline decide gives amido-azo-benzeno and diwetbyl-ammo; similarly, diazebenzene-milide H. N. NHC II. treated with m-phenylenelamine hydrochloride yields di-amido-azoengine C₆H₅,N₂,C₆H₃(NH₂), whilst aniline is free; but diazo-benzene-anilido treated th p-toluidino hydrochlorido gives amido-azosazene and p-toluidine, because the para H of amilino is more readily replaced than the the H of the p-toluidine. Even very weak tresolving most diazo-amides. In the latter the diazo- compounds produced immediately ounds, for diazo- residuos replace the hydrogen the nucleus more readily in phenois than amines (Henmann a. Occonomides, B. 20,

In general it may be said that the reactions of the diazo-amides towards reagents in presence of acids, are simply the reactions of the free diazo-salts; thus SnCl, and HCl reduce them to hydrazines (cf. reaction 7); heating with strong halogen acids gives haloid derivatives of the hydrocarbons (cf. reactions 11, 14, 16, and

17); &c.

It appears to be proved (Griess, B. 7, 1619; Nölting u. Binder, Bl. 42, 336; Meldola a. Streatfeild, C. J. 1887, 102, 43 I) that the diazo-amides of the types X.N. NHY and Y.N. NHX are identical-that is, the same body is obtained whichever of the two amines is diazotised and combined with the other. The resolution of these unsymmetrical diazo-amides quite corresponds to their formation, for they yield a mixture of hoth diazo-compounds X.N.OH and Y.N.OH, and both amines X.NH2 and Y.NH2. For instance, the compound CaH, NaH.C.H, is obtained by combining either diazo-benzene with p-toluidinc, or p-diazo-toluene with aniline; and on treatment with HCl it splits up equally into diazo-benzene, p-diazo-toluene, aniline, and ptoluidine. When only 1 mol. of HCl is present these resolution products will recombine to form one or more amido-azo- compounds according to circumetanees.

Alkalie, even in boiling aqueous or alcoholio solution, usually have no action upon diazo-

amidee. The H of the NH group of the primary diazo-amides appears to have elightly acid properties, and by introduction of NO₂ groups into the nuclei the diazo-amide becomes sufficiently acid to dissolve in aqueous alkalis and form tolerably stable salts X"(NO₂).N₂.NM'.Y(NO₂) (Mrldola a. Streatfeild, C. J. 1886, 624; 1887, 102, 434).

By the action of alkyl haloids upon the primary diazo-amides dissolved in an alcoholic solution of (1 mol. of) sodium ethylate secondary diazo-amides are obtaine ! (M. a. S.; Friswell a. Green, B. 19, 2034; C. J. 1886, 746). When the two aromatic nuclei are the same the secondary diazo-amides obtained by alkylation are identical with those got by direct combination of tho diazo- compounds with secondary amines. But according to Meldola and Streatfeild (C. J. 1887, 134) the compound obtained by ethylating tho unsymmetrical m-nitro-diazo-benzene-p-nitranilido [3:1] $C_0H_1(NO_2)N_2H_1C_0H_1(NO_2)$ [1:4] (which is obtained either from m-nitro-diazobenzene and p-nitrauline, or from p-nitro-diazo-benzene and m-nitrauline) is different from either of the two ethyl-derivatives [3:1] $C_aH_a(NO_a).N_a.NBt.C_aH_a(NO_a)$ [1:4]

[4:1] C.H.(NO.).N...NELC₆H.(NO.)[1:3], obtained by combining m- and p-uttro-diazo benzeno with ethyl-p- and m- intramilines respectively. The two latter ethyl derivatives are split up by acide into m-uitro-diazo-benzene and ethyl-p-nitraniline and p-nitro-diazo-benzene and ethyl-m-intraniline respectively; but the former ethyl derivativo (obtained by ethylation) yields both diazo-compounds and both ethyl-nitranilines in about canal amounts,

Constitution of diazo-amides.—Although a considerable amount of work has been done on

this subject, the matter is still far from settled. The formula X.N-N.Y is disproved by the

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resolution of the alkyl derivatives into diazocompound and alkylated amine, since the latter could not be obtained fr. m a compound of that structure (Friswell a. Green, C. J. 1886, 746, and B. 19, 2034). The formula X.N.NIY does not N

appear to have much probability, since one would expect a body of the constitution C_1H_2NNIHC_1H_5 to give on reduction NII_3 and

hydraxobenzene; these bodies, however, are not formed from diazo-benzene-anilide, even by adding sodium to its boiling alcoholic solution (F. a. G.). Neither the latter formula nor that usually assumed X.N.N.NIY is capable of explaining the existence of more than two isomeric alkyl derivatives.

For further references concerning the diazoanides see Bayer a. Jäger, B. 8, 148, 893, Sarauw, B. 14, 2142; 15, 42; Kölling a. Baumann, B. 18, 1147; Staedol a. Bauer, B. 19, 1952; Wallach, A. 235, 233; Berntisen a. Goske, B. 20, 926.

Imides.—Obtained by the action of NH npon the per-bromides:

 $X.N_2Br_3 + NH_3 = X.N < \begin{cases} N \\ 1 \\ 1 \end{cases} + 3HBr$ (Griess, 4. 137, 65; B. 18, 963). They are also formed by

the decomposition of nitroso-hydrazines: $X.N < NH^2 = X.N < N + H_2O$ (Fischer, A. 190,

92, 96). They are usually yellowish oily liquids, insoluble in water, exploding when heated. When heated with cone. HCl they are converted into chloro-amido-compounds; 11X".N₃+11Cl - X"Cl(NII.)+N₂. Diazo-benzene-imide heated with H₂SO, diluted with an equal volume of water is converted into p-amido-phenol: C₆H₃,N₃+1l₂O - C₆H₄(OII)NII₂+N₂ (Griess, B. 19, 313; Fischer, A. 190, 67; 232, 236; Silberstein, J. pr. (2) 27, 116). Diazo-benzon acid reacts with phenyl-hydrazine, forming the imides of diazo-benzoic acid and of diazo-benzene together with amido-benzoic acid and ani-

Eine together with annoto-benizote acid and ani-
line, thua:
$$2C_0H_1 < \frac{CO_2}{N_1} + 2C_0H_2NH.NH_2 =$$

$$C_0H_1(CO_2H).N < \frac{N}{N} + C_0H_2.N < \frac{N}{N} +$$

$$C_0H_1(CO_2H).N < \frac{N}{N} + C_0H_2.N < \frac{N}{N} +$$

$$C_0H_1(CO_2H).N < C_0H_2.N < \frac{N}{N} +$$

$$C_0H_1(CO_2H).N < C_0H_2.N < \frac{N}{N} +$$

$$C_0H_1(N).N < C_0H_2.N < C_0$$

C₆H₁(NH₂)CO₂H + C₆H₂,NH₂. This equation is general (Griess, B. 20, 1528).

Diazo compounds derived from o-amido-azo-compounds.—Although the so-called o-amido-azo-compounds probably do not contain an NII₂ group, but are hydrazimido-

they are, nevertheless, slowly attacked by nitreus seid, with production of diazo-compounds. The only member of this class of diazo- compounds at present examined is that obtained by diazotising o-amido-azo-toluenc. In some of its reactions it behaves like an ordinary diazocompound, in others quite differently. On heating with water or alcohol it is decomposed with evolution of nitrogen. By zinc-dust and alcohol it is converted into m-p-azotoluene and N2. It is not reduced by SnCl2 or SO2 to a hydrazino, but gives a stable compound C. H. N. probably has which the constitution

C₆H₂(CH₃) N-NH N-N,C₆H₁(CH₃). This body is re-

converted by bremine into the per-bromide of the original diazo-compound. The diazo-imide loses nitrogen on heating, and yields tolyl-

identical with that obtained by oxidation of o-amido-aze-toluene (tolyl-hydrazimido-toluene). Hence the constitution of this peculiar diazocompound is probably

$$N-N.OH$$

 $C_{s}H_{s}(CH_{s})$ (Zincke a. Lawson, B. 19, 1452; 20, 1176).

already stated, the fatty diaze- compounds contain a molecule of water less than the aromatio diaze-hydratos. Their discovery is quite recent, and is due to Curtius. At present only a fow members of the group are known; these are obtained by the action of NaNO₂ upon the hydrochlorides of a mido-fatty-ethers X"H₂CO₂R,

and hence have the constitution X"(N₂)CO.R. They are unstable oily bodies, which exhibit the following reactions:—I. By boiling with water or ditute acids, they are usually converted into oxy-ethers, e.g. CH₂(OH)CO₂Et, with evolution of nitrogen; however, diazo-C(N₂).CO₂Et

succinic ether | by this treatment

CH_CO_Et, gives fumaric ether .- 2. With alcohols they yield alkyl-oxy-ethers, e.g. CH_(OEt).CO_Et.-3. With organic acids they yield alkoyl-oxyethers, e.g. CIL (OAc).CO.Et.-4. With aldehydes they yield alkoyl-ethers, e.g. CH Ac.CO Et. -5. With zinc-dust and acctic acid, they are reduced first to hydrazines, and then to the original amido-cthers .- 6. Iodine in cthereal solution converts them into di-iodo-ethers, c.g. CHI_CO_Et; bromine and chlorine act similarly .- 7. By long boiling with aromatic hydrocarbons, nitrogen is evolved, and condensation-products are formed .--8. By treatment with conc. halogen acids they yield haloid ethers, e.g. CH_Cl.CO.Et.- 9. By treatment of the diazo-others with NII, they are converted into the corresponding amides, e.g. CH(Na).CO.NHa, whose reactions are similar to those of the others. The free diazo acids, e.g. CH(N.).CO.H, and their salts, have not yet been isolated, on account of their instability (Curtius, B. 17, 953; B. 18, 1283; Buchner a. Curtius, B. 18, 2371; 19, 850; Curtius a. Koch, B. 18, 1293; 19, 2460). A. G. G.

DESCRIPTION OF DIAZO- COMPOUNDS.

A large number of diazo-compounds are mentioned under the amido-compounds from which they are formed.

Pscude-diazo-acctamide C, H, N, O, together with diazoacetamide, by prolonged action of strong aqueous NH, in the cold upon the methyl ether of diazoacetic acid CliN, CO, Me (Curtius, B. 18, 1287). Fine crystalline yellow powder, consisting of microscopic quadratic plates. Sl. sol. cold water, dilute 11Cl, and acetic acid. Insol. alcohol, ether, and benzene. Dibasic acid. On warming with water it evolves nitrogen. Aqueous alkalis evelve nitrogen and ammonia. It gives a green colouration with Fehling's solution; on boiling a black pp. is formed. Silvor and mcreury salts are reduced on boiling.

Salts.—A"(NH₄)₂: small yellow tables; sl. sol. cold water [c. 155°].—A"Ag₂ I¹₂aq: yellow microcrystalline pp., v. sl. sol. water. Explodes on heating.—A'Hg: yellow pp.—A"Cu*: sparingly soluble reddish-brown pp.—A"Pb: yellow pp.

Diazo-acetic acid CIIN., CO.H. The free acid and its salts have not been isolated on account of their instability. The alkaline salts are obtained in solution by treating the methyl ether with cold aqueens alkalis.

Methyl ether McA'. (129°) at 721 mm. S.G. 21 1·139.

Ethyl ether Eth'. [-24°]. (144°) at 721 mm. S.G. 22 1.073. Formed by the action of NaNO₂ on a concentrated solution of the hydrochloride of amido-acetic ether. Yellow oil. V. sl. sol. water; miscible with alcohol, ether, benzene, and light petroleum. Explodes when come

LSO, is poured upon it. Reactions.-1. Boilng water gives glycollio sther, nitrogen, and loohol.—2. Boiling alcohol gives CH, (OEt). CO, Et ind nitrogen. - 3. Benzoic acid gives riso to CH. (OBz).CO, Et. -4. Reduces AgNO, in cold squeous solution .- 5. Reduces hot Fehling's plution .- 6. Zine-dust and acctic acid form an instable hydrazo- derivative NII...NHCH2.CO.Et, which is further reduced to ammonia and NH...CH...CO..Et. The hydrazo- derivative reluces cold Fehling's solution. -7. Combines with aldehydes, thus: Ph.CHO + CHN. CO.Et -2 Ph.CO.CH...CO.Et + Na.—8. Changes on standinginto azin succinic other, C. II. N.O.Et. [245]. D. Boiling aniline produces NPhH.CH .. CO.Et .-10. Conc. HCl gives CH. Cl. CO. Et .- 11. Hot cone. KOH forms a crystalline pseudo-diazo-salt whence boiling diluto II, SO, liberates hydrazine, N.H. (Curtius, B. 20, 1632). 12. Boiling aromatic hydrocarbons produce condensation products; thus benzene gives C,H,CO,Et (Buchner

a. Curtius, B. 18, 2371).

Iso-amyl-ether C₃H₁₁A'. (160°) at 720 mm.

A mide CHN...CONH₂ [114°]; yellow tables or prisms; v. sol. cold water or alcohol. Formed by the action of strong aqueous NHa upon tho methyl-other. The aqueous solution decomposes on boiling with evolution of nitrogen and formation of glycollamide. By treatment in alcoholie solution with iodine it is converted into di-iodo-acetamide CHI2.CONII2 with evolution of nitrogen. It gives a blood-red colouration with AgNO₃, and in a few moments a reduction to metallic silver. It also reduces IIg(NO₃)₂ and Cu(OAc)₂. With Felling's solution it produces a red colouration, which becomes green on boiling (Cartius, B. 17, 953; 18, 1283).

Diazo-amido-benzoio acid

compounds.

Salts. - BaHCl; whits six-sided plates. -B.II.Cl.PtCl.: sparingly soluble small yellow trimetric plates. -B.AuCl., HCl: yellow insoluble ncodles (Griess, B. 17, 603).

bines with amines and phonols to form azo-

p-Diazo-aniline salts are formed by diazotising salts of p-phenylene-diamine (Griess, B. 17, 607).—C₆H₄(NH₂)N₂CliHClAu₂Cl₆ is an in-

m-Amido-diazo-benzene imide. Yellowish oil. Volatile with steam. Easily soluble in alcohol and ether.

Preparation .- m-Amido-phenyl-oxamio acid CaH4(NH2).NH.C2O2.OH is diazotised and converted into the tribrounide CaH, (N2Br2).NH.C2O2.OH. By treatment with NH₃ this yields the imids C_aH₄(N₃).NH.C_aO₂OH, which on boiling with aqueous KOH splits off the oxalyl group with

production of m-diazo-anilins imide. On diazofisation it gives a diazo- compound which combines with phenols and amines to form azodyestuffs. Decomposes explosively on heating.

Salts.-B'HCl: white soluble trimetrie plates.-B'.H.Cl.PtCl.; yellow needles (Griess, B. 18, 963).

m-Diazo-aniline piperidids.

Acetyl derivative C.II.(NIIAc).N., NC, H₁₀. [I01°]. From acetyl-m-tolylene-diamino hydrochloride by diazotisation and treatment with piperidino (Wallach, A. 235, 266).

(a)-Diazo-anthraquinone nitrate

C, 11, O, N.NO. Formed by passing nitrous acid gas into a solution of (a)-amido-anthraquinono in dry other (Böttger a. Petersen, A. 166, 150). Powdor, m. sol. water, v. sol. alcohol, insol. ether. When heated with water it gives N. and m-oxy-anthragminone.

Diazo-benzene. References : Griess, Tr. 1864,

iii. 667; A. 113, 201; 137, 39. Hydroxide Ph.N., OH(?). On adding acetic acid to an aqueous solution of Ph.N. OK a thick yellow oil is ppd.; this may be diazo-benzene hydroxide. It is very unstable.

Salts.-Ph.N.OK. A crystalline substanco obtained by adding excess of conc. aqueous KOH to a saturated solution of diazo-benzeno nitrate, and evaporating at 100°. Detonates feebly at 130°. V. sol. water and alcohol, insol. ether .-Ph.Ng.OAg: obtained as a greyish-white pp. on adding AgNO, to an aqueous solution of the preceding; explodes when heated.—(Ph.N_r.O)₂Hg: white pp. got by adding HgCl2 to the potassium salt (Griess, A. 137, 57).

Nitrate. - Ph.N. NO. S.G. 1·37. H.F. -47,400 (Berthelot a. Vieille, C. R. 92, 1074). Prepared by passing nitrous funies at 0° into an aqueous solution of aniline nitrate containing undissolved aniline nitrate in suspension; ppd. by adding alcohol and ether. Needlss; v. e. sol. water, m. sol. alcohol, insol. ether and benzene. Stabls in dry air in the dark, but decomposed in moist air. Explodes at 90° forming CO, CH, N, HCN, CH, and C. The decomposition may be roughly represented thus: C_6H_3 , N_2 , $NO_3 = 3CO + 3C + 5H + 3N$.

Reactions .- 1. Barium carbonate added to its aquoous solution produces Ph.N. C.H.OH and NaOHAq.-2. Aqueous NaOH, added to neutralisation, gives the 'benzene-di-azo-phenol' and a brown substance, C30H23N3O, insol. NaOHAq.-3. Aqueous ammonia produces diazo-benzons. anilids and two amorphous brown substances, $C_{1s}II_{21}N_2O$ and $C_{12}H_{13}N_3O$. The latter is very sxplosivs and is decomposed by boiling HCl into phenol, anilins, and N₂.—4. Boiling dilnte HNO. (1 mol.) forms o- and p- nitro-phenol (Nölting a. Wild, B. 18, 1338).—5. Aqusous K, FeCy, forms a compound C₁₈H₁N₂, [150°] (Griess, B. 9, 132), 6. Potassic ferricyanide gives (C₂H₁N₂)₃(H₂FeCy₃)₂. Sodium nitroprusside gives the compound 8. Nitro-benzyl cyanide and alcoholic KOH give a pp. of C₁₁H₁O₁O₂ [202°] (Perkin, C. J. 43, 111).

Chloride.—Ph.N_xCl. Formed in solution

by diszotising sniline hydrochloride. Combinations. - (C.H., N. Cl), SaCl, : white plates, sol, water, v. al. sol. alcohol and ether (Griess, B. 18, 965], -(PhN Cl) PtCl,: yellow prisms, v. sl. sol. water, insol. afcohol and other. PhN ClAuCl, : golden plates, in ol. water, m. sed. warm alcohed.

Property. Th.N.Br. Formed by adding bromine beaucethers disabilition of diago benzung and the or by washing the perbroadele for a long time with other. Pearly plates, v. e. sol. water,

insid, ether.

Perbrounde, -- Ph.N. Br. Formed by adding bromine discoved in HCIAq to an aqueous coliders of a diazole resear salt. harge yellow places, in ol. water and ether, in, sol, cold ntrohol. (over bronco benzene when distilled with Na CO, or when heated with alcohol.

Sulphote. PloNrSO, H. Ppd. by adding alcolod (3 volu) and other to a solution of diazobeagone nitrate muzed with an equivalent quantity of HSO, Picaus, v. e. pd. water, v. al. col. alcohol, in ol. other. Explodes at 100 '. Diryanole. C.H.S. or C.H.N.CN, HCN.

(69). Formed by the action of a diazo-benzene. ralt on a adution of KCN. Readily decomposes

(Galariel, B. 12, 1637).

Th.N.O.C.H (NO.). Parate. medles, obtained by mixing solutions of diazohenzene intrate and codium pierate. Very explosive (Bacyer a. Jager, B. 8, 981).

Sulphite. The potassium salt, Call, N. SO K, called also potassium diazobenzene sulphonate is ppt. by adding KOH to a mixture of diazolenzene nitrate and K.SO, Aq (E. Fischer, A. 190, 73). It forms explosive yellow crystals. Hromine in conc. HErAq pps. diasolenzeno perbromido. Zinc da 4 and acetis neid reduce it to the corresponding hydrazos compound.

Nitrite. Converted by Cu.O into mitro-benzene (Saichneyer, B. 20, 1407).

Benzeno's sulphinato Ph.N. SO .C.II [76". From sodium benzene sulphinate and dinzobenzene mitrate (Konigs, B. 40, 1532). Orange table (from alcohol); inside gold water, v. sol, alcidiol and ether.

m. Totr azo benzene C. H.(N. OH), [1:3]. Formed by the action of a large excess of nitrona sei l'apon er phenylene chamine în presence of a large excess of HCl. It combines

with 2 mode of an amine or phenol,

Saits. CHANACO-ROLL: small vellow plates; nearly in cluble in cold water and alcohol; heatel with dry Na,CO, it yields dichlora benzene. ... C.H. (N.Cl) Au Cl. : Pp. of sellow nucroscopic needles, explosive (Griess, B. 19, 317).

p Totr-azo-benzeno Call (NaOll), (1:1). Formed by the action of an excess of iditions and upon gephenylene-diamine in presence of * large execuse of neid. C.H. (N.Cl) PiCl; yellow crystandic explosive pp.; by heating with dry No.CO, it yields I di chloro benzene (Griess, $B, 19, 319_6$

Diazo henzone er amido benzeie acid

Call NoNH.C.R.CO R. Formed by mixing solutions of diago-bengene nitrate and in amidobenzoic acid (Griess, A. 137, 62). Small yellow plates (from ether). St. sol. alcohoi, v. e. sol. other, Collin's OH Puch.

Ethyl ciner. Eth's yellow crystals, v. c.

acl, alcohol and other, C. II, N.O.H. P.Cl.

The above diazo-benzene-amido-benzoic acid C.H., N., N.H.C., H., CO.H. is identical with diazo-benzoic-neid-anilide, C.H., N.H.N., C.H., CO.H. (Griess, B. 7, 1619; cf. Meldola, C. J. 51,

Dinzo-benzene-anilide Ph.N. NIIPh. Diazoamidobenzene. Mol. w. 197. [96°].

Formation, -By passing nitrous acid gas into an alcoholic solution of aniline (Griess, A. 121, 258).

Preparation.—A solution of 18 pts. of sodium nitrite is added to a solution 50 pts. aniline, 15 pts, conc. H.SO, in about 1,500 pts. of water, the temperature of the mixture is kept for 15 mins, between 25° and 30°, the pp. then filtered off, washed, and dried; the yield is 98 p.c. of the theoretical (Staedel a. Baner, B. 19, 1952).

Properties. Golden plates (from nlcohol), or har that prisms (from benzene). Explodes hetween 150 and 200°. Insol, water, and dilute solds, m. sol, cold alcohol, v. sol, hot nlcohol, v.

sol, other and beazene.

Exections, - 1. Hot conc. HClAq splits it up into phenol, nitrogen, and amiline; cold IIClAq gives audine and diazobenzene chloride, which, if an excess (more than I mel.) of IICl is not present, recombine forming benzene azo-aniline (q. v.). Unstable chlorides such as anilino hydrochloride or ZnCl2 also effect the conversion into benzene azo amiline.-2. Bromine in HBrAqgives diazolenzene bromide and tribromomiline.- 3. The hydrogen alom of the NII-group can be readily replaced by alkyl radicles by treatment with alcoholic haloids, and sodium ethylate. The alkylated diazobenzene-anilides thus obtained are split up by acids into diazobenzene and the corresponding mono-alkyl-nniline. A proof is thus afforded of the unsymmetrical structure of the amilide, and since the only other possible formula $C_b\Pi$ NNHE $C_b\Pi_b$ is

excluded by the fact that the body is not reduced by alkaline reducing agents to hydranobenzene and MIn the formula Ph.N. MIPh is probably correct (Friswell a. Green, C. J. 49, 716; B. 19, 2034). 4, Phonol at 100° gives benzene-pazo-plienol, Resorcin and the naphthols act similarly (Ifeumann a. Occonomides, B. 20.

Salts. Ph.N. NAgPh; orange needles .-(Ph.N. N11PhHCt), PtCl, : mistable crystals. Di-sulphonamide

 $\begin{array}{ll} C_{\rm e}H_{\rm d}(SO_{\rm e}NH_{\rm e}).N_{\rm e}NH_{\rm e}G_{\rm e}H_{\rm e}(SO_{\rm e}NH_{\rm e}) & \\ Yellow needles, & Fram (C_{\rm e}H_{\rm e}(SO_{\rm e}NH_{\rm e})NH_{\rm e})HNO_{\rm e} \end{array}$ and milrous acid gas (Hybbeneth, A. 221, 206). Cone. HCl converts it into C.H.CLSO,NH, C.II.(NII.)SO,NII, and N.,

Diazo-benzene-azo-benzene-p-sulphonic acid $\mathbf{C}_{\mathbf{a}}\mathbf{H}_{\mathbf{d}}(\mathbf{S}\mathbf{C}_{\mathbf{d}}) = \mathbf{N}_{\mathbf{d}} - \mathbf{C}_{\mathbf{b}}\mathbf{H}_{\mathbf{d}} + \mathbf{N}_{\mathbf{m}}$ Small yellow needles. Nearly insoluble in most solvents. Formed by iong action of nitrous acid on amido-benzeneazo-benzene sulphonic acid. Boiled with water it gives oxy-benzene azo benzene-sulphonic aoid: with alcohol it gives benzene-azo-benzene-sul-phonic acid (Griess, B. 15, 2186).

Diazo-benzene-benzyl-nnilide C₀H₂N₂N₄C₁H₃).C₁H₃ Benzyl - diazo - amidobeuzene. [81]. Yellow needles. V. sol. acetone. m. sol. alcohol, insol. water.

Preparation .- 30 g. of sodium are dissolved

in 300 c.c. of alcohol, a hot solution of 200 g. of hours' cohobation; half the alcohol is then diazo-benzene-anilide in 500 c.c. of absol. alcohol added, and the mixture heated with 140 g. of benzyl chloride for 1 or 2 hours; the product is precipitated by water and recrystallised from alcohol; yield, 200 g.

Reactions.—On heating it decomposes explosively. By excess of HCl it is resolved into diazobenzene and benzyl-aniline (Friswell a. Green, B. 19, 2036).

Diazo-henzeae-bromo-anilide r. Diazo-promo-BENZENE ANILIDE.

Diazo - benzene-p-chloro anilido. from p-chloro-diazo-benzene and auiline. By warming with phenol it gives oxy azo-benzene and p chloraniliue (Heumann a. Occonomides, B. 20, 908).

Diazo - benzene - ethylamide Ph.N., NHEt. From diazo-benzene nitrate and othylamine. Pierate C, Hank, C, H (NO,), OH.

Diazo-benzene ethyl-hydrazide I'h.N., N. 11 Et. From diazo benzene nitrate and ethyl hydrazine (E. Fischer, A. 190, 306). Very unstable oil. Reduced in alcoholic solution by zinc dust and acetic acid to ethyl-hydrazine and phenylhydrazme.

Diazo benzene imide C.H.N. Tri-azo-ben-

Formation. 1. Diazobenzene perbrumide is treated with aqueous NII, and the product distilled with steam, dried over CaCl., and rectified under diminished pressure (Grie.a, Tr. 1864, iii. 700). 2. By warming nitrosc-phenylhydrazine with didute KOH (Fischer, A. 190, 92).-3. By adding Na,CO, to a mixture of diazobenzene sulphate and hydroxylamine.

Properties, ... A heavy oil; insol. water, in. sol. alcohol and other. Detonates when distilled under atmospheric pressure. Not attacked by KOH.

Reactions, 1. Reduced in alcoholic solution by Zn and H_{SO}, to acciline and MH_s. 2. By heating with strong 11Cl it is converted into a nixture of a mol p chloro aniline:

C₄H₂N₃ + 2HCl = C₄H₂NH₂ + Cl₂ + N₂ = C₄H₄Cl.NH₄ + HCl + N₂.

8. By heating with H.SO, diluted with an equal

volume of water it is converted into pannidophonol: $C_0H_1N_1 + H_1O = C_0H_1NH_1 + O + N_2 = C_0H_1(OH)NH_1 + N_2$ (Griess, B. 19, 313). plienol:

Diazo-benzene dimethylamide Ph.N. NMe, From diazobenzene nitrate and aqueous dimethylamine (Baeyer a. Jäger, B. 8, 893). Yellowish oil; explodes when large quantities are heated; vol.tile with steam; insol. water and alkalis, v. e. sed, alcohol, other, and acids, Decomposed by acids into diazobenzene salts and dimethylamine. Aniline hydrochloride forms diazo-benzene-anilide and NMe, H leystrochloride. Picrato Ph.Nq. NMe_C.H. (NO.), (OII): acid gas (Hybbeneth, A. 221, 205). Minute yellow needles.

Diazo-benzene-methyl-anilide

C, H, N, NMeC, H, Methyl diazo amido benzene. Heavy deep yellow oil. Not volatile with steam. Miscible with alcohol, insol. water.

Preparation - 30 g. of sodium are dissolved in 300 c.c. of absolute alcohol and mixed with a hot solution of 200 g. diazobenzene anilide in 500 o.o. of absolute alcohol; when nearly cold 170 g. of methyl iodide are added; a vigorous distilled off and the residue precipitated by water, the oil separated and dried over CaCl,; the yield is 200 grms,

Reactions. - On heating it decomposes explosively. By excess of HCl it is resolved into diazal cazene chlorido and methylanilino (Friswell u. Green, C. J. 49, 718; B. 19, 2035).

Diazo-beazeue phosphonic acid nitrate (?). NO₃N₃C₆H₆PO(OH) 3aq. [188]. S. 58 at 18³; 59 at 80³. Fermed by passing nitrons acid gas into a boiling solution of anaido-benzene phosphonic acid in HNO, Aq (Miche elis a. Benzinger, A. 188, 288). Long white prisms (from HNO Aq). Explodes above 190°. V. sol. alcehol, sl. sol. ether. Not affected by boiling water, even in presence of 11.80,; slightly decomposed by boiling NaOllAq. Salts. Na A" 2aq. KA" aq. BaA" 3aq.—

 $A_{i}^{\alpha},A^{\prime\prime}:=1^{\alpha}bA^{\prime\prime},$

Diazo benzone-piperidide PhN,NC,H,. [43] (Baeyer a. Jager, B. 8, 893; Wallach, A. 235, 241).

Preparation - Aniline (100g.) is dissolved in aqueous 11Cl (210 e.c.) and the solution at 0° is diszotised with NaNO, (74 g.) and then poured into a dilute aqueous solution of piperidino (100 g.) containing KOH (60 g.) cooled with ice.

Properties. - Crystals (from ether or petroleum-ether). Reactions. -1. HCl passed into an ethercal solution gives diazobenzene chloride and piperidine hydrochloride. Aqueous HCI acts similarly. 2. Warm HCl form: N, chlorolengene , and piperidine hydrocldorde; phenol is a hyproduct. 11Br and 111 act similarly. 3. Hot dilute H SO, forms phenol. 4. An ethercal

solution of pieric neid gives dinzo benzeno pierate. o-Diazo benzene sulphonic acid

C.H. SO3 Yellowish tables, obtained by passing nitrous acid gas into water containing o-nmidobenzene sulphovie acid in suspension (Berudsen a. Limpricht, A. 177, 101).

m Diazo benzene sulphonie acid

C. H. SO₂. Prepared by passing nitrons acid gas into a concentrated solution of mounidobenzene sulphonic acid, contaming the free acid in suspension (Meyer a. Stober, A. 165, 165; Berndsen, A. 177, 88). Small columns (from water). Very explosive when dry. V. sol. water and decomposed by it at 60%. Boiling HBr gives m-bromo-benzene sulphonie acid. Beiling aleohol hus no action.

m. diazo-lenzene sulphomide nitrate $\mathrm{NO}_{\pi}\mathrm{N}_{\pi}\mathrm{C}_{\pi}\mathrm{H}_{\pi}\mathrm{SO}_{\pi}\mathrm{NH}_{\pi}, \ \ \mathbf{Fron.}\ \ \mathrm{C}_{\pi}\mathrm{H}_{\pi}(\mathrm{NH}_{\pi})\mathrm{SO}_{\pi}\mathrm{NH}_{\pi}$ by mixing with HNO, and possing in nitrous orange needles.

1. Diazo-benzeae sulphonic acid

C.11 SO₂. Formed by diazotising p-amidobenzene sulphonie acid (Schmitt, A. 120, 144; Facher, A. 190, 76). Small needles (from water). Insol, cold water, v. sol, water at 60°. Boiling water forms phenol p-sulphonic seid. An alkaline solution gives a red colour with aldehydes (Petri, H. 8, 291; Zahn, B. 17, Ref. 290), but reaction soon sets in and is completed by 1 or 2 this is not a characteristic test for aldehydes, as it is given also by many other hodies (E. Fischer, B. 16, 657; O. Loew, J. pr. (2: 31, 136).

Reactions. 1. Boiling alcohol forms benzene sulphonic acid. 2. PCl, at 100° has no action (facar, J. pr. 12, 20, 263).

Ethyl mercuptide G.H.(SO,H).N. SEt. Formed by combining policies between employing acide with antikalme colution of chyl mercaptan. The setoms raft (A.Na) forms yellow gli tening needle, w. ol. water., It is very unstable, readily decemps one with evolution of uitrogen. When builed with alcohol it yields ethyl-phenyl-sulphide p sulphonic moid C.H.(S.C.H.(SO,H) (Sudler, B. 17, 2075.)

Methyt an clide C II (80 (Ib.N. NMcC, II). Korned by combination of p diaze be exencisablyonic ucid with mono methyl andme in nearly neutral zolution. The sodium salt (NaA') forms large redouble (c) plate (c, v, sod. water, from which it is precipitated by white felted needles by alkalic nearly into alcohol. Not affected by boiling with dilute caustic sada. Acid resolve it into it is continued p diazo-benzenestiphonic need and no thyl andine, which when the neid is dilute recombine to form methyl andid benzene (azo-benzene sulphonic neid C(H₁80 (II), N₂ CH₂NIIMe (Beruthsen a, Goske, R, 20, 995).

Piper (dide C.H. (SO H) N. NC H₁₀ (Walbach, J. 235, 370). From sulplumilic acid by dinzofisation and treatment with piperidine (I and), and aqueous NaCH (I mol.). Salt.— NaA': satiny plates, "AgA', Stable in neutral or alkaline, olutions.

$$|Im(dv - C_i\Pi_i(SO_i\Pi), S \leq \frac{N}{N}|\{1;1\}|$$

Trio sheavene psutpienie acid. Formed by the action of phenyl hydrazine upon diazohenzene psudphonie acid suspended in obld water; diazohenzene imide, sulphanilie acid, and anilino, are formed simultaneously;

C.H.(N.)80/H C.H.N., C.H.(N.H.)80/H F.C.H.(N.H.). White deliquescent mediles, V. R. sol, alcohol and water, Sa.H.s. Ba.V.(2004); white aix sided plates, m. sol, list water, Phenyl hydra vine salt C.H.N.H.(Aua); long white glatening plates, m. sol, het water and alcohol, less in the cold, nearly insal, ether und chloroforny decret pseed by allahils, but not by HCI even when buttner (Gries 3, B, 20, 1528).

Diazo-benzene disulphonic acid

C.H.(80/H) SN/N The salts are formed by passing nitrons acid cas at 0 into a solution of the acid salts of C.H.(NH ((80/H).)[1334]). The free acid is unstable (Zander, 4, 198, 24).— KA, (BaA/2a).

Diazo-benzene disulphonic acid

C.H.(SO,H)(SO,N)". Formed by diazotising C.H.(MH,)(SO,H), (1.35c). Slender needles; v. sol. water nod alcohol. Decomposed by NaOH or BaCO. The radis are formed by diazotising salts of the numb henzene disulphonic acid (Heinzelmann, J. 188, 174; 190, 223).—KA'_+—BaA', Saq. 19A', Saq. 19

(Hemzenmann, 1, 1885, 1111, 1895, 2007).— Mag—Bak', Saq. PhA', Saq. Biaro-benzene disulphonic acid C.H. (SO, III) (SO, N.)". Formed by diazotising C.H. (NH,)(SO, III) (12; I).

Salts. — NH.A'. — KA'. — CaA', 2aq. — BaA', 2aq. —PbA', 3aq (Heinzelmann a. Zander. A. 198, 5).

Diazo-bsnzene p-toluide is identical with diazo-toluene anilide (q, v_i) .

o Diazo benzoie acid.

Nitrate NO, N., C., II, CO, H. By diazotisation of counids benzoic acid suspended in dilute INO₂ (Griess, B. 9, 1953). Colourless tables or prisms, v. e. sol. water, m. sol. alcohol. Explodes when heated. Boiling water converts it into safelylic acid. Repeated solution in water and ppn. with alcohol converts it into the so-called semi-initrate:

Perbromide C.H.N.O.Br.

Imide N. C. II. CO.II. Triazo-benzoic acid. [115]. From the perbromide and ammonia (Griess, Z. [2] 3, 165). Long needles. M. solbiling water.

m. Diazo-benzoic acid.

Sulphate SO_iRLN₂C₄R₁CO₂R. Formed by passing nitrons acid gas into a thin paste of the sulphate of m-amido-benzoic acid; ppd. by alcohol and ether. Long white lamina, v. e. sol. water; detonates when heated. Treatment with dilute alcohol gives rises to a 'basic sulphate' C₁I₁N₂O₂2I₂SO₄ (?), or more probably [N₄(C₄I₄CO₄II)], ILSO₄.

Nitrate NO₄N₂C₄II₄CO₄H (Griess, A.

Nitrate NO₂,N₂,C₄H₂,CO₂H (Griess, A. 120, 126). Its aqueous solution left in contact with BaCO₂ forms carboxy-benzene acid. With aqueous Na₂CO₂ it forms an acid C₁,H₁₁N₄O₂.

Hydroxide HO.N.C.H.CO.H. Unstable yellow oil.

Chloride Cl.N.,C.,H.,CO,H. Combinations.—(ClN.,C.,H.,CO,H),PHCI; yellow prisms.— (ClN.,C.,H.,CO,H),MCI;. This salt suspended in abcohol and treated with H.S gives benzoic acid, eldoro benzoin acid, and sulphydro-benzoic acid, HSC,H.,CO,H (Griess, J. pr., [2] 1, 102). Perbromide Hr,N.,C.,H.,CO,H. Colly pp.

Perbromide It_xN₂C₀II₄CO II. Oily pp. Converted by boiling alcohol into m-bromehenzoic acid (Griess, A. 135, 121; Cmze a. Häbner, A. G5, 106).

Ethylether; nitrate. NO, N., C, H, CO Et. Formed by diazotising mamido-benzoic ether dissolved in nitre acid (Griess, A. 120, 127).—Aurochloride (Cl.N., C, H, CO, Et) AuCl; golden prisms (from alcohol).

Amide; nitrate. NO₂N₂C₄H₄CONH₂. Fermed by the artion of nitrons acid gas on a solution of manido-benzamide in alcohol mixed with other (Griess, A. 120, 127). Needles. Platinochlarido (Cl.N.C.H., CONH.). PtCl., Imide N₂C₄H₄CO,H. Triazobenzoic acid.

Imide N. C.H. CO.H. Triazobenzoic acid, [160]. From the perbronide and NH, (Griess, Z. 1307, 161). Thin lamines V. sol, alcohol and ether, in. sol, boiling water. By heating with HCl it is converted into two isomeric chloro-amido-benzoio acids, (4, 3, 1) and (2, 3, 1):

N, C, H, CO, H + 2HCl = H, N, C, H, CO, H + Cl, + N, = U, N, C, H, Cl, CO, H + HCl + N, U, N, C, H, Cl, C, C, H, CO, Ag.

Anilide v. Diazo-Benzene-amido-Benzoic acid.

Bromo-anilide v. Diazo-bromebenzene-anidobenzoig acid.

Ntirite. Nitrate NO_xN_x,C_xH_y,CN. From mandio-benzonitrile (Griess, R. 2, 370). Explosive crystals; m. sol. cold water.—Perbromido Br₁N_x,C,H_y,CN. Crystals.—Imide N_xC_xH_y,CN. [57]. Needles, v. sl. sol. water.

p Diazo-benzoic acid.

Nitrate NO, N. C. H., CO. H. Explosive white prisms (Grices, J. 1861, 453).

Amide. Nitrate. NO N. C. H. CONH. (Griess, Z. 1866, I).

Imide N_pC₆H_pCO H. Triazebenzone acid. [185]. Thin laminas (Griess, Z. 1867, 164). m-Diazo-benzoic-m-amido benzoic acid

[331] COJII, CHAN, NHLCHACO II (153). Formed by passing nitrous acid gas into an alcoholic solution of manido benzoù acid, or by mixing aqueous solutions of manido benzoù acid and the nitrate of maliaco-benzoù acid and the nitrate of maliaco-benzoù acid (Griess, A. 117, 2; Z. 1861, 353). Orange grains, Explodes at 180. V. sl. sol, water, alcohol, and ather. Sol, alkalis and reppd, by acids. Boiling HCl forms manido-benzoù acid and machloro-benzoù acid. Bromine water gives bromo, and tri-bromo, benzoù acid Boiling water and iodine form ioda-oxy-benzoù acid. Nitrous acid passed into a boiling aqueous solution forms nitro-oxy-benzoù acid; untrou acid passed into a boiling alcoholic adution forms benzoù acid. Fuming HNO₂ gives tri-

nitro ovy-henzeie neid.

Saltz. (KH) A''. KA''. – Ag A''.

Mathyl ether Mc A''. 100 / Yellow needles.

Ethyl ether EtA''. [144]. Golden needles.

p. Diazo-benzoic-p-amido-benzoic acid [4:1] CO [11,C] H₁N_sNHACH₄CO [11] [1:1]. Orange powder, v. sl. sal, boiling abendal. Formed by passing nitrons acid gas into an alcoholic solution of p-amido-benzoic acid (Belistein a. Wilbrand, J. 128, 269).

m-Diazo-benzoic-p amido-benzoic acid

[3:1, CO] H.C.H., N., N.H.C.H., CO] H (1:1). From the intrate of m-diago beneson acid and p unidonely disclosed (Giress, J. 1864, 359). An isomeone (2) acid is jet from the intrate of p-diagobenzoic acid, and m-acide benzoic acid.

m.Diazo-bromo-benzene.

Perbroseide C.H.Br.N.Br. (Wurster, A. 176, 173).

p-Diazo-bromo-benzene.

Nitrate C.H.Br.N.NO. Formed by passing nitrons acid gas into an aqueous solution of p-brome anilune mirate (Grees, Tr. 1864, iii. 695). Ppd. by alcohor and ether.

Hydroxide C.H.Br.N. OH: bright yellow needles. Veryexplosive, -C.H.Br.N. OK: From the nitrate and strong KOH; gr. e. the proceeding body when treated with acetic acid. -C.H.Br.N. OAg.

when treated with acctic acid. C.H.Br.N.OAg. Bramide C.H.Br.N.Br: scales, v. sol. water, m. sol. alcohol, insol. ether. (C.H.Br.N. Br), PtBr.

I'erbromide C.H.Br.N.Br.: monoclinic prisms (from alcohol), v. sl. sol. ether, insol. water.

Chloride C.H.Br.N.Cl: from the bromide and moist silver chloride.—(C.H.Br.N.Cl)AuCl.—(C.H.Br.N.Cl).PtCl.

Sulphate C.H.Br.N. SO.H: slender prisms.

Imids C₄H₄Br.N₂. Triazo-bromo-benzone, [20°]. Insol. water, m. sol. alcohol, v. sol. other above benzene. Reduced by Zn and H₂SO₄ to bromo-aniline and NII₃.

Cyanide C₄H₄Br.N₂CNHCN, [128]. From

p-diazo-bromo-benzene nitrate and aqueous KCN (Cabriel, 18, 12, 1638).

Autilide C.H.Br.N.,NH.C.H., or, alternatively C.H.N.,NH.C.H.Br. Formed either from diano beaucine nitrate and p-bromo-milline of from poliniza-bromo-beaucine, nitrate and aniline (Griess, B. 7, 1618). Vellow plates; v. c. sol, ether, m. sol, alcohol. (C.H.,Br.N.,11C.).P.IC., p. Brompo-avillide C.H.Br.N.,NH.C.,II.Br.

[115]. Obtained from p-brome-aniline, --

(C_{1.}H.Br.N₂).H.PtCl₆. Diazo di-brome-benzeue.

Nitrate C.H.Br., N., NO. (2:4:1). Obtained by passing nitrons neid into an aqueous solution of the attrate of distreme aniline (Griess, Tr. 1864, iii, 704). Needles or plates.

Flatinachlaride (C.H.Br., N.Cl), PtCl.: orange plates.

Pertamide C H₃Br₈, N Br₈) slenderneedles, Imide C₄H₃Br₈N_c [62]; needles, Distronacionilide

C.H.Br₉, N.HC₈H₂Br₁, [1687]. From (2,4,1)-di bronn amiline. Golden needler, v. sl. sol. alcohol and other,

Dinzo-tri bromo benzene.

Nitrate C_eH₂Br_xN₀,NO_x [2:4:6:1]. Formed when a rapid current of nitrons acid pastis passed when a rapid current of nitrons acid pastis passed pen ion together with excess of HNO_x. As soon as excepthing is dissolved, ether is added and a height yellow crystalline [1], of the diagoniticate is found (H. Silberstein, J. pr. [2] 27, 1024.

Properfies, - Yellow trimetric plates, Explotes at 85%. Sol. water and HCl. V. sl. sol. alcohol, ether, and benzene.

Reactions, 1. Builed with alcohol it gives tri bromo henzene, N₂, HNO, and uldehyde, 2. Boiled with water, gives off no nitrogen but forms undetermined compounds. 3. Heated with glacial acetic acid gives off Ng and nitrous funces and leaves tri-bromo benzene. 4. Heated with ben sene (1 pts.) it decompoles at 45° forming a diazo di Iraggo idienal (g. r.), tetra bromohousene '98 Land pitrohenzene, . 5. Heated with CHCl, it gives diazo di bromo phenol and tetrabrome-benzene. 6. Hented with concentrated HCl it forms cryctals of the perbrounds of diago. tri broma-benzene chloride, C.H.Br.N.CIBr., Probably chlorice, liberated in this reaction, turies and bromme from some of the tri-bromocompound, which broming then unites with the diszo-chloride. The perbromide explodes at 100% farming chicao tri-bronco benzeno. -- 7. With HBr it gives the brounds (9. v.). --8. With III it gives tri-aromo-iodo brazene. Sulphate G.H.Br.N.SO.H.

Properties: -Colourless prisms. Sol. water, sl. sol, alcohol, insid, ether and benzene.

Reactions.—1. Decomposed by alcohol into Iribromo-benzene, II, SO, and N₂, 2. Boiled with acidebled vater it forms no tri-bromo-phenol.—3. Reated with glacial acclic acid it forms tribromo-benzene.—4. Not affected by boiling ben-

Bromide C.H.Br.N.Br. Small golden tri-

metric tablets, got by adding dilute HBr to a solution of the nitrate. Decomposed by sunlight into N, and C.H.Br., Sl. sol. water, insol. alcohol and ether. Heated with glacial acetic alcohol and etner. Helical C.H.Br., acid it gives unsymmetrical C.H.Br., Formed by

Perbromide C.H.Br.N.Br., Formed by adding cone. fills to a solution of the nitrate, CH.Br.N.NO4; the liquid is filtered from CH.Br.N.Br and evaporated to crystallisation. It forms orange prisms and boliaves very much

like C.H.Br.N.ClBr. (q. v.).

Cll Br, N, Cl. Br2 Chloro-perbromide From tri-bromo diazo-benzene nitrate (q. v.) and HCl. It explodes at 100° forming obloro-tribromo-benzeno. Reactions, -- 1. With NH, gives tri-bromo diazo beazene imido (q. v.) .- 2. With alcoholio dimethylaniline forms tri-bromo-benzene azo dimethylaniline Call Br. Nr. CaH, NMo. 3. With alcoholio methyl-di-phenylamine it forms tri-bromo-penzene azo-methyl-di-phenylaminoCallaBr, N., CallaNMel'h. -4. With mercuric dinhenul it forms chloro tri-bromo-benzono and

India C.H.Br., N. (55°). Formed by adding dilute ammoniu to C.H.Br., N. (50°). Formed by adding dilute ammoniu to C.H.Br., N. (50°). Colour-less needles. May be distilled with steam. Insol. water, sol. warra alcohol, ether, and CHCl, Unlike diazo beuzene-imide, it is not reduced by

Zn and HSO, to NH, and tri-bromo-anilino Zii Sha 11,307, 187 A. [2] 27, 116).

An ilida C.H.Br., NHPh. [104°].

alcoholic anilino (2 mols.) and Call.Br.N.NO. (1 mol.). The pp. is crystallised from alcohol (Silberstein, J. pr. [2] 27, 121). Yellow, glittoring, triclinic prisms. Insol. water, sol. hot alcohol, other, and benzene. Boiled with glacial acetic aoid it gives nitrogen and tri-bromo-aniline.

Tri-bromo anilide

C4H.Br4.N., NH.C.11.Br, Obtained by passing N10, slowly into a cold alcoholic solution of tribromaniline. Some Call Br. N. NO. is formed at the sume time. The product is washed with water and hot nleohol and crystallised from benzenc. The yield is bad (Silberstein, J. pr [2] 27, 120). Insol. water and ulcohol, v. sl. sol. ether. V. sol. CHCl, and benzene. Not attacked by cold acids, but decomposed by boiling with acids with evolution of N.

Dlazo p brome-penzene-amide-benzoic acid C,H,Br.N, NH.C,H,CO,H. Diago benzoic acid p.bromo-anilide. From p-dinzo-bromo-benzene nitrate and mamido-lenzoic acid (Griess, J. 1866, 453). Clusters of needles.

Diazo-bromo-benzene sniphonic aold

 $\mathbb{C}_{\mathbf{t}^{\mathbf{H}_{\mathbf{s}}}}\operatorname{Br}<\frac{\mathrm{SO}_{\mathbf{s}}}{\mathrm{N}_{\mathbf{s}}}>\left[\begin{smallmatrix}\mathbf{s}&\mathbf{i}\\\mathbf{i}\end{smallmatrix}\right]$. From the corresponding bromo amido sulphonic acid (Borns, A. 187, 371). Small yellow needles, v. sol. water, in. sol slochol; explodes when struck or when heated. Heated with alcohol, it gives m-bromobenaene sulphonic acid. Conc. IIBr gives C.H.Br. (SO.11) [4:1:2].

Diazo-bromo benzene disnipbonio acid

From C₄H₄(NH₄)(SO₂H)₂Br [1:4:6:2]. Minute states; v. sol. water and alcohol. Does not explode when struck (Zander, A. 198, 15),—
KA*Saq.

O.H.Br. SO. From O.H.(NH.)Br.(SO.H.)
[1:2:6:4]. Yellowish scales; explodes above [1;2:6:4]. Yellowish scales; explodes above 100°. V. sl. sol. cold water, v. sol. hot water

Boiling water gives di-bromo-phenol sulphonic acid. Heated with alcohol it gives di-bromobenzene sulphonic acid.

Diazo - di - bromo-benzene-sulphonic-acid- dibromo-snlpbanilide

C₄.B₁.(SO₄H).N_y.NH.C₄H.Br.(SO₄H). [70°-80°]. From C₄H₂Br₂(NH₂)SO₄H [2:4:1:5] by warming with alcohol and KNO. Needles (from water). Insol. aloohol (Baessmann, A 191, 229).

Diazo-di-bromo-benzene-disulphonio acid

 $C_sHBr_s(SO_sH) < SO_s > From the nitro-ben$ zeae disulphonic acid [1:3:5]? whose chloride melts at 96°, by reduction, bromination and diazotisation (Heinzelmann, A. 188, 183).

Diaxo-tri-bromo-benzene sulphonic acid

CallBrs SOs From tri - bromo - aniline, C.H.(NH.)Br. [1:3:4:5] by anlphonation and diazotisation (Spiegelberg, A. 197, 291). Minute needles.

Diazo-tetra bromo-benzene p-snlphonic acid C₄Br₄<SO₃>. Crystalline powder not decomposed by boiling alcohol (Beckurts, A. 131, 225).

Diazo-bromo nitro toluene snlphonie acid C.HMeBr(NO₂)< $\stackrel{N_2}{SO_3}>$. Formed by projecting m-bronno p-antido-toluenc o-sulphonic acid into

fuming HNO, (Weekwarth, A. 172, 203). Diazo-di bromo-nitro-tolnene anlphonio acid C.Br.Me(NO2) < N2 >. Formed by projecting

di-bromo-o-amido-tolucne p-sulphonio acid into funning IINO, (Hayduck, A. 174, 355).

Diazo di bromo phenol

 $C_{e}H_{2}Dr_{2} < \stackrel{O}{\underset{N_{2}}{\langle}} [O;N_{2}=1;2].$

Preparation .- Bromine water is added to an aqueous solution of o-diazo-phenol chloride and the pp. is dissolved in fuming HCl, filtered through ashestos and ppd. by water.

Properties. -- Orange crystalline powder, much less stable than the p-compound. When heated it explodes at 128°. It is more soluble in cold water than the p-compound; on warming the solution a resin is formed. The solution gives an amorphous grey pp. with silver nitrate. It is almost insoluble in alcohol, ether and CS, but readily dissolves in CliCl, hot benzone, and benzolino. It does not reduce Fehling's solution, nor form a orystalline body with NaHSO, C.H.Br. (OII) N.Br 1 aq: decomposed by water.

Ethyl, ether, nitrate of. C.H.lir., (OEt). N.NO. Got by passing N.O. into alcohol containing HNO, and di-bromo-o-amidophenetol in suspension, and pouring it into dry ether at 0°. Properties. -Prisms. Explodes at 102°. If its alcoholic solution be diluted with water (10 vols.) and boiled as long as Nz escapes, it is converted into di bromo-phonetol. bromino being replaced by hydrogen, not by hydroxyl (Möhlan a. Ochmichen, J. pr. [2] 24,

U.H.Br. (N. [0:N. = 1:4].

Formation.—1. Obtained by adding bromine water to an aqueous colution of any ealt of p - diazo-phenol: C,H,(OH)N,Cl+2Br,= $C_4H_2Br_2 < \frac{Q}{N_2} + HCl + 2HBr. -2$. From HBr and

diazophenol nitrate (q. v.).

Properties. — A flocculent yellow pp. resombling sulphide of arsenio. Dissolves in boiling water, without decomposition, and orystallises in yellow prisms as the colution ocolo. Almost insoluble in cold water, ether, and CS2, somewhat more soluble in CHCl2. Soluble in alcohol and in nmyl alcohol. May be kept for months in closed bottles in the dark, but, when exposed to air and light, it even turns brown. Heated to 137° it explodes.

Salte. — Unstable, docomposed by water and by alcohol. C.H.Br.₂(OH)N.Br.aq. — {C.H.Br.₂(OH)N.Br.}.Pr!.PtCl₁: triolinic plates, decomposed by water. — C.₂H.Br.₂(OH)N.₂SO₄H (Böhmer, J. pr. 132, 458).

Reactions .- 1. Boiled with water of which the boiling-point, by addition of calcic chloride, has been raised to 120°, it gives off nitrogen and forms di-bromo-hydroquinone:

C_sH₂Br₂ $\stackrel{O}{\underset{N_2}{\bigcirc}}$ + H₂O = C_sH₂Br₂ $\stackrel{O}{\underset{N_2}{\bigcirc}}$ (B5hmer, J. pr. 132, 464).—2. Dissolves in a hot solution of NaIISO₃; and as the solution cools, yellow needles of C_sH₂Br₂(OII)N₂SO₃Na 2aq are formed. These erystals dissolve in ether, benzene and CS₂; do not explode when heated; show Liebermann's reaction; do not reduce Fehling's solution; and give with BaCl₂ golden soales of {C.H.Br. (OH) N.SO₃ | 2Ba5aq. - 3. Reduced by Sn and HCl to di-bromo p amidophenol.

Dlazo . di . bromo . phenol. (?) Identical with the preceding diazo-di-bromo-phenol just desoribed may be converted into this isomeride by first reducing it to dibromo-p-amido-phenol

hydrochloride and again diazotising.

Properties. - Explodes at 145°, has a greyishyellow colour, is insoluble in water. Crystallises from alcohol in much thinner needles than the preceding. With NnHSO, it forms small plates, whereas the sulphonate of ite parent-isomeride crystnllises in needles (Böhmer, J. pr. 132, 471).

Diago di bromo phenol

 $C_4H_2Br_2 < N_2 > .$ [Br:Br:N:O = 1:5:6:3]. Formed by heating tri-bromo-diazo-benzene nitrate with

benzene at 45° (Silberetein, J. pr. [2] 27, 107).

Properties.—Oblique prisms, from water.

Crystallises from alcohol. Explodes at 142°.

Soluble in hot alcohol. Nearly insoluble, in thloroform and ether.

Snlts .- These are very unstable; they are formed by warming with rather strong colutions

of acide, but are saponified by water.

Reactions.—1. Not attacked by boiling water. 1. Hested with otrong HBr forms tri-bromophenol and Nr. -3. Reduced by Sn and HCl to li-bromo-amido-phenol and NH.

Constitution. - The hydrochloride of the di romo-amido-phenol obtained by reduction, gives shmitt's reaction upon the gradual addition of linte bleaching powder, viz.: a violet coleur

followed by a white pp. As this reaction is sharecteristic of p-amido-phenols, this body must be di-bromo-p-amido-phenol, and since it is formed from C.H.(NH.)Br. [1:2:4:8] its con-stitution is as given above. It appears to be different from Böhmer'e compound, exploding at 137°.

Diaso-trl-bromo-phonol.

Ethyl other, nitrate of. C.HBr. (OEt) N.NO. (OEt: N. = 1:2). Triolinic plates, prepared by brommation of o-diazophenetol. Does not explode when etruck. In melting-point tubes it oxplodes at 93°. It is decomposed by boiling water into tri-bromophenetol, the N₂NO, being displaced by H, not by OH (Möhlau a. Ochmiehen, J. pr. 132, 484):

2C₄HBr₄(OEt)N.NO₃ + 2H O =

2C₄HBr₄(OEt)H + 2N₃ + O₂ + 2HNO₄.

Diazo-bromo-tolnene sulphonio acida

CH₂.C₂H₂Br<SO₃>. Five are known:

p-diazo-m bromo toluene o-snlphonlo acid (Weokwarth, A. 172, 196). Red orystals. Hoated with alcohol under pressure it gives bromotoluene sulphonio ncid.

p-diaso brome telene m-sulphonic soid (Pechmann, A. 173, 211). Heated with alcohol under pressure it gives bromo-toluone m-sulphonie acid.

diaso-o-bromo-tolnene m-snlpboale

(Schäfer, A. 174, 360).

disse-p-bromo tolnene m-sulphonio soid (S.). diase-p-breme-tolucue o-sulphonio soid (8.) o-Diazo-di-biemo-teluone p-anlphonio

layduck, A. 174, 352). Diazo-campher v. Camphor.

Diszo-p-chloro-benseno [1:4] C.H.Cl.N. OH. Yellow explosive powder, ppd. from its sats by HOAe. Salts.—C.H.cl.N.NO; white platee.—C.H.cl.N.Br.; yellow prisms.—C.H.cl.N.D.,—C.H.cl.N.Br.; yellow prisms.—C.H.cl.N.D.,—C.H

Anilide v. Diazo benzene p-chloro-anilida. p Chloro-anilida C_eH₄Cl.N₃.NH.O_eH₄Ol [125°].

Diaso-di-ohloro-bonsono.

Salte.— C.H.,CI, N.,NO₂.— C.H.,CI₂.N.₂Br₂.— (C.H.,CI, N.,CI), PtCl., (G.). Dischloro-anitide C.H.,CI, N.,NH. C.H.,CI,

[127']. Needles, v. el. sol. alcohol and ethor.

Diszo-ohlero-nitro-phenol

C.H.Cl(NO2) < N2 >. From ohloro-nitro-amidophenol (Griess, A. 113, 215). Brownish-red eolumns (from alcohol).

Diane-di-ohloro-phenel C.H.Cl. N. From C.H. (OH)Cl. (NH.) [1:3:5:2] (Sehmitt a. Glutz, B. 2, 52). Brown flocculent powder.

Diazo-tri-chloro-phenol $C_sliCl_s < N_2$. From tri-chloro-p-amido-phenol (Lampert, J. pr. 33, 375). Golden needles, explodes at 187°, sol. hot alcohol or benzene, insel. ether.

Reactions .- 1. Boiling alcohol gives ohloro-phenol [54] (253').—2. Como. Nn, BOAq forms C.HCl, (ONa). N, SO, Na, of which the acid C.HCl, (Oli). N, SO, H, explodes at 200°, and forme an orange crystalline barinm salt .- 8. HI

forms C.HCl, I.OH. Diago-chloro-thymol obloride O.HCIMePr(OH).N.CL By the action of N.O. ac a cold alcoholic solution of hydrochloride of | chloro-amido-thymol (Andresso, J. pr. 131, 180). Precipitated by ether. Colourless needles, often

grouped in fans.

o Diazo-cinnamle scid. I'rom amido-cinnamio acid (10 pts.), HCl (9 pts. of S.G. 1-19), water (70 pts.), and NaNO_p. The chloride separates as a yellow powder (Fischer a. Kuzel, B. 14, 478; A. 221, 272). The nitrate C.H. (N.NO.).CH.CH.CO.H. forms clear prisms. Both salts may be boiled with potash without giving off nitrogen, but they are decomposed by boiling water, forming o-coumaric acid. With Na₂SO₃ they form C₂H₄(N₂SO₂Na).CH;CH,CO₂H₄ whence, by reducing with zino and HCl, C, H, (N11.N11.SO, Na). C11:CH.CO, H. The latter forms slender needles. It reduces HgO in the cold, and Fehling's solution. IlCl in the sold converts it into hydrazido-cinnamic acid (q. v.).

p-Dlaze-cinasmic acid. The chloride Call (N.N.Cl).Cll:Cll.CO₂ll is prepared by the action of NaNO, on p-annido-cinnamic acid suspended in 11Cl (Gabriel, B. 15, 2300). Long noodles (containing aq). M. sol. water. Can be dried at a gentlo heat w thout decomposition.

p-Dlase-cresol Me.C., II. diazotising C_all₄Mc(Nll₂)(Oll) [1:3:4] (Wagner, B. 7, 1270) (MeC_all₄(Oll)N₄Cl)₂PtCl₄: powder, m. sol. water.

Disse-y-cumeno-sulphito.

Salt. C. 11, (CH2), N. SO, Na [1:3:4:6]. Transparent prisms (containing 2 aq). Not explosivo

(Haller, B. 18, 90).

Diazo-y-cumeno-camids. Diazoamidocumene. [1:3;4:6] C₆11,Mo₂,N:N.NH.C₆11,Me₃ [6:1:3:1]. [131°]. Formed by the combination of diazocumeno with cumidine (Nölting a. Baumanu, $B_{m{\epsilon}}$ 18, 1117). Yellow tables (from ether). V. sol. benzone, other, and acctone, m. sol. alcohol.

Dlase-cuminic-amido-cumiaic acid $Pr.C_aH_a(CO_2H).N_{j,i}NH_iC_aH_aPr.CO_2H_i$ by passing uitrous acid into an alcoholic solution of amido-cuminic acid at 0 . Minute prisms or leaflets (Griess, A. 117, 62).

Disze ethaue sulphito C. H., N. SO, H. Diazo-

ethane sulphonic sold.

Salt. - KA'. Formed by the action of HgO on the corresponding hydrazo derivative C₄H₂NH.NH.NH.SO₄K (Fischer, J. 199, 302). V. sol. water; p.d. by alcohol. Explodes when heated. Decomposed by boiling neids, N2 and SO, coming off. Reduced by zinc-dust and feetic acid to C.H., NH. NH. SO, K.

Diaso-ethoxane C.H.O.N.,—O.C.H. (?) V.D. 402 (calc. 408). Prepared by the action of silver hyponitrite, AgNO, on Et1 (Zorn, B. 11, 1630). Neutral liquid. Exceedingly ex-II, 1630). Neutral liquid. Exceedingly ex-plosive. Is decomposed by water with production of aldehyde and alcohol: (C.H.), O.N. + H.O. = N₃ + CH₂, COH + C.H., OH + H.O. By tin and acetic seid it is reduced to nitrogen and alcohol; $(O_sH_s)_sO_sN_s+H_s=2C_sH_sOH+N_s$ o-Disso hemipio sold

C.H(OMe),(CO.H) < N. O. Diaro-di-methoxyphthalic acid. Formed by the action of nitrons acid upon o amido-hamipie acid. Yellow mieroerystalline powder. Sl. sol, ordinary solvents. 140°-150°. Converted into hemipic soid by boiling with alcohol. Hydrochloride C.H(OMe),(CO.H), N.Clag: long colourless nesdies.

The sulphate forms small prizms (Grüne, B. 19, 2302).

Diazo-hippurio acid.

The nitrate CO.H.CH, NH.CO.C.H, Na.NO. is formed by diazotising the nitrate of m-amidohippuric acid (Griess, Z. 1867, 165).—Per. bromide C,H,NO,N,Br,: yellow prisms. Imide C,H,NO,N,: tables or needles.

Diazo-leucaniline v. Hexa-azo-tri-phenyl-

niethane (infra).

p-Diaze-iodo-benzeno C,H,I.N,OH: yellow pp. Salts.—(C,H,I.N,Cl),PtCl,-C,H,I.N,NO,-C. II.I.N. SO, II: small plates, v. sol. water, sl. sol. alcohol. - C.H.I.N.Br. - Imide C.H.I.N. (Griess, Tr. 1864, iii. 706).

(a) Diazo-naphthalane (Grices, J. 1866 453). Nitrate C, H. N. NO : formed by diazotising a-naphthylamine nitrate. - Perbromide C₁₀H.,N.Br, : orange erystals. - Platino-chloride (C₁₀H.,N.Cl),PtCl, - Imide C₁₀H.,N₃: yellowish oil (cf. Fischer, A. 232, 242).

A solution of the chloride neutralised by Na₂CO₃ gives a brown pp. Part of this dissolves in alkalis and nppears to be C10 H, NO, the rest is ppd. as minute crystals by adding alcohol to its bouzene solution. Analysis indicates C₅₀H₅₃N₅O₂. Both form crimson solutions in nicoliol, ether, benzene, and glacial acetic acid (P. F. Frankland, C. J. 37, 750).

(β)-Diazo-naphthalene. Obtained by diazo-tising (β)-naphthylamino (Liebermann a. Palm, A. 183, 267). The sulphate forms palo yellow needles, and the perbromide orange needles.

(C₁₀H₁,N₂,Cl)Cu₂Cl₂*: very unstable yellow pp. which is formed on adding Cu₂Cl₂ to a cold solution of B-diazo - naphthalene - chloride. (C₁, H₁, N₂, Br)Cu₂Br₂: red pp.; on boiling with water it ovolves nitrogen yielding (β)-bromonaphthalene (Lellmann, B, 19, 810).

(a) - Diaze - naphthalone - (a) - naphthylamide. [100]. Formed by action of nitrous acid on (a)-naphthylamine, or by ppg. a solution of (a)-diazo-naphthindene chloride with (a)-naph-Brown lamine (from alcohol). Acids resolve it into naphthylamine and diazonaphthalene (Martius, Z. [2] 2, 137).

(a) Diaze-naphthalene sulphonio acld

 $^{*}C_{*0}H_{*} < \stackrel{N:N}{SO_{*}} >$. [1:4]. Got by passing nitrous acid gas into (a)-naphthylamine sulphonic acid (forused by sulphonating (a)-naphthylamine) suspended in water (Clève, Bl. [2] 26, 241; Nevile a. Winther, C. J. 37, 632). Powder, nearly insoluble in cold water. Boiling water converts it into a crimson dye, forming very little naphthol sulphonic acid. Heated with dilute H.SO. (\sigma) naphthol p-sulphonic acid is formed. By heating with strong H.SO., or with water at 160°, (a) naphthol is produced. Dilute HNO, (7 to lo p.c. HNO, forms di-nitro-naphthol, [138]. Cone. HCP forms a chloro-naphthalene sulphonio scid, whence PCl, forms dichloronaphthalene [68°].

Imide C10Ha(SO2H).N<N [1:4].

Explodes on percussion or when heated to by the action of phanyl-hydrasine upon the Triazo-naphthalens-p-sulphonic acid. Formed

sold; disso-benzene-imide, (a)-naphthylamine-p-sulphonic sold, and aniline are formed simul-

taneously: 2C.H. +2C,H,NH.NH,=

C₁₆H₄(N₄)SO₃H + C₄H₅N₄ + C₁₆H₄(NH₂)SO₃H + C₄H₅N₆ + C₁₆H₄(NH₂)SO₅H + C₅H₅N₆H₅. White needles, V. sol. water and alcohol.

Salt.—BaA's: whits silvery plates, v. sl. sol. boiling water. Phanyl-hydrazine salt C4H5N2H4A': long plates; v. sol. alcohol, nearly insol. ether and ehloroform (Griess, B. 20, 1530).

(a) Diazo-naphthalene snlphonio acid

 $C_{10}H_{6} < \frac{N_{2}}{SO_{3}}$. [1:1' or 4']. Similarly prepared from the product of the reduction of (a)-nitro-(a)-naphthalens sulphonic acid formed by sulphonating nitro-naphthalens (Clève, Bl. [2] 24, 512). Yellow crystalline powder. Boiling water gives (1) naphthol (a) sulphonic acid.

(β) Diaso-naphtbalsne snlphonio acid

Microerystalline Formed by diazotising (3) naphthylamino sulphonic acid (formed by sulphonating (B)-naphthylamine). By boiling with HCl, converting into the K salt and heating with PCl, it yields ohloro-naphthalene-sulphonio chlorido [129] (Forsling, B. 19, 1715).

(β).Diazo-naphthalene snlphonio acid

 $C_{10}H_0 < \frac{N_2}{SO_3}$. Formed by diazotising (? $\beta_1\beta_2$). naphthylamino sulphonic acid, itself got by the action of NH, upon Schäffer's (β) maphthol sulphonic acid at 180°. Minute crystals. Converted by treatment with cuprous chlorido into (8) chloro naphthaleno sulphonio acid, whose obloride melts at 110°, and, by distillation with PCl, is converted into (*)-di-chloro-naphthalene, [136°] (Forsling, B. 20, 80).

 $\textbf{Tetrazo-dinaphthy1110.N}_2.\textbf{C}_{10}\textbf{H}_6.\textbf{C}_{10}\textbf{H}_2.\textbf{N}_2\textbf{OH.}$ Formed by diazotising naphthidine. violet dye-stuffs when combined with the sulphonic acids of (3) naplithol. By boiling with

alcohol it yields (aa)-dinaplithyl.

Salts.—C.,H.,N.,SO,*: yellowish plates.—
(C.,H.,N.,Cl.)PCC.; sparingly solublo yellow needles (Nietzki a. Goll, B. 18, 3256).

Diazo-nitro-benzaldoxim chloride C_H_(NO_1)(N:NCl)(CH:NOH) [3:4:1]. Formed. by the action of amyl nitrite and HCl on (3:4:1)nitro-amido-phenyl-acetio acid (Gabriel, B. 15, 837). Plates or needles. Explodes on heating. On heating with alcohol it gives m-nitro-benz-aldoxim C₄H₄(NO₂)(CH:NO11).

p-Diaso o nitro-benzaldoxim obloride C, H, (NO2)(N2.Cl)(CH:NO11) [2:4:1]. Formed, with evolution of CO, by the action of amyl nitrite on a HCl solution of o nitre p amido phanyl-acetic acid (Gabriel a. R. Meyer, B. 11, 826; C. C. 1885, 516). Long red needles. Explosive. By the action of HBr it gives o nitro. p-bromo-benzaldoxim. By hot aleghol it gives o-nitro-benzaldozim C, H, (NO.) (CH:NOH). On oxidation it gives o-nitro-benzaldehyde. Ammonium snlphide redocce it to o amido benzaldoxim

74-Diago-nitro-benzene. Formed by diagotising m-nitro-aniline.

C.H.(NH.).CH:NOH [133°].

Imide C.H.(NO₂).N₂, [52°], p Diaso nitro bensene. Formed by diasotising p-nitro-anilino,

Nitrate C.H.(NO2).N2.NO.: slender needles.

Gives no pp. with PtCl.

Imide C.H.(NO.).N. [71].

m. Diazo-nitro-bensene.p. tthyl tolnide

[3:1] C.H.(NO.) -N.- NELC.H.Me [1:4]. [55°]. From m-diazo-nitro-benzene chloride and othylp-toluidine (Gastiger, Bl. [2] 42, 312). Resolved by dilute HCl into its generators.

p-Diaso-nitro-benssne-p-ethyl-tolnide [4:1] C_sH_d(NO_s) -N_s-NEt.C_sH_sMe [1:4]. [105°]. Yellow needles (Gastiger, Bl. [2] 12, 312). Rosolved by HCl into p-diazo-nitro benzens chloride

and ethyl-p-toluidino.

m-Diaso-nitro-benzene-m-nitro-anilide [3:1] $C_a H_4(NO_2).N_2.NH.C_aH_4(NO_2)$ [1:3]. [195°l Formed by the action of nitrous acid (1 mol.) on m-nitro-aniline (2 mols.). Small red prisms, v. sl. sol. alcohol (Griess, A. 121, 272; Meldola a. Streatfeild, C. J. 51, 107). 1usol. hot aqueous KOI1; but the potassium salt separates as brown erystals from a solution in alcoholic KOH. Cold HClAq gives m-nitro-aniline and m-diazo-nitrobenzene chloride.

p-Diazo-nitro-banzeno p-nitro-anilide [4:1] C, II, (NO.).N., NH.C, II, (NO.) [1:4]. [228°]. Formed by the action of nitrous acid (I mul.) on p-nitro-amiline (2 mols.) (Griess, A. 121, 271; Meldola a. Streatfeild, C. J. 49, 624). Small yellowneedles, m. sol. boiling alcohol. Possesses distinctly acid properties, decomposing Na CO. Cold alcoholio KOH or boiling aqueous KOH form a magenta coloured solution of the potassium salte It forms p-nitro aniline when heated with dilute 11,80, or with water in scaled tubes at 230°. Cono. HCl at 20° gives p-diazo-nitrobenzene chloride and p-nitro-aniline.

Salts.-C.H.(NO.).N.Na.C.11,NO.: forms steel-bluo needles, soluble only in oxcoss of Breet-oldo necentes, Solution only in Carl Albert (C.H. (NO₂).N₁.(C.H. (NO₂).N₂.(C.H. (NO₂)).C.d. (C.H. (NO₂).N₂.(C₁H. (NO₂)).C.d. (C₁H. (NO₂).N₂.(C₁H. (NO₂)).C.d. (C.H. (NO₂).N₂.(C₁H. (NO₂)).C.d. (C.H. (NO₂).N₂.(C.H. (NO₂).N₂.(C.H. (NO₂).N₃.(C.H. (NO₂).N₄.(C.H. (NO₂).(C.H. (NO₂).N₄.(C.H. (NO₂).(C.H. (NO₂

m. Diazo nitro-bsnzsne-p-nitro anllide [3 or 4:1] CaH, (NO2).Ny, NH, CaH, (NO2) [1:4 or 8]. [211°]. From diazotised m-nitro-anilino hydrochloride and p-nitro-uniline; or from diazotised p-nitro-anilins hydrochloride and m-nitro-aniline (Meldola s. Streatfeild, C. J. 51, 103, 439). Yellow needles (from alcohol). Dissolves in NaOHAq, forming a rod solution. Cold squeous 1860 gives m. and p-diazo-nitro-benzeno ohlorides and m. and p.nitro-anilino,

m . Diazo - nitro - benzene . m - nitro - benzyl . anilide[3:1] C.H.(NO2).N2.NC,H.,C.H4(NO2)[1:8]. [142°]. From m-diaz i-nitro-beuzeno-m-nitroanilide, alcoholic KOII, and benzyl chlorids (M. a. S.). HClAq at 100° gives m-chloro-nitrobenzene and m-nitro-benzyl-aniline.

p-Diaso-nitre-benzene · p-nitro-bensyl-anilide [4:1] C₄H₄(NO₄).N_xNC₇H₄,C₄H₄(NO₄) [1:4]. [190°]. From p-diazo-nitro-benzene-p-nitro-benzyl-anilide, alcoholio KOH, and C,H,Cl (Meldola a. Streatfeild, C. J. 51, 112). Minute yollow needles. Conc. HCl decomposes it at 100° into p-nitro-benzyl-aniline and p-chloro-nitro-benide [4:1] C.H.(NO.).N.O.H.O.H.(NO.) [1.8]. [180]. From m-diazo-nitro-benzene-p-nitroinilide alcoholic KOH, and benzyl chloride (Meldola a. Streatfeild, C. J. 51, 114).

m-Diazo-nitro-bonzone-m-nitro-ethyl-anilide (8:1] C,H,(NO,).N, NEt.C,H,(NO,) [1:3]. [119°]. From m.diazo nitro-benzene-m-nitro-anilide by treatment with KOH and Etl. Also from m. diazo. p. nitro benzene und m. nitro-ethyl. aniline. Needles. Cold conc. HCl forms mdiazo nitro benzeno chloride and m-nitro ethylaniline (Meldola a. Streatfeild, C. J. 51, 108,

441). p. Diazo-nitro-benseao p-nitro-ethyl-anilide (4:1) C.H.(NO.).N., NEt.C.H.(NO.) [1:4). [1927]. Formed by the action of Eth and KOH upon p. diazo - nitro - benzene - p - nitro - ethyl - anilido (Meldola a. Streatfeild, C. J. 49, 631). Or from diazotised p-intro-uniline and p-intro-ethyl-aniline (M. a. S., C. J. 51, 111, 442). Yellow needles, v. sl. sol. alcohol, insol. atkalis. Decomposed by cold HCl into p-diazo-nitro benzene chloride and p-nitro ethyl-aniline.

p.Diaso.nitro benzeno m.nitro-ethyl.anllide (4:1) C.H.(NO.).N.N.Kt.C.H.(NO.) (1:3). [187]. From diaxotised p.nitro-aniline and m-nitro-ethyl-aniline (Meldola a. Streatfeild, C. J. 51, 111, 442). Orange needles (from alcohol). al. sol. alcohol. Cold cone. HCl gives m-nitroathyl-amline and p-diazo-nitro-benzene chlorido.

m . Diazo . nitro - bonzeno . p . nitro . ethyl.

anllide (*). [3:1] C,11,(NO.).N,Et.C,H,(NO.) [1:4]. [148°]. Prepared by digesting the potassium salt of mdiazo - nitro - benzene p-nitro-anilide dissolved in alcohol with Etl (Meldola a. Streatfeild, C. J. 51, 105). Small yellow needles. Heated with HCl at 100° it gives m- and p-nitro-ethyl-anilines and m- and p-chloro nitro-benzenes. By cold HOl it is resolved into m. and p. dinzo nitro borzeno chlorides and m- and p- nitro ethyl anilines.

m-Dlazo-nitro-benzene-p-nitro-cthyl-anllide [8:1] C.H.(NO.).N., N.F.L.C.H.(NO.) (1:4). [175°]. From diazotised m-nitro aniline and p-nitro-othyl-aniline (Meldola a. Streatfeild, C. J. 51, Yellow needles (from alcohol). V. sl. sol. alcohol. With cone. IIClAq it forms p-nitroaniline and m-diazo nitro-benzene chloride.

p.Diazo-nitro-benzene-plperidide [4:1] C.II (NO.).N. NC.H. . [97°] (Wallach, A.

235, 264).

m. Diazo-nitro-benseno sulphonic acid 0.11,(0.0) $\stackrel{N_{r}}{\sim} (1)$ $\stackrel{N_{r}}{\sim} (1)$ $\stackrel{N_{r}}{\sim} (0.0)$. Formed by diazotisation

of m nitro aniline sulphonie acid (1:3.6). White microscopic tables. By heating with absolute alcohol under an extra pressure of 400 mm. it is converted into p untro benzene sulphome acid (Limpricht, B. 18, 2186).

Diaso-nitro-bonsene disulphonic acid

C₂H₂(SO₂H)(NO₂) N₂ Formed by diazolising m-nitro-aniline disalphonic acid (Limpricht, B. 8, 289). Boiling alcohol forms mnitro benzeno disulphonio acid.

Diago-nitro-bensoio acid

 $\mathbf{C}_{\mathbf{e}}\mathbf{H}_{\mathbf{s}}(\mathrm{NO}_{\mathbf{r}}) < \frac{N_{\mathbf{s}}}{\mathrm{CO}} >$. Explosive yellow laminæ [Balkowski, A. 173, 63].

formed by dissolving hits amide strategy; bensene (Nietzki a. Benkiser, B. 18, 501).

C₄(NO₂)(N₂OH)(OH)(ONa)O₂aq: long yellow explosive needles.—C₄(NO₂)(N₂OH)(OH)(OH)(OA)O₃; and the constant of the constant yellow explosive plates.

Diazo-nitro-phenol $C_sH_s(NO_2) < N_s > .$ From C₆H₂(OH)(NH₂)(NO₂) [1:2:4] by diazotisation (Gricss, A. 113, 212). Brownish-yellow granular mass; v. sl. sol. hot water, v. sol. alcohol. Explodes at 100°.

Methyl cther.

Nitrate C.H.(NO.)(OMe).N.NO. Formed by diazotising nitro-anisidine (Griess, J. 1866, 459). Platinoohloride | C.H. (NO.) (OMe) N. Cl. PtCl. Porbromide C.H. (NO.) (OMe) N. Br. 1 mide C.H. (NO.) (OMe) N.: needles. Methoxy-nitranilido

C.H.(NO)(OMe).N..NH.C.H.(NO2)(OMe). Formed by passing nitrous acid into ar exchol solution of nitro-anisidine (Grioss, A. 121, 278).

Diazo - di · nitro - phenol $C_6H_2(NO)_2 < \frac{N_2}{O}$. Formed by passing nitrons acid into an alcoholic solution of di-nitro-amido-phenol (picramic acid (Gricss, A. 113, 205). Yellow plates (from nleohol).

Diazo-nitro-8-phenyl-propionio acid [3:4:1] C₀H₃(NO₂)(N₂OH)C₂H₄,CO₂H. Nitrodiazo - hydrocinnamic acid. The nitrate is formed by the action of ethyl nitrite and HNO. on (3:4:1) nitro amido 8 phenyl propionic acid. It forms short colourless needles which explode feebly on heating. Heated with alcohol it gives m · nitro · hydroeinnamio acid (Gabriel, B. 15, 845).

Diazo-nitroso-methyl-nitro-bonzene v. Diazo-NITRO-BENZALDOXIM.

p-Diago-nitroso-oxindole chloride

 $C_sH_3N_4O_2Cl$ or $C_4II_2(N:N.Cl) < \frac{C(N.OH)}{NH} > CO.$

p-Diazo-dioxindole-oxim chloride. Small yellow needles. Prepared by the action of amyl nitrite on a solution of amido oxindole in HCl. Is only slightly attacked by boiling alcohol (Gabriel a. R. Meyer, B. 14, 832; C. C. 1885,

o.Dlazo.p.nitro-tolnone-piperidide C₁₂H₁₈N₁O₂ i.e. [1:4:2] C₆H₃Me(NO₂).N₂.NC₅H₁₆. [51°]. From p-nitro-o-toluidine [107°] by diazotisation of its hydrochloride and subsequent

addition of piperidine (Wallach, A. 235, 218). o-Diazo nitro tolnene p-sulphonio acid

 $C_eH_2Me(NO_2) < \frac{N_2}{SO_2} >$. Formed by dissolving o-toluidine p-sulphonic neid in furning HNO. and ppg. with ice-water (Hayduck, A. 172, 117). Boiling alcohol under 1160 mm. pressure does not attack it.

p - Diago - nitro - tolnene o - sniphonio acid. Formed by dissolving p-toluidine o-sulphonic acid in cold fuming HNO, (Weckwarth, A. 172, 202). Large dark-red prisms. Alcohol heated with it under 1,000 mm. pressure gives nitro-toluone sulphonic acid (Pagel, A. 176, 304).

p.Disao-nitro-tolneno m. snlphonio acid C.II.Me(NO₂)N,SO₂ [1:2:4:5]. Formed hy dis-solving p-toluidine m. snlphonio acid in fuming HNO₃ (Pechmann, A. 173, 214), and also from C₄H₂Me(NO₂)(NH₂)SO₂H [1:2:4:5] and nitrous

said, final, and water; on holing with water comes off; when boiled with Fe.Cl. all comes off. Boiling alcohol does not affect it, but in a sealed tube at 100° it is converted into o-nitrotoluene m-sulphonio acid.

p-Diazo-di-nitro-tolnene o-sulphonie acid

 $C_tHMe(NO_2)_2 < N_2 > 0$ From p-toluidine osulphonio acid and fuming HNO, (Pagel, A 176, 306). Yellowish needles. Not affected by alcohol boiling under 1,000 min. pressuro.

Diago-oxy-acrylic ether (?)
C.H.N.O. i.e. CN.;C(OH).CO.Et (?). (142) at
717 mm. V.D. = 50 (obs.). Yellow oil of strong peculiar odour. Volatile with steam.

Formation .- Gelatine, swollen with water, is warmed with absolute alcohol whilst IICl gas is passed in, it soon dissolves and after disthing off the alcohol a thick brown syrup is left; in latter, which appears to be the hydrochloride of amido-oxy-acrylic ether CH(NH1,):C(OH) CO.Et yields the diazo-ether when its concentrated aqueous solution is treated with NaNO2; it is purified by distillation with steam; the yield is 150 grms. from 400 grms. of gelatine.

Reactions. - Acids eliminate its nitrogen, but it is very stable towards alkalis, with the exception of NII, which even in the cold soon splits off CO, and alcohol. By zino-dust and acetic acid it is reduced first to a hydrazine and finally to an amido-acid. An ethercal solution of iodine converts it into di-iodo-oxy-acrylio ether Cl.:C(OII).CO.Et which on treatment with oold aqueous NII, loses CO₂ and alcohol and yields di-iodo-vinyl-amine Cl₂:CH(NH₂). It reduces AgNO, in the cold (Buchner a. Curtius, B. 19, 850).

Diazo-o-exy-benzeio acid

$$C_{\nu}H_{\nu}(CO_{\nu}H) < {}^{Q}_{N_{\nu}} (CO_{\nu}H) : 0:N = 1:6:3$$
.

C_bH₃(CO₂H)<0 (CO₂H:0:N = 1:6:3).

Diazo-salicylic acid. Formed by passing N₂O₄ into an aqueous or alcoholic solution of hydrochloride of amido-salicy lic acid and concentrating. Crystallised from water (Schmitt, J. 1864, 384; Goldberg, J. pr. [2] 19, 362; P. F. Frankland, C. J. 37, 749). Slender needles. Reactions.—1. Boiled with cono H1 it gives

iodo-salicylic acid [1967] (q. v.). 2. Mercaptan at 170° gives salicylic acid (Schmitt a. Mittenawey, J. pr. [2] 18, 193).
Chloride.— C. H. (CO. II) (OH) N.Cl. — Pla-

\$1 no.oh loride (C.H. (CO.II) (OH) N.CI }.PtCl. Dlazo-p-oxy-benzolc-amido-oxy-benzolc acid.

Dimethyl derivative.

Diazo amido-anisic acid C.H. (OMe)(CO.H).N. NH.C.H. (OMe)(CO.H). Formed by passing nitrous acid gas into a cold alcoholic solution of amido-anisic acid (Gricss, A. 113, 337; 117, 45). Amorphous powder, insol. water, alcohol, and ether. Warm conc. 1980. water, alcohol, and ether. warm conc. HCl converts it into a red acid C₁₀H₁₀O₁... Na_A"1§aq...K_A"2aq...Eth_A": narrow leaflets. Diazo-oxy-cinnamic acid. Methyl deriva-

tive C.H.(N.OH)(OMe).CH:CH.CO.H [5:2:1]. Formed by diazotising methoxy amido-cinnamio acid. The obloride forms yellow crystals, which decompose at about 102°. The nitrate O.H., (N.N.NO.) (O.Me) (C.H., CO.H.) crystallises in medies, which explode at 152°, nearly

a told water, sloobel, and (Schnell, B. 17, 1888).

o-diaso-phonol, Ohlorids C.H.(OH)N.CL By passing N₂O₂ into an alcoholio solution of the hydrochloride of o-amido-phenol, and then adding other (Solunitt, B. 1, 87; Böhmer, J. pr. 132, 460). Rhomboliedra. Bromino water added to its aqueous solution forms a yellow pp. of

diazo-dibromo-phenol (q. v.).

Sulphite C.H. (OH).N. SO.K aq. Golden scales, got by adding KIISO. to C.H. (OH).N.C. (Schmitz a. Glutz, B. 2, 51; Reisenegger, A. 221, 314).

Platinochloride (C.II.(OII)N.Cl),PtCl, m-Diazo-phonol. Ethylderivative (Wagner, J. pr. [2] 32, 70).

p-Diazo-phenol. Nitrate O.H. (OH).N.NO... Formed by passing nitrous acid gas into a cold ethereal solution of phonol (Weselsky, Sits. B. 1875, 9; B. 8, 895), or of p-nitroso-phenol (Jäger, B. 8, 894).

Preparation.—By passing N₂O₄ into alcoholic solution of hydrochloride of p-amido-phenol, adding strong HNO₄ and cooling strongly (Böh-

mer, J. pr. 132, 450).
Reactions. -1. By warming with dilute HBr (15 per cent. solution) it does not give off nitrogen, as diazo-benzene nitrato would do, but forms di-azo-di-bromo benzene in accordance with the equation:

60.41.
 0H + 9HBr = 60.41.
 8C.41.8F.00.4 + 9HBr = 8C.41.8F.00.4 + 9H.4.0 + 9H

Formed by diazotising p. amido -phenol hydrachloride. Converted into hydroquinone by boiling cone, 11Cl or dilute H₂SO₄ (Schuler, B. 9, 1160). Heated with mercaptan it gives phenol and (C.H.) S. & Schmitt a. Mittenzwey, J. pr. [2] 18,

194). -(O_aH₁(01), N_a(D), M(C),

Browide C_aH₁(01), N_aEr. Formed by passing N₂O₃ into an alcoholic solution of the hydrobromido of p-amido-phenol (C. Böhmer, J. pr. 132, 451). Precipitated by ether.

Platinobromide (C.II, (OII)N, Br), PtBr. Needles grouped in spherical segments. Got by adding an aqueous solution of I'tlir, (prepared like PtCl, by dissolving spongy platinum in HBr mixed with HNO, and evaporating) to one of diazo-phenol hydrobromide. If left for several days in contact with their mother liquor, the crystals absorb eight molecules of water of crystallisation changing to blood-rod twin erystals resembling gypsum. These are insoluble in ether, CS, and CHCl, difficultly actable in water. The salt heated with 10 pts. of Na CO, does not yield bromophenol.

Sulphate C.H.(Oll)N. 50,11. by adding dilute H.So, to an alcoholic solution of the hydrochloride of p-amido-phenol, passing in N.O. and adding other. Needles. Does not explode when heated. Converted by boiling HBr into the bromo-phenyl ether of hydro-

quinone (q. v.).
Sulphite HO.O.H., N., SO.K. From p-amido-phenol, HCl, NaNO., and K,SO. (Reisenegger, A., 221, 316). Yellowish plates.

Ethyl derivative "C.H.(OEt)N2OH. Salts. O.H. (OEt) N.Cl. From p. amido-phenetol hydroohloride by solution in alcohol and treatment with N₂O₂. Precipitated by ether as an oil

in the

Crystallises when cooled to -18°. But if H.SO. be added to the alcoholic solotion of the chloride, avoiding rise of temperature, crystals of the sulphate O.H.(OEt) N. SO, H separate. Boiled with water this forms Call (OEt)(OH), hydroquinone mono ethylic ether (q. v.) (Hantzsch, J. pr. 130, 461).

Methyl derivative. The salts are formed by diazotising p-anisidine (Salkowski, B. 7, 1009); C₄H₄(OMe),N₂NO₂, - C₆H₄(OMe),N₂SO₄H₅ Dlaze phenol-carbanic ether. Ethyl deri-

C, II, (OEt) N CO, Et.

Preparation. - By passing N₂O₂ into a solution of the hydrochloride of ethoxy-amidophenyl-urethane, C, II, (OEt) (NII,), NII, CO, Et, HCl (Kohler, J. pr. [2] 29, 273).

Properties. Stender, silvery-white matted needles. Itsol. water, sol. alcohol, ether, and glacial acetic acid. Decomposes below 100°.

Reactions. 1. Does not explode, -2. Not affected by boiling alcohol. -3. Boiled with soda, it is deenaposed with evolution of nitrogen.

Diazo phenol sulphonic acids C.H.(N.OH)(OH)(SO,H) [1:1:3] and [3:4:1] are unstable crystalline acids obtained by diazotising the corresponding amido phenol sulphonic acids (Bennewitz, J. pr. [2] 8, 52).

Diaso phenel disulphonio acid

 $C_a\Pi_z(SO_a\Pi)_z < \frac{N_z}{C}$. From plamido phenol di-

sulphonic acid (q, x) (Wilsing, A. 215, 238). Salt. KA"aq. Small sulphur - yellow needles. Warmed with water it forms hydroquinone disulphonic acid.

p. p. letra-azc-diphenyl

Netrate NO, N. C. H. C. H. N. NO. Formed by passing nitrons acid gas into an alcoholic solution of nitrate of benzidine, and ppg. with ether (Griess, Tr. 1864, iii. 719). White needles, v. sol. water, m. sol. alcohol, insol. ether. Expludes when heated. Boiling water forms p-p-di-oxy-diphenyt.

Perbromide Br, N., C, HeC, H., N., Br, Imide $N_x G_a \Pi_v C_a \Pi_v N_s$, [127]. Whiteplates. Platinochtoride

(CIN, C. II, C. II, N. Cl)PtCl4: yellow plates. Sulphate (HSO,N,C,H,C,H,N,SO,H)H,SO,:

white PhNH.N., C. H., C. H., N., NHPh: Anilide lany-shaped crystals, insol, water. Explosive.

Piperidide. C, II, N.N. C, II, C, II, N. NC, H, From benzidine by diszotisation and treatment with piperidine, Insol, water, st. sol. alcohol, v. sol. ether

(Wallach, A. 235, 271). m.m.Tetra.azo-diphenyl (Brunner a. Witt, B. 20, 1028).

Diago.phenyl-carbamic ether

/N,\ N CO.Et. Formed by diagotising o amido phenyl carbamic ether (Rudolph, B. 12, 1296)

Hoxa-azo-tri-phenyl-carbinel Chloride {C.H.,(N.,Cl)},COH. Diazo-p-resanding chloride. Formed by diazotising prosanilins hydrochloride (B. a. O. Fischer, A. 194, 274).-Ci.H.: N. CI. BAUCI.

Hexa-axo-tri-phenyl-sarbinyl cyanide

Chlorids [C,H,(N,Cl)], C.CN 2aq. From
hydrocyano-p-rosauiline hydrochloride by diazotisation. Slender needles, v. col. water.

Givee, with boiling water, (C,H4OH),C.CN (Fischer, A. 194, 275).

Hexa-azo-tri-phenyl-methane chloride (C,H,N,CI),CH. Diazo-p-leucaniline. Formed by diazotising tri-amido-tri-phenyl-methane-hydroeliloride (E. a. O. Fischer, A. 194, 269). Gives p-rosolic acid (aurin) when boiled with water.

Di-azo-phenyl methyl ketone sulphite of sodinm CH, CO.C, H, N, SO, K. Formed by diazotising o amido acctophonone and treating the product with K.SO.. On reduction it gives the hydrazine salt: CH. CO.C. H. NH.NHSO.K whence IICl forms incthyl-indazol, ,CMe

Colling (v. Indazol). The aqueoue solution of the hydrazine salt changes to tion of the ayarazme sate changes at CMo
Coll N.-N.SO, Na methyl-indazol sulphonate of sodium (Fischer a. Tafel, A. 227, 305).

Hexa-aze-di-phenyl-telyl-carbinel.

Chloride (C,H,N,Cl),C(OH),C,H,MeN,Cl, Diazo-rosaniline. Formed by diazolising rosaniline hydrochloride (Caro a. Wunklyn, Z. 1866, 511; E. a. O. Fischer, A. 194, 279). Gives rise to rosolic acid when boiled with water.—

(C₂₀H₁,N₂(I₂),BPtCl₂fnq. - C₂₀H₁₃N₁Cl₃BAuCl₄.

Hexa.azc.di.phenyl-tolyl. carbinyl cyanide.

Chloride (C₂H₁N₂Cl)₂:C(CN).C₂H₂MeN₂Cl.

Diazohydrocyan - rosaniline. — Gold salt.—

Czoll, (CN)N Cl, 3AuCl,

Diazo-resercin chloride. Diethyl ether C,II,(OEt),N,Cl. From the amido- compound (Pukall, B. 20, 1136). Unslable crystals. Diazo-rosaniline (v. supra).

Diaze-salicylio acid v. DIAZO-OXY-BENZOIC ACIN.

Diazo-succinamic acid

CO.H.CH., CN., CONH. Methyl ether A'Me: [84°]; long yellow prisms (from alcohol). Formed by the action of aqueous NH, npon methyl diazo-succinate. By decomposition with cold slightly acidified water it yields methyl fumaramate and methyl malamate. Heated with benzoic acid at 140°. 150° it gives methyl benzoyl-malamate. Iodine in ethereal solution con crts it into methyl di - iodo - snecinamate — CO. Me. CH₂. CI₂. CONH₄ (Curtiue a. Koch, B. 19, 2460).

Ethyl other C.H.N. (CONH.) (CO.Et .-[1120]; long thin yellow prisms; casily soluble in hot water and alcohol, sparingly in cold water and ether. It is not altered by boiling with pure water, but by acids and alkalie ie at once decomposed with ovolution of nitrogen. Reduced to aspartic other by zinc-dust and acetic acid (Curtiue a. Koeh, B. 18, 1293).

Disso-succinic acid C₂H₁N₂(O₂H)₂. The dimethyl and di-ethyl ethers of this acid are obtained by mixing iced solutions of the hydrochlorides of the aspartic ethers (CO₂R.CH(NI₂Cl).CH₂CO₂R and sodium nitrits, and sodium nitrits, and adding a few drops of dilute H.SO, after wasca sae product is shaken out with other. The ethers form dark-yellow oils which have not yet been obtained in a pure state. By boiling with water or squeous acids they are decomposed with evolution of nitrogen and formation of the corresponding fumario ether. They decompose spontaneously on keeping, evolving nitrogen and forming azinsuccinie ethers (CO,R),C,H,:N.N.C,H,(CO,R), Strong aqueous NH, converts them into diazo succinamic ethers.

p.Diazo-tolnene.

these salts resemble those of the corresponding diazo-benzene salts (Griess, C. J. 20, 86).— C.H.Me.N.NO.; long slender white needles.—
(C.11,Me.N.Cl).PtCl; yellow prismatic crystals.
C.H.Me.N.SO.ll.—C.H.Me.N.Br,
Dicyandle C.H.N., or C.H.Me.N., CN,11CN.
[78°]. Needles or leallets. Formed by the action

of a diazo-toluene salt on a solution of KCN

(3. 42.1638).

Aniliac C. H. Mc. N., NHPh or, alternatively, Ph.N. NII.C. II.Me. From p toluidine and diazobenzene nitrate or Irom aniline and p-diazotoluene nitrate (Griess, A. 137, 60; B. 7, 1619). Narrow yellow leaflets. By warming with phenol it gives a mixture of aniline, p-toluidine, benzeneazo-phenol and p-toluene-azo-phenol; similarly with resorcin (Heumann a. Occonomides, B. 20, 907).

p. Chloro-anilide. Formed from pchloro-diazo benzene and p toluidine. By heating with phenol it gives p-toluene azo-phenol and p-eblorabiline (Heumann a. Occonemides,

B. 20, 909).

Piperidide C12H11N, i.e. C1H1 N2NC H4, [41°]. From CH2C4H2N2Cl and piperidine. Prisms (from alcohol or ether). Insol. water. Dry HCl passed into its solution in petroleumether appears to form an unstable hydrochloride, quickly decomposing into diazotoluene chlorido and piperidino hydrochloride (Wallach, A. 235, 244).

p. Toluide C.H.Me.N. NH.C.H.Me. [116]. Formed by passing nitrous acid gas into a solu-tion of p-toluidine in alcohol and ether (Griess, A. 121, 277; when pure (by digestion with alcoholic (NII.).S) it forms nearly colourless large thin prisms (Bernthsen a. Goske, B. 20, 928).

p. Ethyl toluide Coll, Mo.N., NEt. C.11, Me. Decomposed by acids into ethyl p toluiding and p-diazo-toiueno chloride (Gastiger, Bl. [2] 42,

342).

o-Diazo-tolnene-o-tolnide [2:1] C,H,Me.N,:NH.C,11,Me [1:2]. [510]. Orange-yellow powder of microscopic needles. Prepared by adding sodium nitrite (1 mol.) to an aqueous solution of o-toluidine (2 mols.) and HCl (3 mols.) and then neutralising the HCl with sodium acctate, the temperature being kept below -5° during the whole reaction. It is crystallised by dissolving in cold alcohol and adding ice (Fischer a. Wimmer, B. 20, 1592).

o-Diazo-tolnene-azo-telnene C14H, N.OH i.e. HO.N-N. (?) Obtained by diazotising -N.C.H.

toluene-azo-o-toluidine; the salls erystallise out when a stream of nitrous acid gas is passed into an alcoholio solntion of toluene-o-azo-

toluidine and an excess of sold, or upon aubsequent addition of a little other. On heating with water or alcohol it decomposes, evolving nitrogen like ordinary diano-compounds. By SnCl, or SO, it is not reduced to a hydrazine but gives a stable compound C₁₄H₁₁N, which N-NH

probably has the constitution C.H. N - NC.H. hy bromine this body is reconverted into the diazo perbromide. By zinc dust and alcohol it Salts.—The preparation and properties of its converted into m-teluene-p-azo-toluene [589] with evolution of nitrogen. The diazo-imide loses nitrogen on heating and yields tolyl-

> N/ N.C.11, identical with azimido toluens C,H,<

that obtained by oxidation of toluene-azo-o-toluidine. o-Diazo-teluene-azo-toluene reacts with amines and phenols like an ordinary diazocompound; the products, however, reduce to a diamine or amido phenol and tolyl azimidotolnene. All the salts have a deep orange yellow colour and are tolerably stable. - Cull N. Cl. : convariant and are tolerany sames. - C₁H₁₁N₁Cl³; yollow acionlar crystals. - C₁H₁₁N₁Cl³; yollow acionlar crystals. - C₁H₁₁N₁Cl³; slender red pointed crystals. - C₁H₁N₁S₀Sl₁H²; red interwoven needles. - C₁H₁N₁B₁; [125], long distanting and problems are constant absorbed. glistening red needles or compact crystals.

Imide CallaN,: [85], thick yellowish red crystals; formed by the action of alcoholic NH, upon the perbromide (Zineke a. Lawson, B.

19, 1452).

Product of Reduction Callan, probably N-N11 [168°]. Long colourless

 $C_{:}H_{\bullet} \Big \langle \Big| \begin{array}{c} | \\ N-N.C_{:}\Pi_{\bullet} \end{array}$ V. sol. hot alcohol, sl. sol. ether ncedles. It lus 20 and chloroform, insol. water. lt is not affected basic properties. reducing agents. Bromine in alcoholie or acetic acid solution readily converts it into o-diazotoluene-azo-toluene perbromide. On addition of Ag₂O to its alcoholic solution nitrogen is evolved and m-p-azotolnene [58°] is formed.

Acetyl derivative Cullan, Ac [184°]: glistening white plates (Zincko a. Lawson, B.

19, 1457).

p. Diago-toluene-ago-toluene
[2:1] C₄H₄(CH₂)-N₂-O₆H₄(CH₂).N₂OH [1:3:4].
Prepared by dissolving toluene-ago-p-toluidine in alcohol, adding an excess of 11Cl, diazotising by passing N.O. into the well-cooled solution, and precipitating the diazo-salt with ethor. By reduction with SnCl, or zinc-dust and acetio acid in cold dilute aqueous solution it is split up (without formation of a hydrazino) inte o-toluidine and tolylene-p-diamine. Salts.—*C₁₁H₁₁N₁N₂N₃: slender brownlsh-

yellow needles, m. sol. water and alcohol.-C. H., N. Br.: yellow crystalline pp. which changes on standing to small violet needlos .-C₁,H₂N₂SO₂Na: glistening scales (from alcohol), v. sol. hot. alcohol, sl. sol. water.

Imide C₁₁II₁₁N₁: (60°); long plates; sol. alcohol and acetic acid (Zincke a. Lawson, B.

20, 1181).

o.Diazo-toluene m-sniphonio acid *MeC,H, $\langle {}^{N;N}_{SO_2} \rangle$ [1 2]. Precipitated as

white powder when nitrous gas is passed into a

ecild solution of o-foluidine sulphonic seid. Explodes feebly at 100° (Nevile a. Winther, O. J. 87, 628).

o-Diago-toluene p-sulphonic acid.

Minute monoclinio priems (Haydnek, A. 172, 213; 174, 3il). Boiling alcohol produces Ms.C, H, (OEt).SO, 11 [1:2:4].

p-Diazo toluene o-snlphonic acid. er brown needles (Ascher, A. 161, 8; Jensen, A. 172, 235). Heated with alcohol under pressurs it gives McC, H, (OEt) (SO, H) Remsen a. Palmer, Am. 8, 243).

p-Diazo-telnene m-sulphonic acid

 \times McC_aH_a<N:N> [1 $\frac{4}{3}$]. More soluble than the cerresponding o compound (Nevile a. Winther, C. J. 37, 631). Prepared by passing nitrous acid gas into p-toluidine sulphonie acid suspended in alcohol. Hot alcohol gives toluene m-sulphonie acid (Petermann, A. 173, 201).

p-Diaze-toiuene exo sulpheaie acid

O.H. CIL SO. Heated with alcohol under 1,100 mm. pressure it gives Call, (OEt).CH2.SO411 (Mohr, A. 221, 219).

p. Diaze - telnene - sulphonio - amido - telnene

sniphonic sold. Amids (14:2) C.II.Met 8'.NU.NI.N.,NII.C.II.Met So.NII. (2:1:1). From C.II.,Nic(NII.)SO.JNII. (1:2:1), alcohol, and nitrous acid gas (Physan, A. 221, 211). Decomposod by 11Cl into N₂, C_aH₃ClMe.SO₂NH₂, and C_aH₂(NH₁)Me.SO₂NH₃.

o Diazo-tolnene disniphenio acid

 $O_aH_aMc(SO_aH)$ $\stackrel{N_a}{\underset{SO_a}{\mid}}$ $\begin{bmatrix}1.8:\frac{2}{3}\end{bmatrix}$. From N_2O_a and aqueous o-toluidine disalphonic acid at U^o (Lim-

prioht, B. 18, 2176; Hasse, A. 230, 291). Microscopic needles. Explosivo. Heated with alcohol under pressure it gives C₆H₂Mc(OEt)(SO₄H)₂. Witt HI it forms C₁H₂Mc(SO₂H)₂.

Salts .- KA' .- BaA', 4aq. - PbA' p-Diazo teluene di-salpheaio acid

N, C_aH₂Me(SO₃H) . Formed by diazetisation

of p. toluidine - di - sulphonie acid. Yellowish crystals. By heating with III it yields p-iodotoluene-di sulphonio acid; with HBr it yields p-bromo-tolnene-di-sulph nie neid.

Salts .- A'K: large yellow prisms .- A', Ba: yellowish white needles. - A', i'b; small red prisms (Limpricht, B. 18, 2178).

Call, Me(CO, II). N., NII. C. 11, Me. CO, II. amido-toluic acid and nitrous ether (Griess, 4. I17, 59). Minute yollow prisms (coataining aq); insol. water, alcohol, and other.

o. Diaze. p. teinidine bromide. Accept derivative

C.H.Mc(NHAc).N.Br [1:4:2].
From C.H.Mc(NHAc)(NH.) [1:4:2], cone. HBr

Tom Call, ste(Stract), the state of the stat

PROF CH. MajNHam N.Cl. gives C.H.Me(NHAe)OL

Diazo-m-xylene-sulphonie acid

 $C_{e}H_{e}Ms_{2} < \frac{N_{2}}{SO_{2}} > [1:3:4:6]$. White pp. Sparingly soluble in water. Decomposes at 600-700. Combines with phenols and amines. Formed by diazotisation of m-xylidins-sulphonio acid (Nölting a. Kohn, B. 19, 138).

Diazo-p-xylene-enlphonic acid

 $C_5H_3Me_2 < \frac{N_2}{SO_3} > [1:4\frac{2}{5}]$. Yollowich whits plates. Stable at ordinary temperature, decomposss on heating with water at 60°-70°. Formed by the diazotisation of p-xylidine-sulphonio acid C_aH₂Mc₂(NH₂)(SO₂H) [1:4:2:5] (Nölting a. Kohn, B. 19, 141).

DISAZO- COMPOUNDS. Secondary azo- compounds. Compounds containing two azo- groups of the form C-N2-C. The general methods by which they may be propared are given in the article on azo-colouring matters (p. 368). The nomenclature here used is like that used for azo- compounds. To find the name of a disazocompound, write down the formula, strike ont everything between the two N, groups, remove one of the N₂ groups, and join the remaining parts of the formula together and name the resulting azo- compound as directed on p. 369. Then insert after the word 'azo' the name of the central hydrocarbon, preceded by prefixes representing its substituente and followed by

Di-amide-bsnzene-azo-bsnzens-azo-benzsne salphonic acid

[4:1] C₄H₃(SO₃H)—N₂—C₆H₄—N₂—C₆H₂(NH₂)₂ [1:2:4]. Formed by the combination of diazobenzene-azo-benzene-p-oulphonio acid with mphenylene diamine. Red microscopic needles. V. sl. sol. alcohol and other. In H.SO, it dissolves with a violot-blue colour.-KA'2aq:red glistening plates, al. sol. hot, v. ei. sol. coid, water, dyes silk, wool, and cotton a brownishrcd (Griess, B. 16, 2035).

Amide-sniphe-naphthalene-azo-diphenyl-azenaphthylamine sniphenic acid

[1:4:2] $C_{10}H_3(NH_2)(SO_3H) - N_2 - C_8H_{50}$

[1:4:2] $C_{10}II_{5}(NII_{2})(SO_{4}II) = N_{2} - \dot{C}_{6}H_{2}$ Formed by combining diazotised benzidine with (a)-naphthylamine p-sulphonic acid. Dyes cotton from an alkalino bath scarlet, turned blue by a trace of acid. The aqueous solution is readily reduced by NIL, and zinc-dust, giving benzidine and naphthylene-o-di-amine enlphonio acid (Witt, B. 19, 1719).

Benzene-aze-m-diamido-benzene-azo-benzene O, 11, -N, -C, H, (NH), -N, -C, H, [250°]. ornied by the combination of dinzobenzene with chrysoïdine. Dark red needles or eixsided plates. Sol. hot chioroform and benzene, v. el. eoi. alcohoi and ether, insol. water. Weak base. Salts .- B'fiCl: violet-brown amorphous solid. —B',H,Cl,PtCl,: violet brown amorphous pp. (Grices, B. 16, 2028).
Benzenc-arc-m-diamido-benzenc-arc-benzene

p-sulphonic acid

C.H.(SO.H)-N2-C.H2(NH2)2-N2-C.H2. Formed by the action of p-diago-bensens-sul-phonic acid on chrycoldine. Dark-brown mi-eroscopic crystals. V. al. sol. alcohol insulsther. A'K: reddish-brown plates, sol. hot; al. sel. cold, water (Griess, B. 16; 2032).

Benzene-azo-di-amide-benzene-azo-benzele acid C,H,(CO,H)-N,-C,H,(NH,),-N,-O,H, Formed by combining m-diaze benzeio acid with chryceidine (Griess, B. 16, 2032). Browniehred powder. Insoluble or nearly insoluble in all ordinary solvents. Soluble in elkalis with a brewnieb-red colcur.

(a)-Benzsne-azo-m-di-amlde-benzene-ezo-tolnene C₂H₃-N₂-O₂H₂(NH₂)₂-N₂-C₁H₁. [192°]. Formed together with a small quantity of the (8)-isomeride by the combination of p-diazobenzenc with p-toluene-aze-phenylene-diamine. Dark-red glistening neodles. Sol. ether, chloroform end hot benzene, insel, water and alcohol.

(8) - Benzene - azo-m - dl - amido - bonzene - azotolnene C₆H₁-N₂-C₆H₂(NH₂), N₂-C₇H₃. [225°]. Slender yellow needles. Sol. alcohol and ether, and sel. chloroform, eel. in water (Griess, B. 16, 2029).

Benzane-aze-m-di-amido benzane-p-azo. toluene C, II, -N₂ -C₄II₂(NII₂), -N₂ · C₄II₃. [214°]. Formed by the combination of p-diazotoluene with chrysoïdine (Griess, B. 16, 2030). Dark-red glistening needles. Sl. sel ehleroferin, ether, and benzene.

Benzene-azo-bsnzsne-azo-p-oresol

C.H. -N. -C. II. - N. -C. II. (CfI.) (OH). [160°]. Obtained by the action of diazo-benzene-ezobenzeno oliloride (by diazotising benzene-uzoaniline) on an alkaline solution of p-cresol (Nölting a. Kohn, B. 17, 354). Small brown needles. Sl. sol. alcohol, m. sol. chloroform, benzono, and acetic acid. Dissolves in H2SO, with a reddish violet colcur.

Benzene-azo-benzene-azo-ethyl-(3)-naphthylamine C₄H₄-N₂-C₆H₄-N₂-C₁₆H₄.NHEt. [142°]. Small red needles. Formed by heating ethyl-(B)-nephthyl-nitrosamine with an acetic acid solution of benzenc-azo-aniline (Henriques, B. 17, 2670).

Benzene azo-benzene-azo-(6)-naphthol C₆H₅-N₂-C₆H₁-N₂-C₁₆H₆OII. From diazotised benzene azo aniline and (\$\beta\$) naplithol (Nietzki, B. I3, 1838). Brick red powder.

Banzene-azo-benzene-azo-rasoreln O.H. -N. -C.H. -N. -C.H. (OH). Bythe action

of diazetised benzene-azo amilino on resorcinol two isomerides are formed which are separated by their different solubilities in aqueous alkalis.

(a)-Compound [181°]. Brownish red powder consisting of microscopic tables. Disselves with a carmine rod colour in aqueous NaOH and in H2SO4. Sol. alcohol, ether, and chleroform.

(B)-Compound [215°]. Brown powder. Dissolvee in alcoholic NaOH with a violet-blue colour and in H.SO, with a pure blue colour, v. sl. sol. alcohol, ether, and chloroform, insol. aqueeue NaOH (Wallach, B. 15, 2817).

Benzene-azo-methyl-pyrrol-azo-benzens $C_sH_3-N_2-C_sH_sNMe-N_2-C_sH_s$

C.H.N.C=CH NMe C.H.N.C=CH probably [196°]. Fermed by

the methylation of benzenc-azc-pyrrol-benzene. Red plates (O. Fischer e. Hepp, B. 19, 2258). Voz. L.

Benrens-ano-ary-bennens-ans-benrens

C₂H₂-N₂-C₃H₄(OH)-N₂-C₄H₂. Bensene. (181°). Sormed, together with benzene-azo-phenol, by treeting diaze-benzene nitrate with BaCO, in the cold, or by the action of diazo-benzene nitrate upon a solution of benzeno-azo-phenol in KOHAq (Griess, A. 137, 86; B. 9, 628). Brewn lustroue needles or plates (from aloohol). V. sl. sol. water, v. sol. KOLAG, v. sl. sol. NH, Aq. insol. Na,CO,Aq.

Methyl cther Cullis N. (OMe) [110°], small yellow crystals, v. sol. ether, benzsne, acetone. and hot alcohol.

Acetyl derivative C. H. N. (OAc) [1160]. small yellow needles, sol. alcohol, other, and beuzene.

Benzoyl derivative C14H11N4(OBz) [1392], small yellow needles, st. col. cold clochol (Nölting a. Kohn, B. 17, 368).

Benzene-azo-di-exy-benzene-azo-benzene C_sH_s—N₂—C_sH₂(OII)₂—N₃—C_sH₃. By the action of diazo benzeno obloride on an alkeline solution of benzene-aze-resorein two isomerides are formed which are separated by their solubility in aqueous alkelis. A third isomerido (7). is formed, together with benzene-azo-resoroin, by the action of diazo-benzeno chlorido on resorein treated with KOH (1 mol.) in dilute aqueous solution.

(a) Compound [215°]. Brewn felted needles. Dissolves easily with a red colour in aqueens NaOH and in H₂SO. Sl. sol. alcohol and ether, m. sol. chloroform. Its di-acotyl derivative forms brown glistening needles, [1849] (Wallach).

(B)-Compound [220°]. Mieroscopio needles. sol. alcohol and ohloroform, insol. aqueens NaOH. Dissolves in H.SO, with an indigo-blue colour, and in alcoholio NaOli with a red colour (Wellach, B. 15, 2816).

(7)-Compound [222]. Large red needles Sol. chloroform, v. sl. sol. alcohol. It diesolvez in strong alkalis with a brownish-yellow eclour; in Il SO, with the same colcur. By tin and HCl it is reduced to aniline and disemidoresorciu.

Di-acetyl-derivative C₁₈H₁₂N₄(OAc)₂ [138°], orange needlee (Liebermonn c. Kostanecki, B. 17, 880).

Benzsne-ezo triozabenzene aze-benzene C₄II₄—N₂—C₆H(OII)₅—II₄—C₆H₃, Phloregluein-bi-liazo-benzene, Yollowish-brown leaflets. Prepared by the action of diazobenzene nitrate on phlerogluein (Weselsky a. Benedikt, B. 12 226).

Benzsne-azo-dioxy. venzeno-azo - naphthalene C.H. N. C.H. (OII) N. C. H., [155°]. From diszo-benzeno chloride and an alkalino solution of m di-oxy benzene-azo naphthalone (Wallach, B. 15, 22).

Benzsne-azo-dl-oxy-benzene-aze-tolnene C₀H₃-N₂-C₀H₂(OH)₂-N₂-C₀H₄(CH₃) {1:4}. Prepared by the action of diaze benzenc chloride on an alkaline solution of m-dicxy-benzene-azotoluene, er ef diaze-toluene ehleride en an alkaline solution of benzene-aze-resercin: in either ease the same three isomerides are zimultanecuely produced and ere separated by means of their different solubilities.

(a) Compound [1960]. Golden brown needles. Discolvez with a red coleur in H.SO, end NaOH.

Sel. sloohol and chlorotorm. In all scenario derivative forms yellow needles, [176°].

(c')-Compound [241°]. Dissolves with a red solour in H.SO, and in aqueous NaOH. Its di-acetyl derivative forme yollow needles, [196°].

(s).Compound [206°]. Brownish - black microscopic crystals. Insol. aqueous NaOH, dissolves in 11,80, to a blue solution. Sl. sol. alcohol, m. sol. chloroform (Wallach, B. 15, 2821).

Benzene-azo-oxy cymene-azo-benzene

(2) (1)(4) (3) (5)

\$\mathbb{G}_1\mathbb{H}_4\mathbb{H}_2\mathbb{H Fe.Cl., it is converted into oxy-thymoquinone.

Benzene.azo-o-oxy-toluene-ezo-henzene (1) (3) (1) (5) Q₄H₅-N₅-C₈H₁Mc(0H).··N₂-C₈H₅(115°]. Obtained by the action of (2 mols. of) diazobenzeno oldoride on an alkaline solution of (2 mols. of) diazobenzeno oldoride on an alkaline solution of (3 mols. of) (12 mol). o-oresol (Nölting, B. 17, 364). Reddish-brown plates. V. sl. sol. cold alcohol. Dissolves in alkalis with a yellowish red colour.

Acetyt derivative [121°], yellow needles, v. sol. elcohol, other, and benzene.

Benzene-azo-m-oxy-tolnene-azo-benzene

 $C_4 11_5 - N_2 - C_6 11$, $Me(OH) - N_2 - C_6 11$, [149°]. Obtained by the action of (2 mols. of) diazobenzone chloride on an alkaline solution of m-cresol. Roddish-brown plates. Sol. ether, benzene, and hot alcohol, sl. sol. cold alcohol.

Acetyl derivative [157°], small yellow. ish-brown needles (Nölting a. Kohn, B. 17, 367).

Benzene azo dl. phenyl nree azo benzene O.H. -N, -C.H. NH.CO.NH.C.H. -N, -C.H. (270°). Formed by the action of carbon Formed by the action of carbonyl shloride on benzene-azo-aniline (Berju, B. 17, 1404; C. C. 1881, 871). Small plates. Sol. shloroform and benzene, sl. sol. alcohol.

Benzene are di-phenyl thie nrea are benzene C₁H₄-N₂-C₄H₄,N11.CS.NH.C₆H₇-N₄-C₄H₄. [199°]. Formed as a by product of the action of phenyl-mustard oil on benzene-azo-aniline (Borju, B. 17, 1405). Sl. sol. hot chloroform, zylene, and acctic soid, v. sl. sol. alcohol, bensene, and CS.

Benzene-azo-pyrrol-azo-henzene

 $C_{c}H_{s}, N_{s}-C_{c}H_{s}N_{s}-N_{s}-C_{c}H_{s},$ $C_{c}H_{s},N_{s},C_{s}-C_{c}H_{s},$ $C_{c}H_{s},N_{s},C_{s}-C_{c}H_{s},$ $C_{c}H_{s},N_{s},C_{s}-C_{c}H_{s},$ $C_{c}H_{s},N_{s},C_{s}-C_{c}H_{s},$ $C_{c}H_{s},N_{s},C_{s}-C_{c}H_{s},$ $C_{c}H_{s},N_{s},C_{s}-C_{c}H_{s},$ [131°]. Obtained

by combining (2 mols. of) diazo benzeno chloride with (1 mol. of) pyrrol in alkaline solution. Red crystallino solid. Sublimable. M. sol. other and benzene, el. sol. alcohol, nearly insol. water. Possesses basic properties. Dissolvos in dilute 1tCl with a reddish-yellow colour; in cone. 11,80, with a splendid hlue colour. Its alcoholic solution is turued magonta-red by NaOH, reddish-violet by cone. HCl (O. Fischer Hspp, B. 19, 2251).

Benzene-aso-pyrrol-(β)-azo-naphthalene C_eH₁,—N₂—C₄H₄N—N₂—C₁₆H₇

C,H,N,C=CH NH | Brobably [151°]. Formed C.H.N.C. CH

by the hombination of disco-benefic chievith pyrrol-(6) assembly habitation or of (6) disco-benefic alphibates deal-this chief. in alkaline alcoholic solution. Red plates, with bluish reflection. Sl. sol. alcohol (O. Fischer Hepp, B. 19, 2256).

Tri - bromo - benzene - azo - di - phenyl - diizoindole-azo-tri-bromo-benzene

C₆H₄—N₅—C₆H₄Br₅ [150°]. Grange yellow prisms. Soluble in mean ordinary solvents except water. For med by the action of tri-bromo diazo benzene ohloride on di . phenyl . di - iso · indole. - B"H2Cl2: slender yellowish-brown neodlee (Möhlau, B. 15, 2490).

Di - bromo - oxy - benzene - azo - di - phenyl - di isoindole-azo-di-bromo-phenol.

C.H. -N2 -C.H2Br2(OH) 0,H,O-`он CaHaN Br.O.or HĊ

C.H.-N.-C.H.Br.(OH) [198°]. Yellowish green prisms. Soluble in alcohol, dyos wool orango and silk yellow. Formed by the action of di-bromo-diazo-phenol on di phenyl di isoindole. - B"H.Cl; ahort motallic glistening prisms, insol. water (Mohlau. B. 15, 2192).

ψ-Cumene - azo - m·di - oxy - benzene . azo - ψenmens

C₄H₁Me₂-N₂-C₆H₂(OII)₂-N₂-C₆H₂Me₂. Formed, together with cumene azo resoroin, by combining diezo-cumene ohloride (from antido-pseudo-cumenc (62°)) with resorcin (Liebermann a. Kostanecki, B. 17, 882). Small red needles. Dissolves in H₂SO, with a red resordin colour. Insoluble in alkalis.

p. Di-methyl . amido - benzene . p-azo - benzene-

aro. (8) naphthol

HO. C₁₀H_e—N₂—C₂H_e—N₂—C₂H_e,NMe₂ [210°],

Got by pouring a diazotised solution of p-amidobonzeno-azo-dimethylaniline hydrochloride into on the state of t ln alcoholio NaOH, red; in cono. H.SC, green, turned blue by dilution. An alcoholic solution is turned blue by HCl.

p. Di-methyl-amide-benzene-p-aso benzeneaso.(c).naphthol HO.C.,H.,N.,C.,H.,N.,C.,H.,NMe, Prepared like its (\$\textit{\beta}\) isometries (\$\text{M}.) Its properties are similar, except that the solution in alcoholic KOR is violet. It decomposes below 200°.

Manchyl - unlit - belings - & (HO) O.H.—N.—C.H.—N.—C.H., NMe. Brown powder. Decomposed before melting. Sl. sol. boiling alcohol, the solution being reddishorange and turned first violet, then blue by adding HCl. Sl. col. glacial acetic acid, the solution being red when hot, violet when cold. Insoluble in tolnene. Solution in alcoholic KOH is reddish-violet. Solution in cono. H.SO.

is violet, becoming blue on dilution (Meldola, C. J. 45, 110). Di-methyl-amido-benzene-p-aso-bensene-asophonol

HO.C.H.-N2-C.H.-N2-C.H.NMe2. powder, forming a brown solution in aquequa KCH, and a red solution in alcoholio KOH. Resemblee the analogous di-methyl-amidobenzene · p · azo · benzene · azo · resorcin (q. v.)

(Mcldola, C. J. 45, 111).

Di-mes rl · amido · benzene · azo · tolnene · azo-

(A)-naphthol

(8)-naphthol (2) (1) (4) C₀H₄(NMe₂)-N₂-C₆H₃Me-N₂-C₁₆H₄.OII. From diazotised di-methyl-amido-benzeno-azop-toluidine and (3)-naphthol (Wallach, A. 234, 358). Red needles (from chloroform), insol. water.

Di-methyl-amido-beazeae-azo-tolueae-azophenol (3) (1) (4) C.H.(NMe.)—N.,—C.H.,Me—N.,—C.H.,0M-[160°]. From diazotised C.H.(NMe.)—N.,—C.H.,MeNil., and phenol (Wallach, A. 234, 357). Orange phenol

(a) -naphthalene - azo - pyrrol - (a) - azo - naphthalene $C_{ip}H_1 - N_2 - C_4H_2NH - N_2 - C_{ip}H_1$, $O_{ip}H_1N_2 - C_{ip}H_1$, $O_{ip}H_1N_2 - C_{ip}H_2$

NH | C, H, N, C=CH probably

Formed by adding (a)-diazo-naplithalone obloride (2 mole.) to an alkalino solution of pyrrol (I mol.).
Metallio glistening needlee. Sol. alcohol with a
dark yellowish red colour. Dissolvoe in conc. H.SO, with a blue colour (O. Fischer a. Hepp, B. 19, 2255).

(β)-Naphthalene-aso-pyrrol-(β)-azo-naphthal-C₁₀H,-N₂-C₄H₂NH-N₂-C₁₆H₄. C, H, N, C=CH

NII | . [288°]. Formed by C, II, N, C=CH probably

adding (8) diazo-naphthalene chloride (2 mols.) adding (8)-diazo-napnnanene eniorius (z mois, to an alkalino solution of pyrrol (I mol.). Glistening coppery plates. Sl. sol. alcohol. Tha alcoholie solution is turned reddieh-violet by cone. Hol. Disselves in cone. H.SO, with a bine colour (O. Fischer a. Hepp, B. 19, 2255).

m. Mitro-benzene-p.aso-benzene-(2)-azo-(3)-azo-(3)-azo-(4).

maphthol NO₂C₆H₄-N₂-C₆H₄-N₂-C₁₆H₄-OH. [0.218°]. From diazotised NO₂C₆H₄N₂-C₆H₄NH₄ and (8)-naphthol. Small yield (Meldola, C. J. 45, 113). Orange crystals with green lustre. Solutions in C.H.O. and in toloene are orange; in alcoholic NaOH, violet; in conc. H.2SO, green, turned blue on dilution.

m-Nitro-bensene-(a)-azo-naphthalene-(a)-azo-(a)-maphthol

NO.C.H.—N.—C.,H.—N.—C.,H.OH. A dark amorphous powder. Solutions in tolnene, chloro-form and glacial acetic acid are red; in cone. H.SO. dark indigo violet, becoming blue on the cone in alcobolic potash, greenish-blue Meldels, C. J. 45, 116). 57 Ex

(A)-naphthol

des militares. No.

NO. C.H. N. C. H. N. C. H. OH. From m-nitro-bensene (a) -aso (a) -naphthyl. amine by diazotising and treating with (8).
naphthol (Meldola, C. J. 45, 115). Minute
bronzy needles (from toluene). Blackens at 245°. Insol, alcohol or glacial acctic acid. Solutions in chloroform and in hot suilino are violet; in tolueno red when hot, reddish-violot when cold; in boiling alcoholio Koll, blue; in conc. IL,SO4, olive colour, on dilution, blue and then violet.

m-nitro-benzene-(a) - azo-naphthalene-azoresordin NO₂O₆H₄ N₂—C₁₆H₃—N₂—C₆H₃(OH)₄. Bronzy powder, not very soluble. Solutions in boiling alcohol are reddish; in glacial acotic acid, toluene and eliloroform, orange; in aquecos or alcoliolio KOH, blue; in conc. H.SO., green, changing to bluish groen on dilution (Meldola, C. J. 45, 116).

p. Nitro-benseae-azo-m-xylene-aso- (a) naphthol NO₂C₄H₄-N₂-C₄H₄Mc₂-N₂-C₁₀H₄OH.
Preparation and propertice are eimilar to those of the preceding (3)-compound.
Sulphonic acid

NO, C, H, -N, -C, H, Mo, -N, - C, H, (SO, H) (OH) Similar to the corresponding (8)-compound, but dyes reddish brown.

p - Nitro - benzene - azo - m - xylene - xz - β naphthol

NO_TO₄II₄—N₂—C₆II₄Mo₂—N₂—C₁₀H₄·OH. [278°]. From NO_TO₄H₄—N₂—O₆H₂Mo₂NH₂ by diagotising and treating with eodium (3)-naphthol (Meldola, C. J. 43, 434). Green seales (from tolueno). Scarcely coluble in alcohol or glacial acetic acid. Forms a crimeon solution in boiling aniline or nitrobenzene, and a green colution in conc. ILSO, turned violet by dilution.

Sulphonic acid NO, C, H, N, C, H, Me, N, C, H, (OH) SO, H. Gotby using O, H, (OH) SO, H. Dyes olaret. red.

p. Nitro-benzene.azo-m-xylene-azo-phenol NO₂C₄H₄-N₂-C₄H₄Me₂-N₂-C₄H₄.Olf. Orange powder. Sl. sol. alcohol and bonzeno, v. sol. hot aniline. Solutions are orange. Solution in alcoholic NaOH is roddish-violet. Solution in

cono. H.SO₂ is blae (Meldola, C. J. 43, 436),

p. Mitro-benrene-azo-m.xylene-azo-resoreia
NO. O.H.—N. — C. H. Me.—N. — C. H. (OH),

[2319]. Brown powdor. Forms orange solutions in boiling alcohol, tolueno, and glacial acetic acid. Solution in alcoholic NaOH is red, turned violet by excess of NaOH. Conc. H.SO, forms a blac solution (Meldola, C. J. 43, 430).

p. Oxy-benzene. p. aso benzene. (a) aso. (a) haspa-thol C.H. (OH) - N. = C.H. = N. = C. H. OH.

From diazotised p. amido benzene azo. (a) anaphathol and an alkaime colation of phenol (Meldola, C. J. 47, 665). Dark amorphous powder. Its solution in conc. H. SO, is indigo. blue; in boiling toluene, orango; in alcohol, red; in KOHAq, dull red; and in alcoholts NaOH, deep claret colour.

p. 0xy. bearene. p. szo. benzene. (a). sso. (8). naphthol HO.C., H. N. C., H. N. C., H. OH. (225°). From p. amido. benzene. aso. (8). naphthol by diazotising and mixing with an alkaline solation of phenol (Meldola, C. J. 47, 666). Bed warty concretions. Its solation in conc. H.80. is blnish-green, and becomes violet on dilution

Its solution in boiling toluene, or boiling alcohol. Is red. Its solution in KOHAq is reddish-violet. m-Di-oxy - benzene - p-azo-benzene-(a)-azo-(a)-

naphthol

[4:2:1] C₃H₄(OH)₄—N₂—C₄H₄—N₃—C₄H₄(OH). Formed by mixing diazotised p-amido-benzencazo-(a) naplithed with a solution of resorcin in dilute NaOH (Meldola, C. J. 47, 665). Bronzygreen powder; sl. sol. toluene and acetic acid, forming a red solution. Its solution in KOllAq is blue; its solution in NH, Aq is violet. It is decomposed by heat without fusion.

 $m ext{-}\mathrm{Dl}\cdot \mathrm{oxy.henzene}\cdot p\cdot \mathrm{azo.henzene}\cdot (a)\cdot \mathrm{axo}\cdot (\beta)\cdot$

naphthol

[4:2:1] C_aH₁(Oll)₂-N₄-C_aH₁-N₂-C_{ba}H_cOH. From diazotised p-amido-benzene-azo (β)-naphthol and an alkaline solution of resorein (Meldola, C. J. 47, 606). Bronzy-green powder. Its colu-tion in conc. 11,804 is deep bluish-green; in dilute aqueous alkalis, violet. Sl. sol. toluene forming a red colutioo.

m.Oxy-housene-azo-henzene-p-azo-phenol

[8:1] $110.C_0H_1 - N_2 - C_0H_1 - N_2 - C_0H_1 OH [1:I]$. Dark powder. Easily soluble in alkalis. Formed by diazotising anido benzene m azo-phenol (C_aH₄(OH) -- N₂ -- C_aH₂NH₂) and combining it with phenol (Wallach a. Schulze, B. 15, 3021). p.Oxy henzene azo henzene p.azo phonol

[4:1] $10.2\Pi_4 - \Omega_2 - \Gamma_0 \Pi_1 - \Omega_2 - \Gamma_0 \Pi_1 + \Omega_1 - \Gamma_0 \Pi_1 - \Omega_2 - \Omega_2 - \Omega_1 - \Omega_1 - \Omega_2 - \Omega_2 - \Omega_2 - \Omega_2 - \Omega_2 - \Omega_1 - \Omega_2 - \Omega_2$ alkali (Meldola, C. J. 47, 660). Amorphous brown powder. Its solution in conc. II SO, is violet, unchanged on considerable dilution; v. sol. NH, Aq and KOHAq forming a red solution; insol, boiling toluene; forms an orange solution in hot pluenol or amilino.
m. Dl. exy-henzone. p-azo-henzone. szo-re

zoroln

[4:2:1] $C_a n_a(OH)_a = N_a - C_c H_a - N_a = C_a H_a(OH)_a$ [1:2:1]. From diagorised p-amido-benzene ago-resoroin and an alkaline solution of resorcin (Meldola, C. J. 47, 661). Dull brouze-like powder, v. sl. sol. alcohol, insol. toluene. Its solutions in oone. Il, SO, and in alkalis are violet.

Oxy - carboxy - beazene - azo - beazene - (a) - azo-

(8) naphthol

of (8)-naphthol (Meldola, C. J. 17, 668). Minute brown needles (from boiling audine). sol, boiling toluene, sl. sol, alcohol and glacial HOAQ Its solution in conc. II SO, is greenishldue, changing to violet on dilution. Its colution in KOHAq is reddish-violet.

Oxy-cymone-azo-tri-phenyl-methane-azothymol

(6.3.4.11C, II, MeDr(OH) - N, - C, H > CHPb, [6.3.11(C, II], MeDr(OH) N₀ - C, II > CHPb, [170°]. Formed by mixing diagotised diamidotri-phenyl-methane hydrochloride with an alkalina solution of thymol (Mazzara, G. 15, 41). Amorphous black powder. After reduction and oxidation it gives thymoquinous.

(a) . Oxy . naphthalone. p - azo - henzene. (a) .

naphthol

 $C_{10}H_{a}(OH) - N:N - C_{0}H_{4} - N:N - C_{10}H_{a}(OH)$ Formed by diazotising p. amido benzene azo-(a) naphthol and mixing the product with an

alkaline solution of (a) naphthol (Meidola, O.J. 47, 664). Green lustrous powder; v. al. sol. glacial HOAc, alcohol, and toluene, m. sol. boiling aniline, forming a red solution. Its solution in NaOHAq is blue, in conc. H.SO. blue, turned violet on dilution. With Ac.O and NaOAc it forms on heating a di-acetyl

(3)-Oxy-(a)-naphthalene-p-azo-benzene-(a)-azo-(b)-naphthol

C₁₆H₄(OH). N:N -C₆H₄-N:N -C₁₆H₄(OH). (β) (cover 275°). From diazotieed p-amido-benzene-azo-(β) naphthol and an alkalino solution of (β)paphthol (Meldola, C. J. 47, 664). Dull bronzy powder, or green needles (from hot anlline). Insol. boiling alcohol, or NaOHAq; sol. cold alcoholic NaOH, forming a violet solution. Sl. sol. hot toluene forming a magenta solution.
Conc. H.SO, forme a blue solution, turned violet on dilution.

(a) . Oxy-naphthalene. p-azo - cenzene.azo. (8)naphthol

 $C_{\rm in}\Pi_a(0|1) - N;N-C_aH_a-N;N-C_{\rm is}H_a(OH).$ [23:6']. From diazotised p-amido-benzene-azo-(a)-naphthol and an alkaline solution of (β) naplithol (Meldola, C. J. 47, 865). Dull bronzy powder, v. sl. sol. boiling alcohol; m. sol. boil-ing toluene and glacial acetic noid forming violet solutions; conc. H₂SO, forms a blue colution, becoming violet on dilution.

(a) ·Oxy·naphthalene-p·azo-henzene-azo-(β)· naphthol·di-zalphonio acid

C_nU_((01)-N:N-C_nU_-N:N-C_nU_((SO_nU_)(OH).

From diractised p-amido benzeno azo (a)-naphthol and an alkaline solution of (B)-naphthol di-sulphonic acid. Its sedium ealt is violet and gelatineus; it is nn indigo blue dye (Meldola, C. J. 47, 665).

(3) Oxy naphthalene azo benzone azo (3) naphthol disulphonie aeld. Is similar to the

last body, but of greater stability (M.).

Oxy.dl-sulpho-naphthalene.azo-henzene.azo-(8) naphthol di salphonic acid C₁₀H₄(SO₂H)₄(OH)=N₂-C₂H₄-N₂-C₁₀H₄(SO₂H)₄(OH). (Histening greenish needlos. Dyes wool and silk a deep indigo-blue, which, however, is very unstable to light. Is prepared by diazoticing the mono-acetyl derivative of p-phenylene-diamine and combining it with (8) naphthol-disulphonic acid (modification insoluble in spirit), the product Call, (NIIAc). N. C. I. II. (OH) (SO, H), which is a searlet colouring matter, is seponified, diazotised, and again combined with (\$) naphthol-di-sulphonic acid (Nietzki, B. 17, 344; 1350).

Phenyl amido henzene p.azo henzene azo.

(8) naphthol

• HO.C, oH, -N2 -C, H, -N2 -C, H, NHC, H2. [204°]. From p-amido-benzene azo di phenylannine by diazetising and adding sodium (8)-narhthol (Meldola, C. J. 43, 442). Warty ecales with bronze lustre. Sl. sol, boiling alcohol, v. sol, benzene. The solutione are red. In glacial acetic acide the colution is red when hot, violet when cold. Solution in conc. H, SO, is greenishblue, solution in alcoholic KOH is red but turned blue by HCl (characteristic).

Phonyl - ethyl - amido - henszne - aso-bensene-

azo.(8)-naphthol

HO.C., H. -N. -C.H. -N. -C.H., NELOH.

From p-diaso-nitro-bennene by combining with ethyl-di-phenyl-amine, reducing with ammo-nium sulphide, diasotising the product and treating with (β) - naphthol. Bronzy powder. Solutions in alcoholio KOH, alcohol, and benzene are red; in conc. H.SO, indigo-blue, turned hright blue on diluting. HCl turns the alcoholic solution blue (Meldola, C. J. 45, 111).

p - Sulpho - hanzene - azo - henzene - azo - (β)

naphthol.[\(\beta\)]. Fullphonic acid
(4) (1)(1) (4)(a) (8) (8)
C.H.(\(\sigma\)). N.N.-C.H.(\(\sigma\)). N.N.-C., H.(\(\sigma\)). (8)(1)(01).
From diazotissed p-amido-benzeme-azo-benzems sulphonic acid and an animoniacal solution of sodium (8)-nsphthol 'a'-sulphonio acid (Bayer a. Co., B. 15, 1351). The sodium salt is a scarlet dys (oroccius searlet). Conc. H.SO, forms a blue solution. The absorption spectrum has been studied by Hartloy (C. J. 51, 195).

p-Snlpho - benzene-azo-bsnzene - azo - tolylens

diamine

C.H. (SO.H) N - C.H. N. - C.H. (NH.). Reddish-brown microscopio needles. Formed by diazo-benzeno-azo-benzene-p-sulcombining phonio acid with tolyleuo-diamiuo (Griess, B. 16, 2036).

Sulpho - benzene-azo - di - phenyl-di-isoindoleazo-benzene zulphonio acid

Formed by the action of diazo-benzene-sulphonio acid on di-phenyl-di-isoindole. Metallic glistening brown scales. Very slightly soluble in all solvents. Dyes silk and wool nearly the sams shads as chrysoidino. On reduction it gives sulphanilic acid and di-amido-di-phenyl-di-isoindols.

Salta.—A"Na, aq: yellow plates.—A"Ag; red prisms (Möhlau, B. 15, 2495).

Sulpho - benzene-azo-sulpho-benzene-(a)-azo-

(8)-naphthol

O.H. (SO,H)-N2-C,H2 (SO,H)-N2-C10H4.OII. From diazo-sulpho-benzeno-azo-honzeno sulphonio acid and an alkaline solution of (β) -naphthol (Nietzki, B.13,800). The sodium salt is a rad dys (Biebrich scarlet). Conc. ILSO. formz a green solution. Its absorption spectrum has been studied by Hartloy (C. J. 51, 194).

Sulpho, bennene, ano sulpho, bennene. (8)aso-naphthyl-p-tolyl-amine ('Wool-black')
O,H.(SO,H),N.C.H.(SO,H),N.C.H.,NHC.H.,
The comi, product is a bronzy powder. V. sol.
hot water with a violet-blue colour. Dissolves in cone. H.SO, with a deep indigo-blus colour. Formed by combination of diazotised amidosulpho-benzane-azo-benzene-sulphonio acid with p-tolyl - (3) - naphthylamine. By holling with moderately dilute H₂SO₄ it is decomposed into tolu-(af)-naphthazine [169°] and amidoszo-benz-

ena-di-sulphonic acid: O.H.(SO,H)-Ns-C.H.(SO,H)-Ns-C.,H. NHC,H.= C,H,+C,H,(SO,H)-N,-C,H,(NH,)(SO,H). The Ca and Ba salts are insoluble black pps.

(Witt, B. 20, 579).

p-Toluene - aro-m - diamido - benaene - aro - (8) naphthalena O,H,-N,-C,H,(NH,),-N,-O,H, Formed by the combination of p-diago-toluena with (3)-nsplithaleno-azo-m-phonylouo-diamine (Griess, B. 16, 2031). Small red glistening plates. V. sol. chloroform.

p-Toluene-azo-ethyl-pyrrol-p-azo-tolusue C,H, -N2-C,H2NEt-N2-C,H,,

O,H,Mo.N,,O=CH NEt C,II,Mo.N,,O=Cl1 [1800]. probably

Formed by othylation of tolueno-azo-pyrrol-azotolneno; or by combining diazo p-tolnena chlorido (2 mol.) with ethyl-pyrrol (1 mol.) in alkalins solution. Steel-blue needles. Sl. zol. alcohol (O. Fischer a. Hepp, B. 19, 2254).

Tolnene-azo-dioxy-benzane-azo-tolueno C,H,-N2-C112(OH)2-N2-C,H2. By thonetion of p-diazotolneno ohlorido on an alkaline solution of p-toluene-azo-resorcin two isomerides are formed which are separated by their different solubilities in alkalis.

(a)-Compound [256°]. Yellow felted needlos. Sparingly soluble in alcohol and cold chloroform. (8) - Compound (203°). Brownish - black microscopic needles (Wallach, B. 15, 2825).

Toluene-ago-trioxyhanzana-azo-toluane C₆H₄Me-N₂-C₆H(OH)₅-N₂-O₆H₄Me. Long red needles. Prepared by the action of diazotoluene nitrate on phlorogluoiu (Wesolsky & Benedikt, B. 12, 227).

Toluene-azo-pyrrol-szo-tolusne C, H, - N2 - C, H2NH N2 - C, H, Call, Mo.N., C.-Cli probably NII ... C.11 Me.N., C.=CH

Formed by adding (2 mols.) diazo-p-toluene chloride to an alkaline solution of pyrrol (1 mol.). Red prisms with steel-blue reflex. alcohol. Its alcoholic solution is turned reddishviolet by conc. HCl. Dissolves in conc. H2SO4 with a blue colour (O. Fischer a. Hepp, B. 19,

Tolnene-o-azo-toluene-azo-(a)-nsphthol O₂H₄(CH₂) = N₂ = C₄H₂(CH₂) = N₂ = O₁₂H₄(OH), or C₄H₄(CH₂) | N = N = C₁₂H₄(CH). [210°]. N = N = C₁₄H₄(CH).

Formed by combination of o-diazo-azo-tolusne with (a)-naphthol. Brownish-red needles (from snilino). Sl. sel. ordinary solvents. Insol. aqueous NaOII, sol. decholio NaOII wan a violet-red colour. By SnCl, it is slowly reduced to amido-(a)-naphthol and tolyl-azimido-toluens C.H.; N.C.II., together with small quantities of p-toluidino and tolylone-o-diamine (Zincke a, Lawson, B. 20, 1178).

Toluene-o-azo-tolnene-azo-(8)-naphthol C₂H₄(CH₃)-N₂-O₄H₄(CH₃)-N₂-O₁₈H₈(OH), or $C_{\epsilon}\Pi_{s}(CH_{s}) \Big\langle \begin{matrix} \uparrow & \uparrow \\ N-N-C_{\epsilon}\Pi_{s}(CH_{s}) \end{matrix}$. [177°]. Formed

by combination of o-diazo-azo-toluene with (8) nsphthol. Long dark-red four-sided prisms, with green raffex. V. sol. benzane, and ohlorowith green raflex. form, sl. sol. alcohol, acetons, and petroleum-spirit. Dissolves in alcoholic NaOH only slightly.

By BaCI, it is reduced to (a) amide (8) asphibot and tolyl asimide toluene (, H, N, C, H, together with small quantities of p-tolnidine and tolylenee-diamine (Zincke a. Lawson, B. 20, 1179).

Telnene-p-azo-teluene azo-β-naphthel

C.H.(CH.) -N₂ -C.H.(CH.) -N₃ -C.₁₀H.(OH). [186°]. Formed by combination of p-diazo-azo-tolusne with (B)-naphthol. Deep red needles. V. sol. hot alcohol and benzano. By SnCl. it is easily reduced, giving (a) amido (b) naphthol, o toluidine, and tolylene - p - diamino C.H.Mc(NH2)2 [1:2:5] (Zincke a. Lawson, B. 20, 1182).

Tolnene-o-zze-tolnene-aze-(8)-naphthylamineC₄H₄(CH₂),N₂,C₄H₄(CH₃),N₂,C₄H₄(NH₂),or the corresponding hydrazimido-formula. [203°]. Formed by combination of o-diazo-tolucne-azoglistening plates. V. sol. bonzene and chloroform. By SuCl. it is reduced to tolyl-azimidotoluene C, H_a:N₃C, H₁ and (probably) naphthylene-o-diamine (Zincke a. Lawson, B. 20, 1180).

AZO- COMPOUNDS, TERTIARY.

Oxy-bensens-p-aze-benzsns-(a)-aze-naphtbalene.arc.phenel

Hos. II. N. - C.II. - N. C. II. - N. - C. II. - O.II. oH. From NH₂ C₄II, N. - C. II. - NI. MI₂ by diazotising and adding sodie phenate (Meldoln, C. J. 43, 489). Dull bronzy green powder. Porms orange solutions with beiling aniline or toluene. Solution in alcoholic KOH is violet; in conc. H.SO, indigo.blue.

Di-oxy-henzene.p-azo-henzene-(a)-azo-naphthalene-age-reseroin

(HO)₂C₆H₆-N₆-C₆H₆-N₅-C₆H₆-N₅-C₆H₅(OH)₆ Brown powder. Its solutions are dull red in alcohol, violet in alcoholic KOII, bluish-green ln cone. Il SO.

(β) exy-naphthalsne-p-azo-benzene-(α)-azonaphthalane-azo-(B)-naphthel

 $HO_{cl_{0}}H_{\bullet} = N_{\bullet} = C_{\bullet}H_{\bullet} = N_{\bullet} = C_{to}H_{\bullet} = N_{\bullet} = O_{to}H_{\bullet}, OH_{\bullet}$ Formed by diazotising amido-benzene-azo-(β)naphthylamine Nil. C. II, N. C. H. NH, and treating with an alkaline solution of (B) naphthol (Meldola, C. J. 43, 437). The pp. may be separated by crystallisation from hot aniline into two modifications, one remaining in solution, the other separating as needles with green

Crystalline form [c. 2950]. Insoluble in boiling alcohol, acetono, or glacial acetic acid, hardly soluble in chloroform or benzone. Forms violat solutions with aniline or nitrobenzene. Solution in cone. H.SO, ie deep inky blue. Alcoholie KOH forms, with difficulty, n blus

Soluble form .- Soluble in the above liquids. Its solution in boiling glacial acotio acid is violet when hot, blue when cold. Solution in cone. H2SO, is clear indigo-blue. Alcoholic potash forms a violet solution.

Dl-sulphonic soid.-From (8)-naphtholsulphonic acid and diazotised NH C.H. N. C.H. NH.

(a) Oxy-naphthalene-p-age - benzene \cdot (a) azemaphthalene-azo-(a)-naphthol. Similar to the preceding. Bronzy powder, forming a blue solu-tion in alcoholic KOH, and an indigo blue solution with cone. H,SO.

zyleze an-(s)-aspathal
HC.O.H. N. C.H. M. C.H.Me. N. C.H.OH
From NH.O.H.N., C.H.Me.NH., by diazotising
and adding sodium (s)-naphthol (Meldola, C. J.
43, 430). Small green reading Small green needles (from zylene). Its solution in aniline is reddish-violet; in xylens, violet; in hot alcoholic KOH, blnishviolst; in conc. H,SO, greenish-blue, becoming blue on dilution. It is insoluble in alcohol and in glacial acetic acid. Its sulphonic acid dyes silk and wool dull violet.

Tri-oxy-tri-naphtbalene-hexa-asc-tri-phenylcarbinol HO.C(C₆H₁-N₂-C₁₆H₄OH)₃*. From diazotised para-rosaniline and (α)- or (β)- naphthol in alkaline solution (Meldola, C. J. 47, 608). Orange amorphous powders. Similar compounds may be got from ordinary rosaniline. They dye silk and wool orange.

AZO- COTION DYES. The azo- dye-stuff obtained from benzidine and ite bemologues (Congo-red, benzopurpurine, ditapurpurine, benzaurius, ohrysamine, benzazurine, &c.) have this special characteristic of dycing cotton without a mordant. This property depsuds upon the constitution of the diphenyl molecule, for benzidins itself (and other diphenyl bases) readily combines with the cotton fibrs. This is casily shown by allowing cotton to soak for 24 hours in a cold solution of benzidine hydrochlorids, wringing, drying at the ordinary temperature, and washing thoroughly with hot and cold water; if the cotton thus mordanted with beuzidine is now passed through a dilute solntion of nitrous acid, and finally treated with a solution of (a)-naphthylumine-sulphonic acid it becomes dyed with Congo red (Möhlau, B. 19, 2014).

Tetrazo-diplienyl (diazotised benzidine) forms red dyc-stuffs by combination with (a). and (β) -mono-sulphonic acids of (β) -nuphthol or with the G di-sulphonic acid. The R disulphonic neid (Na sult insoluble in spirit) however oxhibits a very peculiar reaction. One mol. of tetrazo-diphenyl combines with ons mol. of the R di-sulphonate forming a dye-stuff which is red. If two mols. of the sulphonate are taken one med, remains in solution unused. If now the ppd. rsd colouring matter is heated with the mother liquor, the second mol. of di-sulphonate is taken up and a blus colouring matter is produced. A similar reaction has been observed with many other di-amido- bodies (Schnliz, B. 17, 461).

In general, the tetrazo- derivatives of diphenyl, ditolyl, &c. (obtained by diazotising benz. idine and its homologues) can combine with either one or two mols. of a phonol, amine, or their sulphonic and carboxylic acids. componn's with one mol., viz., R" N.OH tolsrably stable, and sparingly soluble; by boiling with water or with alcohol the uncombined diazo group is replaced by OH or by hydrogen. These half-conjugated diazo-compounds readily combine with a second mel. of phenol or amine, so that mixed azo-compounds can be thus obtained. The latter bodies are colouring matters of various shades, and dys cotton direct without a mordant (Lange, B. 18. 1697; Martine, B. 19, 1755).

The and composeds (Heating villor, Heating parties, curetinine, de.) obtained by combining wo mais, of a phenol, amine or respective sulphonic acid, or one mol. of one amine or phenol and one mol. of another, with the tetrasocompound darived from di-p-amido-di-phenylethyleneor its eulphonic acide, have the property of dyeing unmordanted cotton from a soap bath. The compound from (a) naphthol sulphonio scid dyes cotton a bluish violet, (8)-naphthol.R-disulphonio acid a bluc, (a)-naphthylamine-eul-phonio acid a red salicylic acid a yellow, &c. (Bender a. Schultz, B. 19, 3234). The di-amidederivative of fluorene also give colouring matters which dye cotton. To the class of cotton coloure also belongs the compound which has recontly been introduced under the name of 'Primuline.' Thie body dyes unmordanted cotton from an alkaline bath a greenish yellow. It is an amido-compound and may be diazotised upon the fibra. By treatment of the ootton thus prepared with solutions of amines or phenols, fast colours (red, orange, and brown) may be produced (Green, priv. com.).

TRIÁZOL. A name given to the hypo-HN ----N

thatical C.H.N. or HC

(Bladin, B. 19, 2598) (v. PHENYL-METHYL-TRIAZOL and Cyano-PHENYL METHYL-THIAZOL).

AZO - MECONIC - ACETIC ACID v. Dioxy earboxy · methyl-phthalide · Azo-dioxy · phthalide. acetic acid.

AZO-DI-METHYL-HYDROQUINONE v. Di.

oxy-benzene . 120-hydr quinone.
AZO-NAPHTHALENE v. Naphthalene . 120-

naphthalene. TETRAZONES. Compounde of the form R.N.N.N.N.R. obtained by oxidising unsymmetrical di-alkylated hydrazines (q. v.).

AZONIUM BASES. Compounds of the form NH, NRRT"(OII) (Fischer). The name has been also applied by Witt (B. 20, 1183) to compounds

NR(011) of the form X"

AZO-OPIANIC ACID ie amido-homipio anhydride, v. HEMIPIC ACID.

AZOPHENINE C34H29N3. [237°]. Ie formed by the action of a variety of azo. and nitrosotompounds (e.g. benzeue-azo-aniliue, toluene-120-tolnidine, phenyl-amido-benzeno-azo-benzme, chrysoidina, diphenyl-nitrosamine, nitrosoil-methyl-aniline, &c.) upon aniline (Kimich, B. 1, 1026). The best mathod of preparation conusts in warming an acetic acid solution of aniina with a nitroeo- compound (e.g. nitrose-dinethyl-aniline) filtering off the crystale of azoshanine, washing them with alcohol and resystallising from anilina or nitro benzeno. If he anilino ie replaced by ite homologues, the comologues of azophenine are obtained. orms unstable salts of violet colon-. It cannot acatylated. Conc. H.SO, at 100° yields a ulphonio acid cryetallising in violet needlas, thich form brown crystallina salts. By heatng with aniline hydrochloride it is converted sto induline (q. v.). By heating alone at \$60° is converted into aniline, a violet inter-

mediata body, and fluoriodise. The letter is a splendid blue crystalline colouring matter, the solutions of which have a beautiful red fluorescence and a characteristic absorption spectrum. reduction with SnCl, azophenine yields aniline and a sparingly soluble hydrochloride of an unstable base. The latter, when eet free by alkalis, rapidly absorbs oxygen from the air, and changes into a new colouring matter, the salts of which are indigo blue (Witt, B. 20, 1588; Witt a. Thomas, C. J. 43, 112,

AZO-PHENYL ACETIO ACID v. Exocarbory toluene Azo phenyl-acetic acid.

AZO-DIPHENYL BLUE v. INDULINE.

AZO-PHENYLENE, now called PHENARINE (q. v.).

AZO-PHTHALIC ACID benzene Azo-phthalie acid.

AZO PYROMELLITIC ACID C.N. (CO.H).
Tetra ethyl ether A'Et. [1840]; glistening red trimetrio tablee; easily coluble in alcohol, ether, and acetio acid, insoluble ln water; cublimable. Formed by reduction of di-nitro-mellitio other with zinc-dust and noetle acid. It forme an unstable colourless hydroohlorido (Nef, B. 18, 2805).

AZO-RESORCIN v. ILEGORCIN.

AZO-RESORUFIN v. RESORCIN.

AZO-TOLUENE v. Tolucne Azo-toluene.

AZO-TOLUIDINE v. Amido-toluene-AEOtoluidine.

AZOXIMS. Amidoxims, R.C(NOH).NH, are produced by the union of hydroxylamine with nitriles; these react with acid chlorides, R'.CO.Cl, or anhydrides, (R'CO), O, with formation of alkoyl derivatives, R.C(NO.CO.R').NH., which under euitable conditions (application of heat, or boiling, with water) split off water and change into szoxime $B.C < \stackrel{N}{\sim} C.R'$. The azoxims are very volatile in the vapour of other liquids or in air; many of those boiling above 200° are volatile even in vapour of other. They sublime at the ordinary temperature of the air (Tiemann, B. 18, 1000; 19, 1475).

Chloroformio other acting upon bonzamidoxim produces an azoxim which hae the character of a lactam, C,H, C NII >CO.

Phonyl-acet-antitaxim.

Ph CH2-C(NOH).NII2. does not produce azoxims so readily as beuz-amid oxim, Ph.C(NOH).NH, but phenyl acryl-amidoxim (oinnamidoxim), Ph.CH:CH.C(N(1)).NH. produces them with great ease.

m-Amido-benzenyl-azoxim-benzanyl

C.H.(NH2).C.N.O.C.H. [143°]. Formed by reduction of m-nitro-benzenyl-azoxlm-benzenyl with alcoholic ammonium sulphide. Orystal lisce from alcohol or sublimee in long needles. V. sol. alcohol, ether, benzene, and ohloroform ineol. ligroin and water. By nitrons acid it is diazotised.

Salts. -B'HCl: very eparingly soluble.

B',H Cl.PtCl.: sparingly soluble pp.

Benzoyl derivative C, H,ON (NHBs) [218°]; needles; sol. boiling alcohol, ether, a benzene, insol. water and ligroin (School, R. 18, 2478).

Bennenyl-asoxim-bennenyl C, H, N,O i.e. C.H.O.NO C.C.H., [108°]. (290°). Volatile with steam. Sublimes in long white needles. V.D. (H=I) 113 3. V. sol. alcohol, ether, and

benzens; v. al. sol. water. Formation .- 1. By heating the benzoyl derivative of benz-amidoxim - C,H,C(NH2):N.OBz. 2. By heating benz-amidoxim with bsuzoic acid,

or with benzotriohloride.

Reactions. -- It is an extremely stable sub-stance, being unattacked even by strong HNO, or H.SO.. Long boiling with tin and HCl reduces it to bonzonitrile (Tiemann a. Krüger, B. 17, 1694).

Benzenyl-azoxlm-henzanyl-o-carboxyllc aold $C_{\bullet}H_{\bullet}.C \stackrel{N.O}{<} C.C_{\bullet}H_{\bullet}.CO_{2}H.$ [151°]. Formed by melting together benz-amidoxim and phthalic anhydride. White glistoning needles. alcohol, ether, and benzene, v. sl. sol. water and ligroin.

Halts. - A'Ag: white crystalline A'Ba aq: plates or microscopic needles.— A'Cu: bluish green pp.—A'PbOH: white

granular pp.

Ethylether .- A'Et: heavy yellow oil. Amide. -C, II,ON, CONII,: [160°], mioroscopic ueedles (Schulz, B. 18, 2163).

Benzenyl-azoxim-benzenyl-m-carboxyllcacld $C_{e}H_{e}.C \underset{N}{\overset{N.O}{>}} C.C_{e}H_{e}.CO_{z}H. \ \ [218^{o}]. \ \ \text{Formed}$ by heating benzamidoxim m-carboxylic acid with BzCl (G. Müller, B. 19, 1497). orystalline powder; sol. alcohol, insol. water.

Benzsnyl-azoxim-bntenyl

 $C_uH_s, C {\gtrless \stackrel{N.O}{\sim}} C.C_uH_s, \ (265^\circ).$ Oily fluid. Volutilo with steam. Formed by the action of butyrio anhydrido upon benz-amidoxim, or by climinating II O from the butyryl derivative of the latter (Solulz, B. 18, 1085).

Benzanyl-imidoxim-oarbonyl

CallaNaOa i.e. Calla Calla Con and

 $\textbf{0}_{s}\textbf{H}_{s}.\textbf{C}\underset{N}{\overset{N.O}{\leqslant}}\textbf{C}(O11). \quad \textit{Benzenyl-azoxim-carbi-}$ nol. [198]. Carbonises at about 300°. Soluble in alcohol, ether, chloroform, benzene, and hot water, sparingly in cold water. The aqueous solution reacts strongly aoid to litmus. It de-composes carbonates. Not attacked by PCl, or by HCl. Formed by elimidation of alcohol from benzenyl amidoxim carbonic sther or, directly, by heating bonsonyl-amidoxim with chloroformic ether.

Salts.—A'Ag; white pp.—A',Cu: green pp.

Ethyl derivative C,ll,EtN,O, [36'];
soluble in alceliol, ether, &c., nearly insoluble in water; indifferent body (Falck, B. 18, 2468;

Benzenyl-azoxim-ethenyl C.H.N.O i.s. C.H., C≪NO C.CII. [41°]. (244°). Formed by boiling benzsnyl-amidoxim with acstic anhy drids (Tiemann a. Krüger, B. 17, 1696; 18, 1059; Schulz, B. 18, 1084). Flat prisms. Easily volatile with steam. Sublimes at the ordinary temperature in white needles. V. sol. alcohol, ather, and benzene, al. zol. water.

 C_sH_s , $C \leqslant \stackrel{N.0}{N} \geqslant C$, C_sH_s . (255°). Colourless off. Bennenyl-asonim-propenyl Volatile with steam. Formed by the action of propionio anhydride npon benz - amidoxim (Schulz, B. 18, 1085).

Benzenyl-azoxim-propenyl-w-carhoxylic acid $C_{u}H_{u}.C < \stackrel{N.O}{N} > C.CH_{u}.CH_{u}.CO_{u}H.$ [120°]. Formed by fusing benz-amidoxim with succinic anhy-dride. White trimetric plates or prisms. V. sol, alcohol, ether, hot water and benzene, incol. ligroin. It is not decomposed by warming with

Salts. - A'Ag: white cryetalline pp. -A',Ca 3',aq: long glistening soluble needles .-A'Ba aq : short prisms, or monoclinic cryetals. A',Cu : bluish-green granular powder.

A'PhOII: granular pp. Ethyl ether A'Et: (255°), yellowieli oil. Amide C₁₀H₄ON, CONII₂: [168°], elender needles (Schulz, B. 18, 2459).

m.Carboxy-benzenyl-azoxim-benzenyl

 $C_sH_s(CO_sH).C \stackrel{N.O}{<} C.C_sH_s$. [218°]. Formed by heating benzamidoxim-m-carboxylic acid with benzoyl chloride. White crystalline powder. Soluble in acetic acid, alcohol, and other, insoluble in water and benzene. The aqueous solution of the ammonium salt gives pps. with AgNO, and CuSO, (Müller, B. 19, 1497).

m-Carboxy-bonzenyl-asoxim-ethenyl

 $C_uH_s(CO_uH).C \stackrel{N.O}{<} C.CH_s$. [217°]. Formed by heating benzamidoxim - m - carboxylio acid with acctic unhydride. White crystalline powder. Sol. alcohol and ather, sl. sol. water and benzene. The aqueous solution of the ammonium salt gives pps. with AgNO., CuSO., Pb(OAc)., and ZuSO. (Müller, B. 19, 1496).

p-Carboxy.benzsnyl-azoxim-ethsnyl

 $C_{i_0}H_sN_sO_s \ \textit{i.e.} \ C_sH_s(CO_sH).C {\stackrel{N.O}{<}} C.CH_s.$ [218°]. Formed by boiling benzamidoxim-pcarboxylic acid with acetic anhydride. Cryetalline solid. Soluble in hot water and alcohol, more sparingly in other and chloreform. The diluto solution of the ammonium salt gives pps. with AgNO,, Pb(OAc), and CuSO, (Müller, B. 19, 1492).

m. Carboxy - benzanyl - azoxim - propenyl - etarboxylio acld

Calla(COull).Con NO C.CH..CII..COull. [218°]. Formed by heating henzamidoxim-m-carboxylic acid with succinio anhydride. Needles. Sol. hot water, v. sol. alcohol and other, sl. sol. chloroform, insol. benzeno. The aqueous solntion of the ammonium ealt gives sparingly soluble pps. with AgNO₂, CuSO₄, and Pb(OAo)₂ (Müller, B. 19, 1496).

p. Carboxy -hansanyl -azoxim - propanyl -s-carboxyllo acid

C.H. (CO.H.) C.N.O. C.CH., CH., CO.H. Formed heating benzamidoxim-p-carboxylic scid with an excess of succinic anhydride. Sol. alcohol, sl. sol. water, v. el. sol. ether, insol. benzene and ohloroform. Carbonisee at a high tsmperature without malting. The dilute aqueous

solution of the ammonium salt gives insoluble pps. with CuSO, and Pb(OAo),, pps. soluble in hot water with ZuSO, and AgNO, (Müller, B. 19, 1498).

Ethenyl-azoxim-benzenyl

CH,.C≪N.O>C.C.H,. [573]. Long white needles. Begins to sublime at 70°-80°. Easily volatile with steam. V. sol. alcohol, ether, and benzene, sl. col. hot water, insol. cold water and ligroin. Formed by heating ethenyl-amidoxim hydrochloride with benzoyl chloride (Nordmann, B. 17, 2754).

m-Nitro-benzsayl-azoxim-benzenyl

 $C_4H_4(NO_4).C \stackrel{N.O}{\leqslant} C.C_6H_5$. [160°]. Formed by the action of benzoyl chloride upon m-nitrobenz-amidoxim C, H, (NO.). C(NH.,): NOH (Schöpff, B. 18, 1067). White needles. Sublimable. Sol. alcohol, ether, and benzene, insol. water and ligrom.

m-Nitro-ver enyl-azoxim-ethenyl

 $C_sH_s(NO_s).C \stackrel{N.O}{\swarrow} C.C11_s$. [109°]. needlos. Sublimable. Formed by the action of soetic anhydride upon m-nitro-benz-amidoxim C.H.(NO.).Č(NH.):NOH (Schöpff, B. 18, 1066).

m.Oxy-bsnzsayl-azoxim-benzenyl

 $C_{\bullet}H_{\bullet}(OH).C \gtrless_{N}^{N,O} > C.C_{\bullet}H_{\downarrow}$. [163°]. Formed by dlazotising m-amido-benzenyl-azoxim-benzenyl and heating the aqueous solution. needles. Sublimmble. Sol. alcohol, ether, and benzeno, scarcely sol. water, insol. ligroin.

Ethyl ether C, H,ON, (OEt): [71°]; fine felted crystals; soluble in alcohol (Schöpff, B.

18, 2475).

Phenyl-allenyl-azoxim-benzenyl

 $C_{\bullet}H_{\bullet}.CH:CH.C {\gtrless \stackrel{N.O}{\sim}} C.C_{\bullet}H_{\bullet}. \ \ [102^{\circ}]. \ \ Cinna$ menyl-azoxim-benzenyl. Formed by elimination of H.O from the benzoyl derivative of phenylacryl-amidoxim by heating it above its meltingpoint or by boiling it with water (Wolff, B. 19, 1509). Very slender white needles. V. sol. alcohol, ether, chloroform, and benzene, v. sl. sol. oold water. Sparingly volatile with steam.

Phenyl-allenyl-azoxim-ethenyl $\mathbf{C_{e}H_{s}}$.CH:CH.C $\gtrless^{\mathrm{N.O}}_{\mathrm{N}}\geqslant$ C.CH_e. [78°]. Sublimable.

Colourless crystals. Formed by heating phenyl-acryl-amidoxim with acetic anhydride (Wolff, B. 19, 1509).

Phenyl - allenyl - azoxim - propenyl - w - carboxylio scid

 $c_{s}H_{s}$ CH:CH:c N_{N}^{O} >C.CH: $_{s}$ CH: $_{s}$ CO,H: [114]. Formed by heating phenyl-acryl-amidoxim with snocinio anhydride (Wolff, B. 19, 1511). Long white glistening prisms. Sol. alcol.ol, ether, benzene, and hot water, sl. sol. ligroin. -A'Ag: white powder.

Phenyl-ethenyl-azoxim-bsnzenyl

 $C_eH_e.CH_r.C < \stackrel{N.O}{\sim} C.C_eH_e.$ [82°]. Formed by boiling the benzoyl derivative of phenyl acet. amidoxim with water for a long time (Knudson, B. 18, 1070). White needles. Volatile with V. sol. alcohol, ether, and benzene, insol. water.

Phenyl-ethenyl-azoxim-ethenyl C.H., CH., CKN, OCH, (269°). Oil, Volatile with steam. Formed by boiling the acetyl derivative of phenyl-acet-amidoxim with water for a long time (Knudsen, B. 18, 1070).

Phenyl - ethenyl - azoxim - propenyl - w

carboxylio acid

 $C_*H_*CH_*C\leqslant NO$ С. $CH_*CH_*CO_*H$. Formed by fusing together phenyl-acet-amidoxim C.H. CH. C(NH): NOH and succinic ambydride. Prismatic plates. V. sol. alcohol and other, sl. sol. cold, water, m. sol. hot. It is a strong soid. Salts. -A'Ag: white pp. - Λ'_2 Cu: bluish green pp. (Knudsen, B. 18, 2483).

Phenyl-oxy-ethenyl-azoxim-etbsnyl

 C_sH_s .CH(OH).C $\stackrel{N.O}{<}_N$ C.CH_s. [65°]. Formed by heating the acetyl derivative of phenyl-oxynect-amidoxim C.H.,C11(O11).C(N11,):NOAo with water for some time. Transparent needles. Distils undecomposed, and is volatile with steam. V. sol. alcohol, ethor, and benzene, sl. sol, cold water.

Acetyl derivative

 $C_a 11_s$. CH(OAc). $C < \frac{N.O}{N} \geqslant C.C 11_s$. [52°]; fine white needles; volatile with steam; sol. aloohol, ether, and hot water, nearly insol. cold water (Gross, B. 18, 1076).

p-Tolnenyl-azoxim-banzanyl C, II, N,O i.e. C,11,Me.C<\(\frac{N.O}{N}\)>C.C,11, [103°]. Formed by heating the benzoyl derivative of p-toluamidoxim, 11,0 being split off (Schubart, B. 19, 1490). Long slender white needles. V. sol. ether, benzene, and chloroform, sl. sol. hot water, invol. cold.

AZOXINES .- Compounds whose molecular formulæ may he written X" \ NII > Y", such as

Phenazozine Call, NII Call, and Naphthazoxine C1elle NII C1elle

They may be formed:—(1) By heating aromatic imide sulphides with copper exide:

C_{1s}H_s<^{NH}_S>O_{1s}H_s + CnO =

C_{1s}H_s<^N_S>O_{1s}H_s + CnS;

(2) By heating o-amido-phenols with o-di-oxy-

compounds: Coll CoH + Oll > Coll =

 $C_{e}11_{e} < \frac{NH}{O} > C_{e}11_{e} + 211_{e}O$

(Bernthson, B. 20, 942).

AZOXY- COMPOUNDS, compounds whose molecular formula may be represented by

X-N-N-X. They are formed by treating nitro-compounds with alcoholic potash or solium amalgam. They may be reduced to hydrazo-compounds, X—NII—NII—X, and finally to two molecules of an amine, X—NE, The products of nitration of azoxy- compounds;

are often unsymmetrical, X-N-N-Y.

0.H.(CHO)(CO.H).N . N.C.H.(CHO)(CO.H).

ormed by reduction of nitro-p-aldehydo-benzoic with aqueous KCN. Colourless needles. V. sol. ether, alcohol, and chloroform, sl. sol. ligroin, nearly insol. water. Decomposes at about 280°. It gives the ordinary reactions of

Phenyl hydraside: small golden yellow prisms (Homolka a. Low, B. 19, 1090).

o.Amlde-benzene-o-azoxy-aniliae

Di-benzoyl derivative

(C.H.NIIBz), N.O. [195°]. Prepared by acting on bonzoyl-o-nitranilide dissolved in alcohol with zinc-dust and adding ammonia and platlnie chloride. Bright yellow mass, insol. water, sl. eol. boiling alsohol (Mixter, Am. 6, 26).

m-Amido-bsnzene-m-azoxy-aniline [c. 2720 Di-benzoyl derivative .- Prepared by dissolving m-nitro-benzanihde in boiling alcohol and adding alcoholic ammonia and powdered zine together with a trace of platinum. Very light powder with pale yellow colour. Insol. alcohol, ether, and benzone (Mixtor, Am. 5, 5).

p-Amide-benzene-p-azoxy-aniline (C_aH₄NH₄)N₄O. [182°-184°]. Prepared by the action of potassic cthylate on its discotyl de-Sol. alcohol, giving a red solution. rivative. Sl. sol. boiling water, from which it separates on cooling as a fibroue yellow mass (Mixtor, Am. 5, 4).

Di-acetyl dorivative (C. H.NHAc), N.O. [275°-278°]. Prepared by the action of powdered zinc and numonia on p-nitro-acetanilide in alcoholic solution. Hair-like particles with light golden-yellow colour. Sl. scl. boiling light golden-yellow colour, alcohol,

Di-benzoyl derivative (C.H., NHBz), N.O. [310°]. Prepared by the action of zine and ammonia on p-nitro-benzanilide. Light yollow colour. Insol. alcohol and water (Mixter, Am. 5, 284).

o-Amido-toluone-azexy-o-toluidine [1:2:4] C.11,Me(N11.).N.O.C.H.Me(NH.) [4:1:2] Asoxy o toluidine. [168°]. From nitro-o-toluidine in alcoholic solution by sodium amalgam (Limpricht, B. 18, 1405; Oracif, A. 229, 344). Long orange silky needles (from alcohol), or yellow needles (from water). V. sol. alcohol, sl. sol. water. Converted by cone. H.SO., by molecular change, into amido-toluene-azo-amido-crosol.

Saltes-B"H,SO, laq: needles.—B"2HOL, B"H,Cl,PtCl,.—B"2HBr.

p-Amide tolnene-azoxy-p-tolnidine [1:4:2] C_aH_aMe(NH₂).N₂O.C_aH_aMe(NH₂) [2:1:4]. [148']. Yellow needles. Sol. alcohol and hot water. Prepared by the action of sodium amalgam on an alcoholie solution of nitrop · toluidine. -- B"(HCl)2; sl. sol. B"H.Ol.PtCl. (Buckney, B. 11, 1451).

Bensoyl derivative

C.H.Me(NIBz).N.O.C.H.Mo(NHBz). From C.H.Mc(NHBz)(NO.) [1:4:2] by treatment with xine and ammonia (Mixter, Am. 5, 285). Light yellow eubstance, insol, water and alcohol.

Bearese-p-asory-aniline C.H.(NH.).N.O.C.H. [189°]. S. 4-27 at 21°. Formed, together with bensene-aso-aniline, by

p-Aldohyde bennets asony - p aldshyde ben the action or assumentation withhide on the alcoholic solution of benzene asoxy-nitro-be ene (G. Schmidt, A. 122, 174; Z. [2] 5, 419). Large pale-yellow tables (from dilute alcohol), Sl. sol. hot water, v. sol. alcohol and ether; Reduced by tin and HCl to aniline and pphenylene diamine.

Salts .- B'HCl: silvery lamine, eaponified

hy water .- B', H, PtCl.

Benzene-azoxy-benzeae C.H. N.O.C.H. Mol.

w. 198. [86°]. S. (alcohol) 17.5 at 16°. Formation.—1. From nitro-benzene by the action of alcoholic KOH (Zinin, J. pr. 86, 98; Rasenack, B. 5, 364; Schmidt a. Schultz, A. 207, 328; Wilsing, A. 215, 228), or sodium-amalgam (Alexejeff, J. 1864, 525; Moltchanoffsky, J. R. 1892, 350).—2. From aniline by oxidiaing with KMnO₄ (Glaser, Z. [2] 2, 808).—3. From benzene-azo-benzene by oxidising with CrO₄ (Petrieff, B. 6, 577).

Preparation.—1. Byreducing nitro-benzene in alcoholio solution by means of sodium-amalgam. The yield is 87 p.c. of the theoretical (Mottehanoffsky, J. R. 1882, 224; Bl. [2] 38, 551).—2. By boiling nitro-benzene with sodium methylate, prepared from methyl alcohol (250g.) and sodium (10 g.), the reaction being as follows: 4PhNO₂+3CH₂ONa = 2Ph₂N_O + 3HCO₂Na + 8H₂O (Klinger, B. 15, 865).

Properties.-Pale yellow trimetrio needles insol. water, sol. alsohol, and other. quantities may be volatilised with steam.

Reactions.—1. When mixed with neutral substances (e.g. NaCl) and distilled it gives anilino, azo-benzeno, and other producte.— 2. Ammonium sulphide has hardly any action upon it in the cold, but on warming it reduces it to hydrazo-benzeno. -8. SnCl, und HCl reduce it to aniline, vory little benzidine being formed hydrochloride at 230° gives violaniline (v. Dechend a. Wiehelhaus, B. 8,1614).—6. Diphenylamine hydrochloride heated with it gives triphonyl-violaniline (Girard a. Caventou, B. 12, 290).—7. Conc. HBr at 250° gives di bromo-aniline (Sendzink, Z. [2] 6, 266); HI gives benzidine.—8. PBr, gives yellow crystals of C₁₂H₁₁N₂Br, which are converted by aqueous AgNO, into benzene azo benzene (Werigo, Z. [2] 6, 387) .- 9. PCl, added to an ethereal solution gives benzene-azo-benzene (Werigo, A. 165, 202). 10. Sodium amalgam gives hydrazo-benzene .-11. Sulphurous acid forme benzidine sulphate .-12. Nitric acid forms three benzene-azoxy-nitrobenzenes (q. v.) and also a tri-nitro-derivative C₁,H₁(NO₂)N₂O, [152°] (G. Schmidt, Z. [2] 5, 421). This is converted by CrO, mixed with conc. HNO₂ at 200° into C₁,H₁(NO₂),N₂O, [102°], and C₁,H₁(NO₂),N₂O, [52°] (Petrief, B. 6, 558). Benzene-areay-benzene-arsulphenic acid

C_eE., N₂O.C_eH₄(SO₂H). [60°-70°]. Very deliquescent redlish-brown tables. Formed as a byproduct in the exidation of m-amide-benzene-snlphonic acid by KMnO₄.—KA' aq: long soluble tables (Limpricht, B. 18, 1420).

Benzene-asoxy-benzene-p-sulphonic acid C₄H₂N₂O₄O₄H₄(SO₂H), [below 100]. Red scales V. sol. water. Formed as a by-product in

ridd is about I p.a.—KA saq: small yellow cyntals (Limpricht, B. 18, 1420).

Benzene-azoxy-bromo-benzone sulphonic acid C.H., N.O.C.H.Br(SO.H). Formed as a byproduct of the exidation of bromo-amido-benzens-sulphonic acid CaH,Br(NH,)(SO,H) [4:3:1] with KMnO, -KA'2aq: small red six sided tables (Limpricht, B. 18, 1423). V. sol. water and alcohol.

Benzene-p-azoxy-nitro-hanzono

C.H., N.O.C.H., NO. [1:4], [153°]. Formed together with the following body by the action of HNO. (S.G. 1.45) on benzene azoxy-benzene (Zinin, A. 114, 218). Hair-liko yellow needles. Reduced by alcoholio ammonium sulphide to benzeue pazoxy-aniline.

Bonzsns-szoxy-nitro-beazens

C₄H₃N₂O.C₄H₄.NO₂. [49]. Needles or prisms; propared as above. Alcoholic ammonium sulphide forms a compound C1.11, N2O(?) [85°].

Bsazsac-azo-y-nitro-beazeno

C.H., N.O.C.H. (NO.) (1:2). [127°]. Formed by adding fuming 11NO, (25 30 c.c.) to a solution of benzene azo benzene (20 g.) ia glacial acetio acid (100 c.c.) at 75°. Red rhombic plates. Sol. alcohol, ether, and acetone. Alcoholio KOl1 gives au emerald green colouration; by long boiling or by treatment with sodium amalgam it is reduced to a compound Callia N.O (Janovsky a. Erb, B. 20, 361).

Bearoic o-azexy-benzoic acid

[2:1] $C_sH_s(CO_2H)-N_2O-C_sH_s(CO_2H)$. [1:2].

Mol. w. 286. [237'-212°].

Formation .- 1. By the action of KCN on onitro-benzaldeliyde (Homolka, B. 17, 1902).--2. From o-nitro-heuzoic acid hy treatment with oodium-nmalgam or alcoholio KOII (Grices, B. 7. 1611) .- 3. Together with o-nitro-tolueue by boiling o-mitro-benzyl alcohol with aqueous KOH (Jaffé, H. 2, 57).

Properties .- Small colourless prisms; m. col. hot alcohol, sl. sol. ether and boiling water. Reduced by sodium-amalgam to carboxy-benzene-azo-benzoic acid, and finally to hydrazo-

benzoic aoid.

Salt .- BaA" dag.

Benzoio m-azoxy-heazoic acid [8:1]C.H.(CO.II).N.O.C.H.(CO.II)[1:3]. Formed by boiling m-nitro-benzoic acid with alcoholic KOH (Griese, A. 131, 92). Minute needles or plates. Insol. water, sl. sol. alcohol and etner. Reduced by tin and HCl to di-amido-diphenic lusol. water, sl. sol. alcohol and ether:

scid. Dlazoxy-henzelc acid

Formed by reducing di-nitro benzoic acid dissolved in NaOliAq with codium amalgam (V. Meyer a. Michler, B. 6, 746; Michler, B. 7, 420; A. 175, 150). An amorphous black powder, insol, alcohol, ether, benzene, chloroform, aad glacial acetic acid. Reduced by tin and HCl to diamido beazoic acid. HNO, gives an amorphous nitro- derivative.

Salts. AgA: black pp. eol. NH,Aq.—
BaA's: black pp.—ZnA's: brownieh-black pp.
An isomeric acid, resembling the above, is formed from (1, 2, 4)-di-nitro-benzoic acid.

m Bronto-Seures (8:1) O.H.Br.N.O.C.H.Br [1:8]. [118]. From bromo-nitro-bensene and alcoholic KOE (Gabriel, B. 9, 1405). Bright yellow prisms; v. sl. sol. alcohol.

p-Bromo-bonzone-p-azoxy-bromo-benzene [4:1] C.H.Br.N.O.C.H.Br (1:4). [172°] (Hotmann a. Goyger, B. 5, 919); [175°] (Worigo, A. 165, 198). From p-bromo-nitro-benzene by treatment with alcoholic KOII or codium-amaigam. Yollow loaflets, v. sol. hot alcohol. Nitrio

acid forme a tri-nitro- derivative [174°]. Brome - bonzene - azoxy - hrome - henzene eul-

honic acid C.H.Br.N.O.C.H.Br(SO.H). Formed as a by-product of the oxidation of di-brome-amide-benzene-culphonic acid C.H.Br.(NH.)(SO.H) [1:3:6:4] by KMnO, -KA'2aq: very small yellow scales (Limpricht, B. 18, 1425).

m-Chiere-benzone-m-azoxy-chloro-bonzene [3:1] C.H.Cl.N.O.C.H.Cl [1:3]. [97°]. Formed by boiling m-chloro-nitro-benzene with alcoholic KOH (Laubenheimor a. Winther, B. 8, 1623). Yellowish-brown flat needles. V. el. sol. sloohol. Troated with funing 11,80, it is chiefly converted into m-chloro-benzone-azo-oliloro-phonol, only a vory small quantity of m-chloro-bonzeneazo chloro-benzene being formod (Schultz, B. 17, 464).

p. Chloro-henzeae. azo xy-chloro-bonzene [4:1] C.H.Cl.N.O.C.H.Cl [1:4]. [155°]. p-chiloro-nitro-benzeno hy treatment with alcoholic KOH (Heumann, B. 5, 910; cf. Willgorodt, B 15, 1002), sodium-amalgain (Alexejeff, Z. 1866, 269), or (in ethereal solution) with codlum (Hofmann a. Geyger, B. 5, 916). 1'alo yellow needles, el. sol. cold alcohol. Treated with furning 11.80, it is chiefly converted into p-ohlerabenzene-azo-chloro benzene only forming traces of a chlorinated benzeno-azo-phonol (Schultz, B. 17, 45I).

Di-chloro-beazene-azoxy-di-chloro-benzone [3:5:1] C.H.,Cl., N.O.C.,H.,Cl.,[1:3:5]. [172°]. From [3:5,1]. di. chloro. aitro-benzeae and alcoholie KIIS (Beilstein a. Knrbatow, A. 197, 84).

Di-ohloro-honzene-azoxy-di-chloro-bensone [2:5:1] CaHaCla.NaO.CallaCla[1:2:5]. [112°]. From p-di-chloro-nitro-benzene and alcoholic KOH (Lanbenheimer, B. 7, 1600; 8, 1623). Small bright yellow needles.

p.Chlero.henzene-azoxy-chloro-nitro.benzene [4:1] C.H.Cl.N.O.C.H.Cl(NO.) [1:4:?). [184°]. From p-chloro-benzene-p-azov-chloro-benzene and HNO, (Heumann, B. 6, 912; 18, 1185). Bright yellow floculent substance. V. al. solvining alcohol; reduced by alcoholio ammo-nium gulphide to p-chloro-benzene-azo-chloronitro benzene.

Chloro.tolueno.azoxy.ohloro.tolnene [6:3:1] C₄11₄MeCLN₂O.C₄H₄MeCl [1:6:8]. formed by the action of Na on an ethereal solution of chloro-nitro-toluene (Hofmann Geyger, B. 5, 919). Small needles.

m-Iodo-benzeno-m. azoxy-lodo-bonzene [8:1] C,H,I.N,O.C,H,I [1:3]. From m-iodo-nltro-benzenc and alcoholio KOH (Gabriel, B. 9, 1408). Flat yellow needles; el. sol. cold alcohol.

p. Iodo-hencone-p. azory iodo bensene [4:1] C.H.I.N.O.C.H.I [1:4]. (200"]. From a iodo-nitro-benzene and alcoholic KOH (G. Light rellow plates or scales. Sl. sol. hot slocked

M. methyl.amide-bensens.asoxy.di.methyl., aniline [4:1] C.H.NMe.N.O.C.H.NMe, [1:4]. From p.nitroso.di.methyl-aniline and alcoholic KOH (Schraube, B. 8, 619). Glittering brown erystals; sl. sol. water, m. sol. hot alcohol and The ealts are decomposed by water. -B"H,PtCl, aq.

m-Mitro-beazene.m-azexy-nltre-benzene

'3:1] C₄H₄(NO₂).N₂O.C₄H₄(NO₂). [1:3] [142°].

Preparation.—A solution of 2 or 3 pts. of n-di-nitrobenzano in about 15 pts. of methyl ilcohol is mixed with a solution of sodium asthylate prepared by dissolving 1 pt. of sodium in 20 pts, of methyl alcohol. A vigorous reaction cets in, which is completed by 48 hrs. cochobation; large yield. Long needles. V. sol. benzsns, m. sol. ether and CS, v. sl. sol. cold alcohol. By heating to about 140° with strong Il, SO4 it is converted into the isomeric di-nitrooxy-azo-benzene Calla(NO.), Ny. Calla(NO.) (OH) (Klinger a. Pitschke, B. 18, 2551).

p-Nitro-diphenyl. p-azexy-nitro-diphenyl C.H. (NO.). C.H., N.O. C.H., C., H., NO., [225°]. From p-di-nitro-diphenyl by acting on its alcoholic colution with sodium amalgam (Wald, B. 10, 137). Brick red crystalline powder; terms a red solution in cone. 1LSO, Insol. most solvents. Reduced by alcoholic ammonium sulphide to benzidine.

Nitro-oxy-beazene-azexy-di-nitro-phenel

Diethyl ether C.H.(NO.)(OEt).N.O.C.H.(NO.).OEt. [168°]. From HNO, and the diethyl other of p-oxy-benzene p-azo-phenel: the product is exhausted with water, and then treated with alcohol. On cooling, the alcohol deposits the body in long yellow needles grouped in stars. Sol. ether, CHCl, Call, and glacial acctic acid (Andreac, J. pr. 129, 337).

An isomeric body. [1870]. This forms the greater part of the product of the nitration, and is left undissolved whon the former body is extracted with alcohol. It is crystallised from acetie other, in which it is very soluble.

o-Oxy-benzeac-o-azexy-pheael Call (O11). N.O. Call OH.

Ethyl other {C_eH₄(OEt)}₁N₂O. Accryphenetol. [102°]. By reducing a cold (0°) solution of a nitro-phenetol (1 pt.) in alcohol (7 pts.) by adding sodium amalgan; on adding water a pp. is got; this is freed from azo-phenetol by washing with strong HCl as long as the latter ls coloured (Schmitt a. Möhlau, J. pr. 126, 201).

Properties. - Colourless triclinic plates. In soluble in waier, but melts in boiling water.
Slightly soluble in cold alcohol, insoluble in hot alcohol. Not volatile with steam.

Diphenyl-axexy-diphenyl C.H., O.H., N.O.C.H., C.H., [205°]. Small yellow plates. Insol. water and alcohol, sl. sol. cetic scid. Prepared by the action of alcoholic KOH on p-nitro di-phenyl (Zimmermann, B. 13.

Phenyl-glycellle-o-axexy-phenyl-glycellle acid ON₂(C₂H₁O.Cl₁CO₂H)₂. [187°],

Preparation.—o-Nitro-phenyl glycollic acid (18°6 g.), water (140 g.) and Na₂CO₃ (5 g.) are treated at 55° with sodium-amalgam (165 g. of 4 per cent. sodium), added in small portions. On cooling, crystals reparate. These are dissolved in water and decomposed by HCl.

The precipitated acids are recrystallised from alcohol. If the mixed acids now melt above 162°, they are etherified by alcohol and HCl. The ether of the azoxy- said is lsss soluble in alcohol than that of the azo- acid, it is crystallissd from alcohol and then saponified (A. Thato, J. pr. [2] 29, 152).

Properties.—Crystallisee, from aqusous or

dilute alcoholic solutions, with aq as short prisms or as scalenohedra; but if lsft in conthat with the mother liquor these change to rhombohedra, taking np jaq. Both these forms ars snlphur-yellow. At 130° they become whits and anhydrons. Dissolves in alkalis, HCl and glacial acetic acid. The solutions are yellow, Forms red solutions with HNO, and H.SO,. Insoluble in ether and in benzons.

Reactions .- 1. Lead acetate, a ysllowishwhits flocoulent pp .- 2. AgNO, and BaCl, no pp. in hot solutions, on cooling a crystallins pp.

Salts.—(NII.).A": obtained, as a yellow micro-crystalline pp., by passing NH, into a solution of the acid in absolute alcohol. Its aqusous solution gives yellowish whits pps. with BaCl, and Pb(OAc), yellow pps. with AgNO. and Fe_{...}Cl_s, and a green pp. with CuSO₄.—Ag₂A".
—Agl1A": more soluble than the neutral salt. BaA" 2aq.

Ethyl ether.-Et,A"; [114°]. White silky

m-Snlpho-benzsne-m-azexy-benzene sulphonic scid [3:1] C.H. (HSO.).N.O.C.H. (HSO.) [1:3]. [125°]. Prepared by the reduction of m-nitro-benzene-sulphonio acid with alcoholio m-nitro-benzene-sulphonio acid with atconolio KOH (Brunnemann, B. 11, 1048; A. 202, 240). Yellow needlos. V. sol. water and spirit. A"K, 4aq: nocales.—A"(NH₂), 23aq: rhombic pillars.—A"Ba aq: difficultly soluble prisms. A"Ca 33aq: difficultly soluble needles.—A"Pb aq. Chloride [138°]. Yellowish-red pillars. Amide [273°]: sl. sol. hot water.

Snipho-naphthalsns-azexy-naphthalsas aniphenic acid C, H (SO, II). N, O.C, oll SO, H. From (α)-nitro-naplithalene-(α)-sulphonic acid and

water; cone. II, SO, forms a violet solution.

Salts.—K.A" aq: trimetric tabular crystals. Na,A" 2aq: tables. — BaA'aq. — CaA' 2aq. —

PbA' 2aq.

Tersphthalle-azexy-tersphthalie acid [6:3:1] C.II.(CO.H).N.O.C.H.(CO.H). [1:6:3].
Yellowish plates. Sol. hot, sl. sol. oold, water,
v. sol. alcohol and ether. Decomposes between 250° and 280°. Obtained by oxidation of aldehydo-benzoic-azoxy-al lehydo-benzoic acid N.O(O.H₃(CHO)CO.H), with alkaline KMnO₄.

Nalts.—A''(NH₄)₄*: long yellowish prisms.

A''Ag₄: yellow pp. (Homolka a. Low, B. 19, 1091).

.Tolnene-axoxy-bromo-tolnene C4H4Mo.N.O.C4H4BrMs. [74°]. From p-toluene-p-azoxy-foluens and bromins. Bright yellow tables; v.sol. alcohol and sther (Molms, B. 3, 551). Telnene-agexy-nitro-toluene

C.H.Mc.N.O.C.H.(NO.)Me. [84°]. Formed by nitration of p-tolucne-p-nzoxy-tolusne (Potrieff, B. 6, 557). Yellow needles.

Telnene-o-azexy-telnene C.H.(OII.).N.O.C.H.(CH₂). [60°]. Formed by passing Cl₂O into an othereal solution of e-hydrazo-tolnens (Petricff, B. 6, 557).

Preparation .- 10 pts. of o-nitro-toluene are

added gradually to a solution of 5 pia. of sodium in 50 pts. by volume of methyl alcohol, the mixture being finally cohobated on the water-bath for 8 or 4 hours.

Properties.—Large yellow needles or plates; the crystals belong to the dimetric system, 4:0 - 8416:1.

Reactions .- By distillation with fron powder it yields o-azotoluene and a little toluidine. It also yields o azotoluene by heating with H.SO. (Klinger a. Pitschke, B. 18, 2553).

p. Tottens. p. axoxy. toluene C.H.Me.N.O.C.H.Me. [70°] (M.); [59°] (P.). From p-nitro-toluene by reducing it in alcoholic solution with sodium amalgam (Melms, B. 8, 551; Petrieff, Z. [2] 5, 264; [2] 6, 30; B. 6, 557), V. sol. alcohol and ether.

Bremine gives a bromo-derivative, [74°], and a di-bromo-derivative (139°).

Nitric acid gives a nitro-derivative (84°), a di-nitro-derivative [145°], and a tri-nitro-derivativo [201°].

AZOXYLENB v. Xylene-Azo-xylene.

AZOXY-NAPHTHALENE v. Naphthalene-AZOXY-naphthalene.

AZOXY-PHENOL Oxy-benzene-azoxyv. phenol.

AZOXY-DIPHENYL Diphenyl-Azonyø. diphenyl.

AZOXY-TOLUENE v. Tolueno-AZOXY-taluene. AZULENE or Azulin. Dlue colouring matter present in essential oils of chamomile, wormwood, and millefolium. Causes these and other oils to give an absorption spectrum, viz., three A. Fleissner, M. 3, 705; 4, 234, 788; B. 15, bands in red and orange (Hock, Ar. Ph. [3] 2136; 16, 1421; Nüking, B, 18, 1143). 21, 17).

arulatic acid C.H.V.C. Plotocies brown pp. formed together with examide ar oxamic said by passing syanogen into squeet ammonia. Si sol pure water, with viole fluorescence; acid or alkaline solutions fluoresc green. Boiling water clowly converts it int mycomelic soid O.H.N.O. Nitrio seid ork MnO oxidises it to azulmoxin C.H.N.O., an orange powder, insol. water, sol. conc. H.SO, the

solution having a deep-green fluorescence.

Hydrazulmin C.H.N. is formed by mixing dry cyanogen with dry NH. It forms black leaticts, converted by water at once into NH, and azulmic soid (Emmorling a. Jacobsen, B. 4, 927). By the epontaneous decomposition of an aqueous solution of HCN containing a little NH, a brown pp. is produced which, according to Gautier (A. Ch. [4] 17, 168), contains an exulmis seid of the formula O.H.N.O.

AZURIN G. H., N.O. [250-5°]. Small colour-less tables forming solutions which have a splendid blue fluorescence. Prepared by heating salicylia aldehyde with o-tolylene-diamine

(Ladenburg, B. 11, 596).

AZYLINES. Azo-compounds of the form R.N.C.H.,N:N.C.H.,NR, prepared by passing nitric oxide into alcoholic solutions of tertiary aromatic aminos; thus, di-methyl-amiline-azyl-ine is described as di-methyl-amido-benzeneazo-di-methyl-aniline, di-amyl-aniline-azyline is described as di-sayl amido benzene-azo-di-amvianiline; and di-ethyl-aniline-azyline as di-othylamido-benzene azo-di-ethyl-anilino (Lippmann

Acacia. tannin.

BACCARINE. An alkaloid in Baccharis cordifolia or 'Bio-Mio.' Needles, al sol. water, sol. alcohol, amyl alcohol, and ether. Its aqueous solution is neutral to litmus (Arata, Ph. [3] 10, 6).

BACTERIA.—The name given originally to a common roll-like form which is assumed in the course of growth by the minute plants to which Nageli (6) in 1857 applied the term Schizomycetes; hance the term 'Bacteria' is very frequently used to designate the whole of this group of organisms.

The Bacteria, Dacteriacem, or Schizomycetes ato a group of plants of extreme simplicity of structure and very minute in size. Like larger fungi, they are destitute of chlorophyll, and accordingly are unable to decompose carbonic acid in the presence of sunlight; as a consequence their nutrition resembles in some respects that of animals, since they are dependent on the complex chemical substances produced by other organisms. The variety of substances containing either C or N, or both, which they can althout and make contributory to their sus-

SABLAR. The fruit of several species of tenance is very great, whilst the chemical rots. The seeds and husks are rich in changes which they bring about in these subchanges which they being about in these substances are no less varied and remarkable. The exact nature of these changes and the relation of the Bauteria themselves to the substances upon which they feed form an enormous flaid of inquiry which has only recently been looked at by chanists, and that, as yot, very oursorily. The study of the forms presented by different kinds of Bacteria in the course of their growth is also as yet in an incomplete state, and whilst it is certain that tilere are kinds of Bacteria characterised each by its particular forms, its particular pasulum or chemical food, and by its particular chemical operations resulting in the formation of definite chemical products from the breaking up of the appropriate pabulum, we do not yet know in any large number of cases whether a particular form is constantly associated with particular chemical conditions and results, or whether it is possible under medified conditions for a given form to change its chemical and physiological activities. In a certain number of cases we do know that modified chemical and physical conditions will cause a given form in the course of its growth to acquire a very marked modification of form. Hence it is at present im-

me to discriminate with assurance different 'species' of Bacteria, although botanists use par-ticular names to designate those which; so far as sur information yet goes, are characterised by the constancy of a certain rauge of form, or in addition to this, by the conetancy of chemical and physiological activity. By 'species' the naturalist understands a group of organic forms the members of which may present very little or very great differences of form and even of activities as compared one with another, but of which it is true, either that they actually are connected with one another by natural processes of reproduction which have occurred within human experience and observation, or that there is good reason to suppose that they might be so connected within human experience. Forms which are separated from one another by an Interval the passage of which has not been witnessed and recorded by observers in the past, or defies experiment at the present day, are distinct species. We have not by experimental breeding produced a horse from an ass or an ass from a horse, or both from a third form, and we have no record of observations leading to the inference that such a passage has occurred within human experience, lience the horse and the ass are distinct species. On the other hand, we have traditional and experimental evidence of the production of the varieties of fancy pigeons from the Hock Pigeon, and conversely we know that from the most fantastic of fancy pigeons the Rock Pigeon can be produced in the course of a few generations: hence the Rock Pigeon and the Tumbler, Ponter, Fantail, Carrier, &c., are all variously modified members of one species.

It is necessary to addid to the question of species here because the progress of our know-ledgs of Bacteria in the immediate past has consisted in an important degree in the recognition of the fact that a great variety of microscopic forms may helong to one and the same species of Bacterium, and bocause we have to expect the most important advances in the future from the endeavours of bacteriologists experimentally to breed by change of conditions one kind of Bacterium from another, and even to create experimentally new kinds; and this in spite of the fact that it has been unjustillably assumed that the forms of Bacteria at present recognised are of the nature of species and immutable.

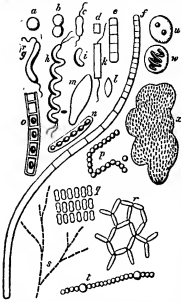
Classificatory position of Bacteria.—The issarest allies of the Bacteria among chlorophyll-bearing plants are the Oecillatorin and certain green-codured organising (the so-called B. chlorinum, B. virens, B. viride) which, whilst identical in form with some of the Bacteria, differ physiologically from them in possessing chlorophyll. The distinction between these plants and the Bacteria is not by any means a wide one, and there can be no doubt of the close genetic relationship of the green and the greenless Schizophyta, the Bacteria having, as is the case in other groups of plants, lost their chlorophyll and acquired parasitio or caprophytic (refuse-sating) habits pari passu.

eating) habits pari passu.

Ferms of Bacteria.—The Bacteria present themselvee either as swarming accumulations of detached cells or as linear aggregates (filaments or chains) of cells. Frequently the cells or plastids are loosely packed side by side and

ambedded to the party of the second ances on manning agreement and a second a second and a second a second and a second a usually extremely minute, being only 001 mm. er even less in diameter, though they may be larger. The cells consist of a homogeneous protoplasm in which no nuclens can be detected; cell wall, sometimes extremely delicate, bounds the surface of the cell, consisting of 'mycoprotoin, rarely of cellulose. When the cell-wall swells up, imbibing water, a jelly is formed in which the cells are eet at intervals ('zooglosa' condition). The ultimate chape of the celle of the Bacteria varies: it may be spherical (coccusform or micrococcus), biscuit-shaped or keyholeshaped, like two epheree partially fased (clithridium-form or bacterium sensu stricto), cuboid, bacillus form), rod-shaped (bacillus-form), carved like a bont rod (comma-form), twisted like a fraction of a corkserew (spirillum-form; if the spiral is not etrongly marked, vibrio-form). The most characteristic feature of these cells is their power of rapid growth and multi-plication by fission into two equal portions. Brefold has observed a Bacterium, formed by fission, grow to the size of the parent cell and itself divide into two in the course of half an hour, each of the daughter cells repeating ths process in half an hour. In the course of 24 houre there are thas produced from a single Bacterium more than a billion individuale like itself. The constant and rapid process of binary fission is what has led to the use of the names Schizophyta and Schizomycetes. All the forms of cells which we have enumerated as being assumed by Bacteria exhibit this phenomenon. But it is not necessary that the results of the fission should separate entirely from one another. Frequently such separation occurs, and in the forms known as clithridium (or bacterium sensu stricto) bacillus, vibrio, and spirillum, a filament of naked protoplasm is frequently observed hanging from each end of the fissionproduct, and by its lashing movements causes an active 'swarning' movement, or darting progression of the separate cells. Contrasted with this locometive swarming phase we have to note the phase of aggregation or continuous growth. As the result of variation in their pabulum, Bacteria which were at one hour separating from one another after fission-may remain in the next hour of growth in contactheld by their unraptured cell-walls. Thus are produced, in place of motile ewarming individual cells, aggregates or colonies which may be (1) linear; (2) tessellate; (3) branched; (4) reti-form; (5) massive. Any of these forms of aggregation may be exhibited by any of the different forms of cells. Linear aggregates of micrococci are called rosary-chains or streptococoue; linear aggregates of micro bacilli form longer bacilli and so-called leptethrix filaments; linear aggregates of comma shaped segments form spirilla; and small spirilla and vibrice when aggregated end to end form larger spirilla. Branched aggregation is seen in the so-called Cladothrix dichotoma, where a leptothrix filement breaks su as to allow a new line of growth to start at the broken surface, but without separation of the original continuation of the file which takes up a lateral position as a fair

branch [13]. Minitarly meah works (reseabling those of the green hydrodictyon) are produced (fig. r), and very regular teasellate aggregates (fig. q). In the latter, hacillus or olthridium forms may be arranged with absolute symmetry forming little plates of twenty or more cells, in rows of fivo or more (merismopedia form). In the sarcina form the grouping is cuboid, 'packets' being produced instead of



FORMS OF BACTERIA. 4. Microsceux; 5. Diplococcus; c. Ciltiridium or biscuit (with fagells); d. Microbacillis; c. Bacillus (built up of microbacillis; A. Microbacillis; c. Bacillus (built up of microbacillis; A. Leptothrix or classestous form, homogeneous at one eud, divided into havillis in the middle and into microbacilli at the other end; g. Spirillum (vibrio); h. Spirilum (close spini) with fagolis; t. Comma (exceeding the control of spirilum); h. Homogeneous bacillus with fagolis; f. Uvoid or double-come form; m. Large irregular form: these may occur of great size and various shapes, as flat diese (macrophasts) in Bact. Tubesceus.—Lead.; n. Bacillus with moniliform protoplasm, not spores; p. Rosary-chain; linear aggregate of micrococci; g. Merismopodia-form or tablet; teaselate aggregate of ciltiridis; r. Hydroidotyon-form: ratiform aggregate of bacilli; d. Gladothrix-form: faiso-branching linear aggregate of bacilli; d. Cladothrix-form: faiso-branching linear aggregate; larger cocci occur at intervals in a ohan of smaller occoi-observed in outivations of B. anthrecio; w. Leuconostoc-form: a spirilium with felly-like curelope; w. Leuconostoc-form: a spirilium with felly-like curelope; w. Z. Zoogica of cilthridis: olithridis embedded in felly-like matrix.

'plates.' Lastly, where the cell-wall swelle up and forme a jelly, we may have the bacterian cells of any one shape adhering by the jelly to one another (fig. x), and forming spherical or irregular masses of jelly (zooglos). These masses often are as large as the hand of a man, and are found on putrelying liquids and solids.

There is no doubt that all the forms of cell and of cell-aggregates which have been above

sessribed, and others to boot, may be exhibited by one and the same species of Bacterium. The Bacterium rubescens of Lankester [1] forms portwine coloured pellioles on decaying organic matter in fresh-water ponds and in salt-marshes, the protoplasm of the cells being coloured by a peculiar insoluble pigment bacterio-purpurin.

Lankester found all the varieties of aggregation and of cell-form (except spirilla, siuco observed by Warming [2] and by Giard [8]), in a small tank in which this organism was flourishing; their connection with one another was proved by their all containing the peculiar colouring matter and by transition-forms of growth. Tha accuracy of these observations has been conaccuracy of these concreations has been obtained by Zopf [4], and Lankester's conclusions adopted by him as well as by De Bary [5]. The species of Bacteria are said to be 'pleomorphie' or in Lankester's phrascology 'protean.' Nevertheless it is exceedingly probable that not all bacterian species exhibit so wide a range of form as dees B. rubeseens. Seme seem te be limited to the micrococcus and ellthridium cell-forms, and to exist either as free swarming cells of those shapes, or as linear aggregates of the eams. Others again are possibly limited to the microceccus ferm, though it is necessarily extremely difficult to be sure that under appropriate conditions of cultivation the cell-form and aggregation-form will not change altogether, and, until experiments have been very carefully made in each case with the object of breaking down the limitation of form usual to this or that species of Bacterium, it will not be justifiable to dogniatically characterise a species of Bacterium by reference to its shape.

Spore-fermation. The Bacteria reproduce with enormous rapidity by fission, but some faw are knewn to produce special reproductive bodies which have the property of registing the injurious effects of desiccation and heat.

In one sense of the word 'spore,' every eeg ment into which a previously unbreken plastid or cell of a Bacterium divides is a spore. A more epecial justification of the use of the term is found when occasionally one of the products of division is larger or more refringent than its fellows. Such 'spercs' are recognised in the cultivations of Bacterium (Bacillus) tuberculosis. None of these are sufficiently specialised as reproductive particles to justify thoroughly the use of the term 'spere' in regard to them. In certain species, however, e.g. Bacterium subtile, Bacterium authracis, and B. megaterium—the fermation of well-defined endospores is characteristic. The protoplasm within each member of a linear aggregate of bacillus forms separates centrally from itself an ovoid mass (fig. o), on the surface of which a coat of dense mysoprotein* is produced. The bacilli themselves die away and decompose, but the evoid spores remain, and have the power when dried of resisting an exposure to boiling water for as much as fifteen This property in the spores of B. minutes. subtile, which are common in old hay, has led to erroneous inferences as to the 'spontaneous generation, or 'abiogenesis,' of Bacteria. It is ossible, as soggested by De Bary [5], that the possible, as soggested by Do Dary Lop, which Bacteria which produce endospores are widely separate (as to their origin from groen significant and the separate of the separate from the other Bacteria which have no see

ised spores. He divides the Bacteria into Endospores and Arthrospores.

Tlassification and nomenolature of Escteris.

For the present De Bary's division of the Bacteria into Endosporea and Arthrosporea may be accepted. The various generic names in use, such as Streptococcus, Ascococous, Cladothrix, Begglatoa, Myconostoc, Leoconostoo, have no logical basis, and produce a good deal of confusion by a false appearance of order. It is probably sufficient at present to limit generic distinctions to the three terms Micrococcus, Bacterium, and Sporobacterium. The genus Micrococcus comprises those Bucteriaces which are not at present known to exhibit any form of plastid or cell other than that of minute spheres; the genus Bacterium contains only those Bacteriaceas which are known to exhibit in the course of growth rod-like forms of plastids, as well as in many cases micrococcus forms and spiral and straight filamentone forms; the genus Sporobacterium Includes only those forms which produce endo-spores, the so-called Bacterium (Bacillus) authracis, B. subtile, and B. megaterium. hering to this nomencluture, we still make use of the terms vibrio, spirillum, bacillus, clithridium, ascococous, zooglesa, &c., &c., to describe conditions of growth or varieties of cell form.

With regard to the use of specific names, it is well that every form or group of forms of Bacteriacem which definitely recurs in certain conditions, and scenis to be, so far as observation has gone, distinct from other known forms or groups of forms, should receive a name. Seeing that many of these names are probably but of temporary significance, it would be well that they should be as downtery descriptive of scroe feature of the supposed species as possible. The Bacteriaces should be named according to their chief properties, place of occurrence, or character

of growth, and not ofter persons.

The following is a list of some of the chief supposed species of Bacteriacen which have been described, with an indication of the mode of occurrence. It is by no means an exhaustive lest, and it is quite certain that some of the few supposed species here enumerated will, on further inquiry, be found to be phases of growth of other species.

Section A.: Endosporea.

Gonus SPOROBICTERIUM.

Species: S. subtile common in hay; S. anthracis, in the blood of cattle, sheep, and man, causes the disease known as splenic fever; S. megaterium, observed on boiled cabbage; S. butyricum, the butyric ferment, occurs in cheesemaking, and has been confused with S. subtile.

Section B. : Arthrosporca.

Genus Bacterium.

Species: B. termo, the commenest form in putrefying vegetable infusions, but not yet isolated and characterised; B. lincola, a larger form occurring in foul pends and sowage; B. rubescens, the protoplasm is wine red in colour, the plastids and aggregates are of the most varied forms, occurs in ponds on vegetable refuse; B. dichotoma, forming brnnehed aggregates (cladethrix) and straight and spiral tilaments, common in river water on dead leaves; B. Kuhniana, in wells and drain-pipes (Crenothrix); B. mesenteroides, forming masses like in the organic compounds forming either the

frog-spawn on the best-root jules of sugar re-fineries; B. suberculosis (fig. s.), in the diseased growths of men and animals suffering from taberoular consumption or phthisis; B. lepra, in the diseased skin of persons suffering from leprosy; B. mallei, in men and horses affected with glanders; B. typhosum, in the spleen and intestinal glande in fatal cases of typhoid fever; B. acidi lactici, in sour milk, tho manufacturer of lactic acid; B. cyanogenum, in milk, causing it to turn deep blue; B. pyocyaneum, in pus in badly dressed wounds, preducing an emerald-green colouring matter; B. alvei, causing a disease in bee larve knewn as foul-brood; B. urea, in urinals, causing the ammoniacal fermentation of urea; B. aceti, the vinegar ferincht, causing the conversion of ethylic alcohol into acetic soid, occurs in vinegar factories; B. prodigusum, causing blood-red staining of bread, milk, &c., leading to public alarm, and regarded as a portent; B. ovatum, causing the silk-worm disease known as 'pebrine;' B. cuniculicidum, eausing a specific septicemia in mice and blrds: B. chalerce gallinarum, in the blood and in the intestines of fawls suffering from chicken cholcra; B. pneumonie cronpose, in the exudation in croupous pneumonia of man; B. Kochii, Koch's comma-hacillus, found in the intestines of persons dead of Asiatic cholera (this is a spirillum form which breaks into comma-shaped segments; it is not proved to have any causal relation to cholera); B. Finkleri, similar to the last but larger, occurs in ordinary diarrhoa; B. buccule and B. Lewisii, spiral and filamentous forms breaking into commas which occur in the healthy human month.

Genus Michococcus.

Species: M. pyogenes, in acute obscesses; M. erysipelatosus, the cause of erysipelas in man; M. variole, to the pustules of small-pox; M. genorrheer, probably the cause of genorrhea; M. bombycis, enusing the disease in silk-worms known as flaccidezza; M. ventriculi, in the human stomach, observed in vomit, the 'sareina ventriculi' of Goodsir; M. scarlating, probably the cause of scarlet fever in man, and of a discase of the udder in cows; M. rabidorum, the cause of rabies, not satisfactorily isolated as yet; besides a list of twenty or thirty more causing special kinds of pyamia in such animals as rabbits and mice, or producing well-marked cofouring matters in colourless vegetable or animal infusions, green, blue, red, yellow, purple.

For a complete cuameration of the supposed species' of Bacteriaces which have been described, together with a description of each species and many illustrative figures, the reador is referred to the extremely useful and trustwithy treatise by Dr. Edgar Crookshank, entitled A Manual of Bacteriology, published by II. K. Lewis, London, 1887. Dr. Crooksbank gives complete references to the original description of overy known species and to the subsequent literature.

Chemical relations of the Bacteriacsm. - The above incomplete list gives some idea of the importance attaching to these minute organisms. It is an importance entirely depending on the variety and peculiarity of the chemical decompositions and reconstructions which they excite

tiving or dead bodies of higher plants and animals. Without Bacteris there would be no such thing as putrefaction, and therefore no circulation of the organic elements from their more stable compounds to the condition of albumens, fats, and sugars, and back again to the stable results of putrescence. The earth's surface would be cumbered with the dead bodies of former generations in which the carbon and nitrogen now serving as the feed of plants would be permanently locked up. All the evil smells which are not directly due to the chomist, are, with few exceptions, due to the action of Bucteria. Many valuable commercial products, such as acetic acid, lactic acid, and flavouring compounds such as butyric acid, are obtained through their agency. The pungent finnes of stable refuse ure caused by their action on area. It is almost certain that they too are the agents of nitrification in the soil - one species of Bacterium (or Microeocens?) converting the ammenia produced by another, into nitrates and nitrites. Some Bacteria produce highly poisonous bodies by their action on the albumens of dead animula and plants; amongst these poisous are the ptomaines, which have recently excited the attention of chemists [6]. Other Bacteria make their way into living animals and plants and there produce poisonous decomposition-products from the albuminous constituents of the organism, which are recognised in their effects under such names as splenic fever, scarlet fever, phthisis, rabies, &c. It appears that there are many kinds of Bacteria which are parasitic in and on the bodies of men and of other animals. the results of whose chemical activity is not injurious, whilst other kinds (or possibly the same kinds under changed conditions) produce deadly results. Other kinds again, it new seems cortain, are not merely innocuous but actually necessary to the healthy life of the animal they inhabit. The digestion of food in the alimentary canal of man and other animals is largely aided by the Bacteria which are present in the intestine in countless myriads, and it appears that the products of digestion owe their chemical characteristics in no small degree to the fineteria. In the absence of the normal parasitio Bacteria the products of digestion in the human intestine would, it appears highly probable, be of such a nature as to act poisonously when absorbed into the blood. When to these considerations we add the fact that the Bacteria are nbiquitous, abounding in the dust of the air. in all natural waters, and upon all surfaces whether of animate or inanimate objects which have not been chemically cleansed within a fow seconds of their examination, some idea may be formed of the immense importance which belongs to the study of the Bacteria in the immediate future.

Methods of Study.—At present the state of knowledge of the chemical relations of the Bacteria is extremely fragmentary. They were originally discovered by Leenwenhoek [7], the Dutch naturalist, in the fluids of the mouth, and various forms were subsequently seen with the microscope in natural waters, ponds, &o., and described by Ehrenberg [8] and others. It was Theodore Schwann [9], however, who, in 1838, demonstrated by a simple experiment that *Yor. It

the Basteria cause the putrefaction of organic substances, and that without them there is no putrefaction. Later, Pasteur [10], in opposi-tion to Liebig, extended Schwann's observations and conclasions, and established the doctrine of organised ferments, which has proved of immenss practical importance, and is as yet only at the commencement of its history. The foundation of the experimental demonstrations of Schwann and of Pasteur lies in the fact that the living protoplasm of the Bacteria is destroyed that is to say, undergoes an irrevocable chemical change -- when subjected to a temperature below or about that of boiling water. Consequently it is possible, by the action of heat, to destroy the Bacteria present in an experimental vessel and its contents, and to protect the contents from the further accession of Buetaria. By this method, and by this method alone, it has been possible to prepare organic infusious, as well as solid golatine, allumen, &c., which, whilst capable of supporting the life of Bacteria, are yet free from their presence for the time being. Such substances are said to be 'sterilised, They can be inoculated at pleasure with Bacteria and the effects of the inoculation studied. In order to procure the Bacteria for inoculation in a state of parity, special methods have been devised. So abundant and varied are the kinds of Bacteria present in nearly all matural organic material, that any rough process of inoculation will introduce many kinds of Bacteria simultaneously into a sterilised medium. To separate the various kinds of Bacteria for the purpose of study of each in its isolated condition, three principal methods are employed. The first applies, as A. Maknown, to but one kind, the Sperobacterium (Bacillus) subtile. The dry spores of this Banterium resist the destructifteer minutes, whilst all other known Bacteria are destroyed by it. flence we have only to boil old hav in water for a few minutes in order to obtain a pure cultivation of B. subtile. The second method (due to Nögeli [1f]) is that of fractional dilution. Given a liquid swamping with a mixture of various Bacteria, of which it is estimated by inspection that one individual in twenty is of the kind it is desired to cultivate. Dilute the liquid to such an extent that one drop of it should contain but a single bacterium. Then it is probable that every twentieth drop. will contain a single isolated individual of the desired Bacterium. Fifty tubes of sterilised nutrient material are prepared, and it to each a single drop of the culated Bacterium-holding tluid is introduced. One, or possibly more, of the tubes will thus be inoculated with an isolated example of the de ired Bucterium, which will multiply in the sterilised nutrient material and thus yield a pure cultivation, and can be recognised by the microscope. The third method is due to Brefeld, of Berlin. By streaking with a needle point a minute drop of fluid containing various Bacteria, over a surface of solid sterilised gelatine, the various Bacteria will be locally. isolated along the course of the streak. They will remain thus separated from one another and commence to multiply in situ. With a low power of the microscope and a fine needle samples can be now removed from the various

further cultivation and study. Similar isolation is effected by mixing liquid gelatine with a dilute infected liquid; when the gelatine solidifies, the various bacteria are embedded apart from one another, and grow in isolated patches,

which can then be removed and scparately studied by further cultivation. Conditions of life required by Bacteria.

General results. (a),

1. The first general result of these methode of study has been to determine the ubiquity of a large number of different kinds of Bacteria, and the comparative rarity of others. More will be said below as to the study of the distribution of Bacteria in air and water.

2. The Bacteria are found to differ from one another in their relation to free oxygen; the aërobic (Pasteur) will only multiply in the presence of free oxygen; the anaerobie will not flourish oxcept in the absence of free oxygen, or at any rate are indifferent to its presence. Thus B. anthracis is eminently acrobic, whilst the Bacterium of malignant ædema is anaërobic. The hay-bacillus (B. subtile) is aërobic, the butyrie bacillus of cheese (very similar to the former in appearance) is anaërobic.

8. The source of nitrogen required by Bacteria for building up their protoplasm ie various. Very many can take it in as low a form of combination as ammonia. Others require it in higher combination, and some either require it in the form of albumen or at any rate can take it from albumens. It is from albumens that some of the most remarkable products fermed by Bacteria result. The be little doubt that the first steps in this process are comparable to the digestion of albumen by animal cells. It is not ascertained that all and any Bacteria can attack albumens. The exact range of the chemical quality of the nitrogenous food possible to each species of Bacterium has yet to be determined.

4. The carbon required by Bacteria may be taken in as low a form as nectic acid by certain species; others can take it from tartarie acid; others can do with nothing lower than a angar; others again require glycerin or a similar body, and others apparently require their carbou us well as their nitregen to be presented in the form of a proteid. Thus it results that many Bucteria can be nourished by solutions of ammonium tartrate alone, whilst the limits of complexity of necessary food-compounds has various ranges in other species, all of which require accurate determination by the chemist. Little has as yet been ascertained in this direction, but recently Dr. Roux [12] of the l'asteur Institute, has made an oxtremely important observation showing the necessity for extended research of the kind. It was found by Koch extremely difficult to oultivate the Bacterinm tuberculosis, even npon blood-serum kept at the normal temperature of the body. Renz found that the addition of a minnte quantity of glycerin to the serum led to the rapid and abundant growth of the B. tuberculosis supplied with that mixture; and further, that an ordinary

salebas of growth and placed in the pure con- minute quantity of glyperia acts as a moss dition in tubes of sterilised nutriont material for effectent nutrient medium for this species. It is highly probable that other such special requirements in regard to the chemical nature of their food, exist in respect of other species of Bacteria, whilst others again are more catholic in their nutrition.

> 5. Water is necessary for the growth of Bacteria as of all living things. Most Bacteria will flourish in the presence of that small amount of water in proportion to solid matter which suffices to constitute mere dampness or moietness. Bacteria are not killed by partial desiccation, but none resist thorough desiccation. In this respect important variations have been determined in different kinds. The spores of the Endosporea have a special power of resisting desiccation.

> 6. There is an optimum temperature favouring the growth of Bacteria, which ranges in various species from 10° to blood heat. Exporiments have been made priving that certain species of Bacteria are killed by extreme cold, whilst all are arrested in growth during exposure to the freezing temperature of water. The most careful observatione have been made in regard to the effects of exposure to high temperature. Exposure to a temperature of 100° for five minutes kills all Bacteria except those belonging to the Endosporea, the spores of which can resist the effects of this exposure for half-an-hour, and possibly longer. Many Bacteria are killed at lower temperatures (e.g. 80°), but careful experiments are wanting.

7. Experiments as to the effects of diminution and increase of atmospheric pressure upon the life of Bacteria have been made, but without reference to particular species. Diminution of pressure is not known to have any influence, whilst experiments made by the writer show that a pressure of thirty atmospheres does not hinder the development of putrefactive Bacteria appreciably, though modifying the chemical results of their life-processes. Extremely high pressures are stated to be destructive of Bucteria.

8. The influence of light is, according to the experiments of Downes [13], inhibitory to the growth of certain Bacteria, but the species so nffected have not been determined. This is in accordance with the absence of protective pigment in most species, and the general fact of their growth within turbid liquids and beneath the surface of solid bodies awny from the light.

9. Like the yeast-plant, which creates a poison (alcohol) in the nutrieut fluids in which it grows, which after reaching a certain percentage causes the arrest of growth and the subsidence of the yeast-cells—so the Bacteria are limited in their growth by the existence of products of their own formation. These products have not been investigated by chemists. But it appears to be established that putrefactive Bacteria growing in a nutricut medium flourish for a time abundantly, then suddenly cease their growth and sink to the bottom of the vessel in which they have been growing, although the nourishing material is not exhausted. further and exact investigation of this phenomeat broth which alone cannot serve as pabulum menon by the chemist in regard to various for the B. tuberculosis, when mixed with a species of Bacteria must lead to results of the

greatest value in relation to the pr preventive incentation for disease.

10. A condition of the life of a given species of Bacterium is found in the presence of other species of Bacteria. Frequently one species of Bacterinm is the indispensable friend and associate of a second - preparing by its chemical activity the pabulum on which alone the second can thrive. An association of the kind is seen in what is called the vinegar plant, where Myooderma prepares from starch the alcohol which the Bacterium aceti converts into acetio scid. So, teo, the Bacterium of animoniacal fermentation is the antecedent of the Bacterium which converts ammonia into nitrites and nitrates. Equally important is the inhibition and possibly the destruction of one species of Bacterium by another. Very little has been ascertained on this important matter, but it appears that the presence of certain putrefactive Bacteria in a nourishing medium will actually prevent the development and growth of certain pathogenic species of Bactsria, although these are present in small numbers. Apart from the question of possible specific incompatibility of two Bacteria, it appears that the question of quantity (r. Cheyno [14]) is important. A species of Bucterium which is at the commencement of an inoculation experiment one hundred times more numerous than a second species, may by its rapid development and numbers prevent altogother the growth of the second species

11. The question of the conditions of life of the Bacteria involves the very important one of their tolerance of the presence of various chemical substances in the liquids in which they grow, those substances the presence of which is not telerated by the Bacterium being called 'germicides' or 'antiseptics.' On account of the practical importance of destroying or inhibiting the development of putrefactive and pathogenic Bacteria, a good deal of attention has been given to this subject by chemists, but unfortunately it is only recently in the laboratory of Koch [15] that experiments to determine the germioidal action of chemical substances have been made with the necessary discrimination of the opecies of Bacteria which were the subject of experiment. The fact is now definitely established that some species of Bacteria are killed by chemical substances which do not injuriously affect others, and that the amount of such substances which is effective varies in the case of different species. The inquiry has only as yet been com-menced, but it is of immense practical importance, since it may be possible to discover 'ger-micides' of a generally innocuous character which are specific poisons for esrtain diseaseproducing Bacteria, whilst harmless to other Bacteria and harmless to the higher animals in whose tissues the pathogenic Bacteria flourish. Thus weak solutions of quinino sulphate are poisonous to the Bacterium urea, whilst not ininrious to putrefactive Bacteria. Such a solution can be injected into the lawnan bladder without causing irritation, and thus the inflammation resulting from the ammoniacal decommation resulting from the annual position of the urine in the bladder by Backerinm ures, which sometimes gains access thareto, can be entirely arrested. In this inquiry the difference between actual destruc- but the organism gets into ite food instead

tion of the life of the Busteria, and there are or inhibition of growth due to the presence of the antiseptic chemical, have to be distinguished. It is also needful to inquire how far such 'antiseptios,' without killing or inhibiting Baoterla, may modify the physiological processes and chemical results brought about by the latter. The most powerful and generally effective poison for Bacteria appears to be corrosive anblimate. The presence of as little as 1 in 10,000 of this salt in a nutrient fluid has been found to kill Hacteria present. Phenol is also a general and powerful germicide. Boracic acid also and common salt in large quantities are effective. The nature of their action and their effectiveness in regard to different species of Bacteria have yet to be accurately determined. Antiseptic surgery, the future treatment of zymotic disease, and the preservation of perishable articles of food, donand upon the further discoveries of chomists in regard to this matter. It is not improbable that the most effectivo and useful germieides will be found in chemical substances which, like quinine, resemble those inhibitory products which are produced by the Bacteria themselves and act as the natural obstacles to their excessive multiplication. The more general question of the tolerance of or necessity for the presence on the one hand of free acid, on the other of free alkali in the nutrient fluids suited to different Bacteria, belongs here. It has been studied in regard to many Bacteria in a rough and ready way. Some Bueteria will not flourish in acid media, others will; but accurate quantitative investigations are still wanting.

The products of the activity of Baoteria When a species of Education graws in a nutrient fluid of known chemical composition with access to a definite and limited volume of atmospheric oxygen-under given conditions of temperature, pressure, and illumination-certain ohemical interchanges occur in the materials contained in the apparatus. These can be accurately determined in certain instances, and the variation of the quantity of change in relation to time oan be stated. Various factors of the process, auch as temperature, presence or absence of initial chemical substances, &c., can be varied, and the results stated and compared. In no case has such an experiment as yet been accurately made by a chemist. Nevertheless, we know roughly that, in the supposed experimental apparatus above indicated, there will be after a certain time an increase in the weight of mycoprotein and albu-mens existing in the form of Basteria, and corresponding diminution in the C, II, N, and O of the other material in the apparatus. Not only this, bat we find certain new chemical compounds present outside the actual substance of the multiplied Bacteria which result from and secompany the growth and life of the particular species experimented upon. The same general statement is true of any higher organism in relation to its necessary pabulum; but where in large multicellular organisms the resulting: products of this life of the organism are term porarily or permanently held within the man of the body, in the minute unicellular Bacteria there is no taking in or onvelopment of the materials to be acted upon by the living thing

the food getting into it : consequently processes comparable to the digestive and even to the more deep-seated metabolic processes of higher organisms take place in the nutrient liquid in which the Bacterium lives, being initiated at the surface of the swarming cells constituting the colonies of these miaute plants, and serving their economy equally as well as though they occurred In an alimentary canal or in a series of bloodvessels and tissue-spaces. The chemical changes induced by Bacteria should be studied from the same point of view as that taken by the physiclogist in regard to the activities of the various calls of the tissues with their diverse and specific functions. We are not yet in a position to treat the subject from this standpoint but we can distinguish with more or less certainty results traceable to respiration, digestion, assimilation, secretion and exerction; the cheminal correlatives of these processes are changes described as de oxidation, oxidation, specific fermentations, specific syntheses.

The obvious results of the activity of Bacteria (setting aside the probably universal evolution of CO₂ and coasumption of free O, common to the Bacteria and all living protoplasm) though by no means necessarily the most important in regard to their own physiology, are the production in the liquids in which they grow of (1) substances having distinctive smells and flavours; (2) substances having lirilliant coloms; (3) substances having enimently poisonous properties; to these may be subled such remarkable results of exidation as the manufacture of intutes in soil, if act tie neid in vinegar factories, and the manifestation of light—the phosphorescence—of decaying—m. nones, and other organio refuse.

The obenical nature of the substances which are thus produced, the by-products which accompany them, and the nature of the processes by which they are originated, have not yet formed the subject of chemical investigation to any large extent. Such knowledge as we have is due to Pasteur [16], to Fitz [17], and one or two others.

It seems probable that we may distinguish amongst these results those which are due to synthesis, by the Bacterium acting on lower compounds taken into its substance, and those which are due to analysis resulting from the action of ferments and other agents secreted by the Bacteria and acting on surrounding material of a high degree of chemical complexity. Of the nature of these ferments we know nothing; their existence is hypothetical but highly probable. To the first entegory belong certainly many of the brilliant pigments which the Bacteria produce; in most cases these pigments are soluble and pass out from the protoplasm into the surrounding water. In Bacterium rubescens the wine-red pigment is not soluble, and remains where it is manufactured in the cells of tho plant. The remarkable smelling substances formed by putrescent Bacteria also belong to The remarkable smelling substances this group of built-up products, and it is probable that the poisonous products of some pathogenous Baoteria, though not of all, are thus elaborated. The chief experimental reason which we have for concluding that these bodies are built up by the Bacterium from lower com-

pounds is this, that they are formed when the Bacterium is cultivated in a pure solution of ammonium tartrate (with traces of mineral salte) often called Pasteur's or Cohn's solution. Thus the Bacterium of hlue milk can be grown and made to produce its blue colour from ammonium tartrate, the Bacterium of green pus similarly, and many of the chromogenic Micrococci, whilst some of the specially active putrefactive Bacteria manufacture foul-smelling products from the same salt when experimentally nourished with it.

In regard to the second group, that of suhstances resulting from a breaking down of higher chemical bodies brought into relation with the Bacterium (and that probably by the action of a secreted ferment which may be minute in amount and possibly nover separated from the surface of the Bacterium-cell), we have to note first of all that the ferment itself belongs to the previous group. Secondly, that various species of Bacteria have been shown in produce cthylic and other alcohols in this way -from sugar and similar bodies-as iloes the yeast-plant (Succharomyces). Fitz [17] has shown that a certain Bacterium converts glycerin into ethyl alcohol, whilst another converts it iato butyl alcohol. Other Bacteria have been shown to convert sugar iato gum or into manuite, producing the so-called 'ropy fermentation syrups, winc, and beer. Urea is converted into carbonato of ammonia, hippuric acid into benzeie neid and glycocoll. Albumens are broken down into bodies which have not been determined in many cases, but include the ptomaines, nearidine, and trimethylvinyl-animonium hydrate. Various Bacteria as well as the specific B. lactici, produce small quantities of lactic acid from various substances, such as grape-sugar, milk sugar, and glyceria, whilst possessing other ferment-producing action also. Butyric acid is frequently produced in these processes by other Bacteria as well as by the B. butyricum of cheese-factories. Exact knowledge is, however, sadly deficient in these matters, owing to the fact that hitherto chemists have not been careful to ascertain what succies of Bacterium is present in the fermentations studied by them. Owing to this we do not yet know whether in different nourishing fluids and under different conditions of access of oxygen and of temperature, the same Bacterium can produce different fermentations. Such knowledge as we have tends to a positive answer to the above question. One of the best researches with a known species of Bacterium is that of Vandevelde [18], on the hay bacillus (B. subtile).

Since it is probable that there is this change of obenical activity under changed conditions, it is also probable that a Bacterium which is harmless under ordinary conditions of growth may, when specially cultivated in albuaninous media, acquire the property of living in the animal body as a parasite, and there cause deadly disease by its fermentativo action, or by the secretion of poisonous products. Buchure [19], starting from this theoretical consideration, has endeavoured to produce the deadly B. anthracts of splenic fover from the hay bacillus (B. subbille), and conversely to restore the parasitic form by cultivation to the primitive state. His experi-

ments, though of extreme interest, are not comelusive.

It is difficult to hazard a guess as to whether the poisonous effects of any given Bacterium proved to be concerned in the production of disease, are due to the secretion of a poison by the Bacterium or to the production of one by its fermentative action upon the constituents of the blood, tissoes, or intestinal contents. The chomical theory of the antidotal action of inoculation with various 'vaccine' cultivations, which is that favoured by their chief discoverer and investigator, M. Pastcur, would seem to involve the hypothesis that whilst the effective poison ie a secretion of the Bacterium, the antidotal material is a chemical compound resulting from the fermentative action of the Bacterium, quite distinct from the poison. This fermentationproduct hy its accumulation inhibits the development of the Bacterium as alcohol inhibits the further growth of the yeast plant by the fermentative action of which it has been formed. The phagocyte theory of Metschnikow [20] in relation to preventive inoculation does not involve thie distinction (see below).

The exidising action of Bacteria must be considered merely us a special form of their fermentative action. By the latter they produce intermediate chemical substances which are readily oxidised by the free atmospheric oxygen. It is probably thus that acetification proceeds when B. accti converts ethylic alcohol into vinegar or when organic nitrogenous bodies and ammonia in the soil are converted into nitrites and nitrates. There is no evidence of a specific oxidising action on the part of the Bacterium. The phosphorescent sobstance produced in stule fish, old bones, meat, &c., under certain conditions by certain Bacteria (as yet not precisely determined) pury be regarded as an example of one of these intermediate oxidisable substances. The oxidation in this case is accompanied by the evolution of light.

Special Study of the Occurrence and Distribution of Bacteria In the Atmosphere and in Potable Waters .- The ubiquity of Bacteria has been demonstrated by the use of sterilised nutrient fluids. If such a fluid be touched by a glass rod or by the finger or by any surface not chemically cleansed, Bucteria are conveyed into the fluid multiply there with enormous rapidity. Gelatin has been used as a means of studying the number of Bacteria present in the atmosphere or in a sample of water (v. Percy Frankland's rescarches [21]). However modified, the process is essentially this: a given volume of air is passed through a liquid so as to arrest all Bacteria previously floating in the air. The liquid is then mixed with gelatin, warmed to liquefy the gelatin, and rapidly cooled as a thin layer on a plate. The Bacteria develop at various separato points in the gelatin, giving rise to spherical growths or nests. These are then counted, and the species present may be discriminated by forther oultivation. Where water is the subject of investigation the gelatin is directly mixed with a given volume of the water. The results thins obtained have only a subordinate value from ths point of view of the hygienist. The majority of Bacteria are perfectly innocnous, and their presence is not-as has been too readily as- ately accompanied by practical results of ener-

-an indication of the probable p of pathogenic Bacteria. No such natural asso-ciation of pathogenic and innocuous Bacteria in definite proportions has been accertained, and its assumption is not warranted. It is necessary in all cases, if the results are to have hygienic value, to distinguish the kinds or species of Bacteria present and to ascertain their properties. Further, it is quite certain that all species of Bacteria will not flourish in gelatin even when mixed with peptons or such hodies. For instance, one of the most important pathogenio Bacteria -that concerned in tuberenlar consumption -will not do so. A special pubulum is needed for this Bacterium, and its presence would not be indicated by the ordinary gelatin cultivation of the contents of a given volume of air. Hence it seems necessary that in addition to careful discrimination of the Bacteria obtained by such experiments on atmospheric and aquatio distribution, there should be a systematic use of various cultivating media for the purpose of demonstrating the presence of various kinds of Bacteria. No doubt many kinds can be secured by the peptonised gelatin method, but if tha results of such studies are to have uny qualitative hygienio significance, other cultivating media must be simultaneously made use of. All the work at present done on this subject requires doing afresh from this point of view.

Special Study of Pathogenie Bacteria .-- A largo number of most important observations have been made of late years by pathologistsespecially by Lister, Pasteur, Koch, Klein, and their pupils-demonstrating not only the presence of Bacteria in the blood and tissues of mun and other animals when in a state of disease, but also preving in a certain number of cases that the Bacterin are the cause of specific diseasc. The proof, which is sufficient, and has been furnished in a limited number of instances, consists in-1. The constant presence of a definite form of Bacterium in the diseased unimal and in the specially-diseased parts of it. 2. Its successful removal from the diseased animal, and its pure cultivation on media free from all contamination by particles of the discased animal. - 3. The experimental introduction of the aultivated Bacterium into the body of a healthy animal liable to the disease in question but free from it -4. The subsequent development of the disease in the inoculated animal, This proof has been furnished in regard to the connection of B. anthracis with splenic fover in cattle and slicep, and malignant postule in man; in regard to 11. tuberculosis and consumption or phthisis in man and animals; in regard to B. cholera gallina and the cholera of fowls; in regard to Micrococcus erysipelatosus and erysipelas of man; in regard to certain Bacteria and septicionic and pyremic conditions in rats, mice, rabbits, and birds; and in regard to some other diseases of animals. Such a connection is strongly suspected, but not yet proved in the complete manner formulated above in regard to certain observed Bacteria or Micrococci, and the following diseases, viz. small-pox, scarlatina, diph. theria, typhoid fever, cholera asiatica, malaria, yellow-fever, gonorrhea, &c. The first definite researches in this direction, which were immedimous value, were those of Sir Joseph Lasyer, who showed that the blood-poisoning so frequent in crowded hospitals after surgical operations was due to the access of Bacteria to the wounds where they multiplied and manufactured poisonous products (sepsine, ptomaines?) which were absorbed into the blood. Lister adopted measures for preventing the access of these Bacteria, chiefly by the use of phenol and great clemniness in instruments, dressings, &c., and thus established the antiseptic system of surgery.

It is remarkable that the researches which have been made on the relation of Bacteria to disease have been mostly of a purely empirical character. Almost the only investigator who has carried the matter further (and with the most brilliant results) is the French chemist Pasteur. The fact is that the question us to what the Bacteria do after ordering un animal hody is, like the question of their action on substances extornal to the body, essentially a chemical one. Following up the observations of Toussaint, Pusteur [22] was hed to the discovery that the Bacterium outhracis when cultivated in broth could be made to assume a condition in which its virulence was greatly diminished. Nevertheless when introduced into the tissues of a sheep. the oultivated Bacterium multiplied, and as a consequence of its growth reinfered the sheep so treated resistant to the attacks of the virulent Bacterium anthracis taken from another unimal'e bloo.l.

It was known that an animal which had survived an attack of the virulent B. anthracis was thereby rendered 'immuno' to subsequent attacks, just us one attack of small-pox renders its survivor 'immune to the to that disease. Pastour conceived the theory that the Bacterium causing the disease in all such cases produces as a by-product -independent of its specific specific poison -a chemical substance which inhibits its inrther growth (as in the case of the alcohol produced by the yeast plant) and that this substance remaining in the anunal bedy protocts it from being the seat of further growth of the pathegonic Bacterium. The medified cultivated variety of B. anthracis equally produces this substance, and consequently acts as a protective against the incursions of the virulent lorm. Similarly cow-pox is to be regarded as the result of the growth of a medified small-pox Microccoous, and thus the protective effects of inoculation with cow-pox are to be explained. Applying this conception Pasteur has successfully protected while against fowl oholers, and has been led to his greatest triumph, the prosection by inoculation against rabies and the mecessful treatment of persons bitten by rabid logs.

It is extremely interesting and important to obsorve that the discoverios which have been made in this subject are due to chomical consequences. Novertheless there is much probability in the view put forward by Motschnikow a distinguished zoologist, now director of the Bacteriological Institute of Odessa) to the offect hat protective inoculation does not depend upon the development within the inoculated animal of a germioidal poison, resulting from the growth if the very germ which is killed or inhibited by that poison, but is rather due to the education

corpusales of the blood, which he has prevent attack and feed upon Basteris which are multiplying in the blood and tissnes. This property of the white corpusales leads Mctschnikow to term them 'phagocytes,' and, according to him, preventive inoculation depends for its efficacy on the fact that, having learnt to resist and destroy a weaker modification of a pathogenous Bacterium, they are able to deal subsequently with the more virulent variety; whilst it has been suggested that in the struggle between the phagocytes and the inoculated modified Bacteria, there must be a survival of the fittest and a consequent strengthening of the later generations of phagocytes in the protected animal.

However this may be, it is obvious that both the direct study of the chemical history of pathogenio Bacteria and the indirect euggestions derived from further knowledge of the chemical history of Bacteria of all kinds, are of an importance to human life and health which cannot

be over-estimated.

In connection with the study of the pathogenic Bacteria which attack man, it is necessary to insist that at present no one has attempted to determine the various kinds of Bacteria which are normally present on the surface of the human body, in the mouth, stomach, and intestines. There appear to be twelve or more present in the healthy human mouth (v. Vignal, [23]). So strangely has this matter been neglected that Koch of Berlin was ignorant, when he discovered the so-called 'comma-bacillus' in the intestines of cheleraic subjects in India, that an identical form occurs in the healthy human mouth, as shown by Lewis [24].

Spontaneous generation or abiegenesis .-Twenty years age experiments and observations were brought forward by various more or less competent observers [25] which were interpreted as proving the sudden formation of Bacteria aa living things in thuids containing the elements of protoplasm where no germ or living thing previously existed. It is sufficient to say here that these views had a valuable effect in stimulating the investigation of the life-conditions and activities of the Bactoria, but have been definitely proved to be erroneous and to have arisen from the imperfect state of knowledge as to the ubiquity of Bacteria and the pewer to recist the destructive effect of boiling water poesessed by the spores of Bacterium enbtile-the hay bacillus.

Conclusion.—An endless field of investigation is open in connection with the Bacteria.
It eccuis certain that in the near future we shall
be able to control the disease-producing forms,
whilst the euggestion presses itself that it may
be possible to entityate and intensify the activities of those which act as ecavongers and even
to lead some by appropriate methods to the
acquirement of new powers, or to the development of activities at prosent scarcely recognised.
There is no reason, from the point of view of the
biologist, why these lowest plants should not be
oultivated and specialised as breeds and varieties
for the cervice of mankind, as the peach and the
strawberry, the wheat and the cabbage have

Roblingreithy (referred to by ministrals in

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BALANCE v. ANALYSIS.

BALATA. A substance resembling caoutchone and gutta percha, obtained from the dried milky juice of the Bully-tree (Sapota Muelleri) (Sperling, Z. [2] 5, 480).

BALSAM. This term was originally con-

fined to a single substance called Balm of Gilead or Balsam of Judea, but is now employed gencrally to denote any liquid resin with aromatic They are composed of solid resins mixed with essential oils. Balsams of Peru, Tolu, Liquidambar, and Storax, contain cinnamie seid; Copaibs balsam, Mecca balsam, and

turperstine, do not.

PARABANIC ACID v. PARABANIC ACID.

from barwood (Baphus satista); Plates or needles (from ether), insol. water, v. sl. sol. benzene. Its alcoholio solution is turned red by air. Boiling squeous KOH converts it into baphic acid C₁H₂O₄ (?), baphinitin (O₄H₂O), and baphinitone, C₂H₂O₆. The latter gives a tri-bround derivative C₂H₃Br₂O₆.

BAPTISIA TINCTORIA. The root of this plant contains a crystalline alkaloid (Greens,

Ph.~(3)~10, 584).

BARBALOIN r. Alois, 1, 140, BARBATIC ACID C. H. O. [1860]. Occurs, to sether with usnic acid, in a lichen, Usnes barbata (St nhouse a. Groves, C. J. 37, 405; A. 203, 302). Needles or plates (from benzens). Decomposed by boiling milk of lime into CO. and (B)-orein.

BARBITURIC ACID

C, II, N, O, i.e. CII, < CO, NII > CO.

Malonyl-area. Formed by heating malonic acid with area and POCt, at 100° (Grimma, C. R. 88, 85; A. Ch. [5] 17, 276), or by reducing di-bromo-barbituric acid with sodium amalgam or HI. Prepared by heating alloxantin (1 pt.) with Il SO, (3 or 4 pts.) at 100° as long as SO, comes The product is poured into water.

Properties. Trimetric prisms (containing 2aq) from water. St. sol. cold, v. sol. hot,

water.

Reactions .- 1. Boiling aqueous potash Iorms malonic acid and urea (CO, and NH,). - 2. HNO, forms nitro barbiturie (diliturie) acid. -8. KNO forms nitroso barbituric (violurie) acid. -4. Br forms di-bromo-barbiturie seid .- 5. Heated with glicerin it forms dibarbiturio acid, urea, and ammonium maloburati. 3. Gyanogen Iorms a compound C₁II₄N₂O₃(CN)₂ aq, whence potash forms wyamuromalic acid.

Salts.-NII, HA": needles, v. sl. sol. cold water. — Na,A" 2aq. — K11A". — Ball,A" 2aq. — PbA". — Cull,A", 3aq. — AgHA". — Ag₂A".

Acetyl derivative C.H.AcN.O. product in the preparation of barbiturio sold from urea, malonic acid, and POCla. Powder. sol, hot water.

Di-methyl derivatives .- I. Malonyldimethyl-urea CO NMe.CO CH2 [1280].

From malonic acid, di-methyl-urea and PCl, or from di-methyl-urea and cyano-acetyl chlorids. CN.CH. CO.Cl (Mulder, B. 12, 466). Flat needles, v. sol. water. Forms a di-bromo-detivativo [175°-180°].

II. Di-methyl malonyl-ured

CO<NH.CO CMe, [265°]. From silver bar biturate and MoI (Conrad a. Guthzeit, B. 14 POCI, (Thorne, C. J. 89, 545). Plates (from water). V. al. sol. cold water. Boiling ROB Iorms di-methyl-malonic acid.

Salt .- Ag A" laq.

Ethyl derivative CO NH.CO CHUR [190°]. From ethyl-malonic soid, POClar urea. Gives a bromo-derivative.

Di-ethyl derivative CO NH.CO [182°]. From silver barbiturate and Etl.

Beneyl derivative
CO NH.CO OH.OH.Ph.

[2069]. From

benzyl-malonic acid, POCla, and ures.

Additional References. - Finck, A. 132, 304; Basyer, A. 130, 136; Conrad a. Guthzeit, B. 14, 1648 ; 15, 2844.

Di-barbiturie acld C.H.N.O. The ammonium sait is formed by heating barbiturie acid with glycerin at 150°. The acid is an insoluble powder; it gives a di-bromo- derivative.

Salts.-NH,HA". Na,A" 2aq. -KHA" xaq

(Baeyor, A. 130, 145).

BARIUM. Ba. At. w. 136.8. Mol. w. unknown, as V.D. not determined. Very little known of properties; denbtful if approximately pure Ba has yet been obtained. S.G. abt. 3.5-4 (sinks in H₂SO₄). S.V.S. abt. 36.5. Chief lines in spectrum are 5850, 5538, 4934, 4553, \$140 (Huggins, T. 154, 139).

Occurrence.- Not as metal; chiefly as sulphate (heavy spar), and carbonato (witherite); also as silicate in combination with silicates of Sr. K, or Al, and as oxide in combination with oxide of Mn. Ba compounds occur in many mineral waters; in the ashes of certain plants (Eckart, A. 100, 204); und in small quantities in sea water (Diculafait, A. Ch. [5] 15, 540).

Formation. - An amalgam of Ba and Hg is prepared in several ways:—(a) by electrolysing BaCl, mixed with a little very dilute 11ClAq, using an amalgamated Pt wire as negative electrode (Bunson, A. 92, 248); (b) by electrolysing moist BaO using Hg and Pt as electrodes (Davy, T. 1808, 303); (c) by bringing hot BaO or BaCl, into contact with vapour of K, and treating the product with Hg or by heating BaO with Na and treating with Hg (Kert. C. N. 31, 214); (d) by the action of Na anualgam on conc. warm BaCl, Aq (Crookes, C. N. 6, 194). By heating Ba amalgam in an atmosphere of H, or of hydrocarbon vapours, metallic Ba was supposed to be obtained; but Donath (B. 12, 715) asserts that it is impossible to remove all the Hg even at a white heat, and that the residue contains as much as 62 to 77 p.c. 11g.

Preparation .- By electrolysis of fused BaCl2 mixed with NII, Cl, in a porcelain crucible in an atmosphere of II; the positive electrode consisting of a cylinder of coke, the negative of an

iron wire (Matthiessen, C. J. 8, 291).

Properties and Reactions.—It is very doubtful whether the properties enumerated by different chemists as characteristic of Ba were determined by experiments made on even approximatelf pure metal. Davy described Ba as silver-white; Bunsen and Matthiessen as a golden yellow, slightly lustrous, somewhat malleable, metal, which melts at a red heat, but cannot be distilled. It is very easily oxidised, decomposes cold H.O readily, and burns in the oxyhydrogen flame. As no gaseous compounds of Ba have as yet been obtained, and us the spec. heat of the metal has not been determined. the value to be given to its atomic weight, the combining weight or equivalent having been determined, is arrived at chiefly by considering the analogies between the compounds of Ba and those of other allied elements, chiefly Ca, Sr, Mg, Zn, and Cd. These analogies lead to the formule BaX and BaY, for the compounds of

Bs. where X=0, B, SO; CO, de., and X=0. NO; Olo, de.; these compounds belong to one series. The valency of the atom of Ba in gaseous molecules is unknown. That the atomic weight of Ba is represented by a number the most probable value of which is about 136.8, provided the composition of the Ba compounds is expressed by the general formula BaX, where X = a monovalent atom or group of atoms, was established chiefly by analyses of barium chloride conducted by Marignae (A. 68, 215; 106, 165) and Dumas (A. Ch. [3] 05, 137). Ba reacts as a strongly positive metal; the salts obtained by replacing II of acids by Ba are stable and well marked; BaO and BaO₂H₂, BaS and BaS₂H₂, exhibit no acidic characters; BaO₂H₂ is distinctly alkaline, its heat of neutralisation is the same as that of soda and potash (Th. 1, 332) [BaO'H'Aq H SO'Aq] = 31,150; [BaO'H'Aq, 2HC|Aq] = 27,640. Ba combines with O and the hulogens with production of much heat and fermation of very stable compounds :- [Ba,O] = abt. 124 900; [Ba,Cl2] = 194,700; [Ba,Br2] = 170,000 (Th. 3, 266); these numbers are approximate only; they were determined indirectly, except that for BaO, but the Ba used was not free from Hg. Barium is very closely related to Ca and Sr, and less closely to My (v. art. Alkaline Eanths, metals of the).

Baryta was obtained by Scheele in 1774 from heavy spar; Davy in 1808 decomposed baryta by electrolysis; the metal was obtained approximately pure in 1855 by Bunsen and Matthiessen.

Combinations .- Very few compounds of Ba have been formed directly from the metal. It forms alloys with a few metals; that with mercury (v. supra, Formation) is a silvor-white body which rapidly decomposes water and cannot be separated into Ba and Hg by heat alone. Beketoff (A. 110, 375) obtained an alloy with aluminium, as a greyish solid with a tinge of yellow, by heating Al with BaO₂H₂ and a little BaCl, it decomposed H,O rapidly, but the water did not acquire an alkalino reaction. described alloys of Ba with lead, bismuth, antimeny, &c., obtained by the action of alloys of these metals with Na on molten BaCl. (A. 111.

Detection .- Many salts of Ba are soluble in water; some are insoluble; aqueous solutions in water; some are insolution, aqueous solutions of Ba salts are ppd. by cono. IIClAq or conc. IINO₂Aq. Insoluble Ba salts are decomposed by fusion with alkaline carbonates, giving BaCO₄ which dissolves in dilute acids. Fusible salts of Ba impart a pale yellowish-green colour to the non-luminous flame; the colour appears blue-green through a green glass. The emissionspectrum of Ba is characteristic; it contains very many lines in the green; about 1000 mgm. Ba may be detected by the spectroscope. Dilute sulphuric acid, or a dilute aqueous solution of sulphates, upt. white BaSO, insoluble in alkalis and dilute seids; 1 part Ba,2NO, in 100,000 parts of water gives an immediate pp.; one part in 400,000 gives a cloudiness on By this reaction Ba salts are disstanding. tinguished from Ca salts, and to some extent from salts of Sr.

Estimation .- 1. Bais usually determined as Baso, which is ppd from fairly cone solutions, containing a little HCl or HNO, by dilute H₂SO,Aq, the pp. is collected, well washed, and strongly heated, hafore weighing. If salts of Os are present, CaSO, may be removed from or to are present, CaSO, may be removed from the pp. by long washing with very dilute HClAq, or by digesting with Na,S,O,Aq (Dlehl, J. pr. 79, 30) which dissolves CaSO, but not BaSO, PbSO, if present, may be removed from the pp. by washing with solution of potash, or of ammoninm tartrate.-2. In presence of salts of Os and Sr. Ba is best estimated as BaSiF,, which is ppd. by freshly prepared H.SiF.Aq, followed by alcohol; after standing 12 hours, the pp. is collected, washed with a mixture of equal volumes of alcohol and water, dried at 100°, and weighed (v. also Rose, P. 95, 286, 299, 427)

Barinm, alloys of, v. Barrum; Combinations.

Barium, antimonates of, v. ANTIMONATES, under ANTIMONY, ACTOS OF.

Barium, arsenates of, v. assenates, under ABSENTO, ACIDS OF.

Barinm, arsenites of, v. ABSENITES, undor ARSENIC, ACIDS OF.

Barinm, bromide of. BaBr. Mol. w. nn-known, as compound has not been gasified. [abt. 812°] (Carnelley, C. J. 33, 280). S.G. 4.23 (Schiff, A. 108, 21). H.F. (Ba,Br²] = 169,960; [Ba,Br²,Aq] = 174,940 (Th. 3, 266).

Formation .- 1. By acting on BaO, H, or BaS, with HBrAq. -2. By adding BaS to an aqueous solution of Br. -3. Along with BaBrO,

by the action of Br on BaO₂H,Aq.

Preparation.—Aqueous HBr is neutralised by pure BaCO, the liquid is boiled down and allowed to crystalliso, and the crystals of BaBr, 2H₂O are hoated in a stream of dry air to 100°.

Properties and Reactions .- Crystallises with 2H₂O in white trimetric plates (Rammelsberg, 1'. 55, 237); according to Hauer (J. pr. 80, 230) and Werther (ibid. 91, 167) the crystals are monoclinie; the hydratod salt is perhaps dimorphous. H.F. [BaBr",2H O] = 9,110. [Ba,Br',2fI'O] =179,070 (Th. 3, 266). Heated to 75°. BaBr., H.O remains, and at 100° BaBr. is bable, 113 beauty 114; (60°) 123; (80°) 135; (100°) 149. It is also easily solublo in alcohol. S.G. 3-69 (Schiff, A. 108, 21). BaBr, is completely decomposed by heating to redness in dry O (Schulze, J. pr. [2] When cono. aqueous solutions of BaBr, and BaO are mixed so that the salts are present in the ratio BaBr.BaO, crystals of BaBr.BaO.5H.O (= BaBrOff.2H.O) separate out (Beelmann, J. pr. [2] 26, 388 and 474).

Barinm bromide, hydrated, v. Barness, pro-MIDE OF; Preparation.

Barinm, chloride of. BaCl. Mol. w. un-known, as compound has not been gasined. Enorn, as compound has not been gasified.

Bot. 860°, Carnelley, S.G. 375–389 (Schröder,

P. 107, 113). S.H. (16°-47°) 0902 (Kopp, T.

155, 71); (14°-98°) 0896 (Regnault, A. Ch. (3)

1, 129). S. (5°) 32·2; (30°) 38·2; (50°) 43°6; (80°) 52·4; (100°) 58·8 (Mulder; v. Michaelis'

Lehrbuch der Anorgan. Chem. 3, 0°0). S. (alzerrouen aer anorgan. Onem. 5, 0-0). S. (al-cohol 99 p.c.: 14°) 01; (alcohol at B.P.) 06 (Fresenins, A. 59, 127). H.F. [Ba,Cl⁻] = 194,740; (Ba,Cl²,Aq) = 195,810 (Thomsen). The following data apply to the hydrate BaCl, 2H₂O: much heat; Guntz (A. Ch. [6] 8, 5) gives 8.G. 8-052 (Schiff, A. 108, 21). S.H. (18²-46²) the values [BaOAq, 2HFAq] - 84,800; and

171 (Kopp, T. 185, 71). S. (18") 49-5; (108") 78. H.F. (Ba, Cl. 12HO) = 201,740; (Ba, Cl. 12HO) = 7,000 (Th. 3, 266). C.E. (cub. abt. 15"-900") 1000548 (Playfair a. Joule, C. J. 1, 131).

Formation.—1. By the action of Cl on hot BaO (Weber, P. 112, 619).—2. By passing HCl over hot BaO; light is evolved as well as heat: or by adding conc. HClAq to BaO, boiling down, and drving at 100°.—3. By dissolving BaS in cone. HClAq, boiling down, and drying at 100°

Preparation .- 1. Powdered withcrite (BaCO,) is added little by little to HUlAq; the solution is digosted in absence of air with more BaCO, (to remove iron, &c.), and is then poured off, evaporated to drynoss, and the rosidne heated to for some time .- 2. Two parts of finely powdered heavy spar (BaSO,) are heated in a crucible to redness with 1 part dry CaCl, and 2 parts iron filings; the fused mass is digested for a short time with 6-8 parts boiling water (by long digestion BaSO₄ and CaCl₂ are re-formed), the liquid is filtered from FeS, CaS, and undecomposed BaSO₄, made slightly acid by HClAq, and evaporated to dryness at 100° 3. The solution of MuCl, which is obtained in making Cl from MnO, is noutralised by BaCO, or CaCO, and ovaporated to dryness; the residue is heated with heavy spar and conl; the mass is lixiviated (MnS, FeS, and some BaSO, remain), the liquid is treated with a little MnCl,Aq to decompose any BaS present, HClAq is added, and the whole is evaporated to dryness (Kuhl-

mann, C. R. 47, 403, 464, 674).

Properties.—White salt, easily soluble in water, [BaCl², Aq] = 2,070 (Th. 3, 266); slightly soluble in alcohol (v. supra); solution has a bitter taste and is noicenam. Melts at red heat and cools to an opaque mass. Reactions, —1. Heated in steam, HCl 4s

evolved, and residuo has an alkaline reaction .-2. Partly oxidised by fusion with potassium chlorate, but unchanged by heating in dry oxygen (Schulze, J. pr. [2] 21, 407) .- 8. Completely decomposed by fusion with silicates.

Combinations, -1, Conc. solution of BaCl. mixed with cone BaOA, pps, thin transparent plates of BaCl, BaO.5H, O (= BaClOH.2H,O) (Bechmann, J. pr. [2] 26, 388, 474).—2. Combines with water with production of heat, [BaCl-21f-O] = 7,000, to form the hydrate BaCl_2H,O. This hydrate crystallises in white flat trimetric plates, which are not efflorescent; they lose 2H,O at 100°, but take it up again la moist air. A cono. solution is decomposed to Ba.2NO, and NaCl by heating with NaNO,
Barium chloride, hydrated, v. Barium,

Barium corrice, hydrated, v. Danion, culcance or; Combinations, No. 2.

Barium, cyanide of, Bar(CN), Obtained by action of HCNAq on BaO, II. (v. Cyanidas).

Barium, fluochloride of. BaFCI (v. Barium,

PLICATION (INC.)

Barium, finoride of. BaF. Mol. w. unknown, as compound has not been gasified,
[abt. 908°] (Carnelley, C. J. 33, 280). S.G.

134.55 (Bödeker).

Preparation.—1. By ppg. Ba2NO,Aq by NaFAq.—2. By the action of HFAq on BaOAq, or on freshly ppd. BaCO,, and evaporating. The former action is attouded with the production of much heat; Guntz (A. Ch. [6] 8, 5) gives

Properties and Reactions .- White, finely, granular, crystals; scarcely solublo in water but easily in HNO, Aq, HClAq, and HFAq. Not decomposed by heat alone.

Combinations. -I. With BaCl, to form BaF, BaCl, (-BaFCl); obtained by adding NH, Aq to a solution of BaF, in HClAq; also by fusing 1 part Naf with 6-8 parts BaCl,, and digesting with water, also by adding KFAq to BaCl_Aq and ovaporating. Forms white granular crystals, more soluble in water than Bak.: partly decomposed, with loss of BaCl, by lour . continued washing with water .- 2. With BF, to form BaF, 2BF, 2H,O (=Ba(BF,), 2H,O); obtained by acting on BaCO, with IIBF, Aq, and evaporating; boric acid separates, and afterwards the double salt (v. bosoficombes under Boson, FLUORIDE OF). 3. With SiF, to form BaF, SiF, (-BaSiF.); obtained by adding Il.SiF.Aq to a solution of a Ba salt. White solid, very slightly soluble in cold water S. (17°) '03 - and only slightly soluble in HClAq. S.G. 4.28. Leaves Bar, when heated; heated with NII,Cl gives residue of BaCl, (Stolba, J. pr. 96, 22) (v. SILICOPLEORIDES, under Silicon, FLUORIDE OF).

Barlum, hydrexide of. BaO.H. (Caustic baryta). Mol. w. unknown, as compound has not been gastifed. S.G. 4-195 (Filhol, A. Ch. [3] 21, 415). S. (0°) 1.5, (5°) 1.75, (10°) 2.22, (15°) 2°89,(20°) 3°48,(25°) 1·19, (30°) 5·0,(35°) 6·17, (40°) 736, (45°) 912, (50°) 1175, (55°) 1471, (60°) 1876, (65°) 21·67, (70°) 31°9, (75°) 56·85, (80°) 90·77 (Rosensthiel a. Rühlmann, J. 1870, 314). H.F. (Ba, O, H°O) · 146,500, value only appr (Th. 3, 266), approximate; [DaC; [110] = 22,260

Formation. -1. By heating heavy spar with earbon, dissolving BuS formed in hot water, filtering, adding CuO or ZnO to decompose the BaS, illtering, evaporating to dryness, and heating to redness (Müller, J. pr. 82, 52; Stablachmidt, D. P. J. 182, 30; Nicklès, W. J. 1869, 271). 2. By heating Fe with Ba.2NO, to redness, dissolving in water, filtering, evaporating, and heating the residue.—3. By the action of stoam on BaCO, (Lenoir, W. J. 1867.

256).

Preparation. - 1. Water is added little by little to BaO (q. v.); the product is heated to dull redness in a silver dish .- 2. Aqueous solution of pure NaOlI, S.G. about 1.1 to 1.15, the quantity of NaOH in which is accurately known, is thereof to boiling, a quantity of powdered Ba2NO, is added equivalent to the NaOH used, the liquid is boiled for a little, if solution is not complete water is added, the hot llquid is filtered quickly and allowed to cool in a closed vessel when crystals of BaO.H. 8H.O are deposited; these crystals are separated, recrystaillsed from boiling water, and heated gradually to redness in a silver dish (Mohr, Ar. Ph. [2]

Properties and Reactions .- A white powder, dissolving in water (v. supra) to form an alkaline, caustio, liquid; melts at a full red heat and earstallises on cooling; not decomposed by heat sleue, but by heating in a stream of air BaO and H₂O are produced. Aqueons solutinn is warkedly alkaline, and neutralises acids with

BaO'H', 2HF) = 71,400 (solid BaF, is produced) production of same quantity of less as KOHLE by action of gascons HF on solid BaO,H.). Cl, except in presence of H₂O; action is then probably 6BaOAq + 6Cl₂ = 5BaCl₂Aq + Ba(ClO₂),Aq (Weisberg, B. 12, 846). Is not acted on by CO, (Scheibler, B. 19, 1973).

Combinations .- With water with production of heat [BaO'H',8H'O] = 27,470 (Th. 8, 266) to form crystals of BaO,H,28H,0 (Beokmann, J. pr. [2] 26, 388, and 474; Filhol found 7H,O, Noad and othors 9H₂O). (For preparation of these crystals v. supra.) Those crystals lose 7H₂O over Il, SO, in vacuo, or by heating to 75°, and tho eighth H2O at a red heat; they dissolve in about 3 parts boiling water, and 20 parts water at 15°. The solution is attended with disappenrance of heat [BaO²H².8II²O,Aq] = -15,207 (Th. 3, 263), Crystals of BaO₂H₂,H₂O nelt at 83°-85° (Voley, C. J. 49, 371). According to Bechmann (J. pr. [2] 26, 388 and 474) pure BaO is obtained by heating BaO₂H₂,8H₂O in a stream of ().

Barium, iedide of. Bal. Mol. w. unknown, as compound has not been gasified. S.G. 4-92 (Filhol, A. Ch. [3] 21, 415). 11.F. [Ba,I'Aq] = 144,520 (Th. 3, 266).

Formation and Preparation.-Similar to methods for BaBr₂ (q. v.): also by action of gaseous Ill on BaO.

Properties and Reactions .- A white, nondeliquescent, solid; easily soluble in water or alcohol; not decomposed by heat in absence of air, in prosence of air BaO is formed and I evolved; wholly decomposed by heating in O (Schulze, J. pr. [2] 21, 407); aqueous solution absorbs CO₂ from air.

Combinations .- 1. With water to form Bal. 711.0 (Croft, J. pr. 68, 402; Thomsen, B. 10, 1343; Werther, J. pr. 91, 331, says the crystals are Bal. 211.0). This hydrate forms needle-shaped orystals which deliquesce, with partial separation of I, in moist air, and melt on heating; heated in absence of air Bal, on heating; neared in absence of air Baight remains, one H.O is lost at 100°, 5H.O at 125° and the seventh H.O at 150°. Thomsen gives these data [Ba, 1², 711²0] = 151,370; [Bai?-11²0,Aq] = -6,850,—2. With baryta to form BaO.BaL.5H.O(= BaIOH.2H.O) mann, J. pr. [2] 26, 388 and 474); this salt crystallises from a mixture of cone. colutions of its constituents, in the ratio BaO:BaI,.

Barium iedide, hydrated, v. Barium, rodida or, Combinations, No. 1.

Barium, oxides of. Ba forms two oxides, BaO and BaO,; the former is produced by the action of dry air, or O, on Ba; BaO heated to about 450° combines with O and forms BaO. which is again reduced to BaO at a higher teraperature, or by reducing the pressure at 450°. Dry BaO₂ is stable, but the presence of water brings about slow decomposition to BaO₂H₂+O; Berthelot (A. Ch. [5] 14, 438; oomp. C. R. 85, 880) gives these data [BaO,0] = -6.050; [BaO',H'O] = 2,760 (giving BaO₂H₂ + O). BaO is a strongly basic oxide; BaO, evolves O (or H₂O₂) and forms the same salts as BaO when acted on by acids.

I. BARIUM MONOXIDE (Baryta) BaO. Mol. W. nnknown as compound has not been gasified. S.G. 4.65 (Playfair a. Jonle, C. S. Mem. 8, 84) S.G. crystals 5.722 (Britgelmann, W. 2, 466;

Soheele distinguished baryta from lime in 1774; Gahn recognised the presence of this earth in heavy spar ; Bergmann called the earth terra ponderosa; Kirwan gave the name baruta; Davy, in 1808, proved it to be a metallie oxide.

Formation.-1. By the action of dry air on Ba .- 2. By strongly heating BaCO, best with does not again combine with the BaO. 3. By strongly heating Ba.2NO,; Rammelsberg (B. 7, 542) says that an oxide with the composition Ba,O. (- 2BaO.BaO.) is thue produced; Brugelmann (W. 2, 466; 4, 277) obtained hexagonal crystals of BaO by this method .- 3. By strongly heating BaCl, or BaSO, to white heat, in a ourrent of steam.

Preparation .- 1. Ey strongly heating Ba(IO,)2 in a porcelain cruck to until all I is removed .--2. By heating dry Ba.2NO, in a capacione porcelain vessel (best a retort), gradually raising the temperature when the salt melts, again raising the temperature to full redness when the residue in the vessel re-solidities; the heating must be continued until all nitrate is decomposed, but no longer, as on long-continued heating CO. is absorbed; the portions in contact with the porcelain take up a little SiO, and Al₂O₃.

Properties.—A grey-white powder, very poisonous; melts at white-heat; takes up H.O. and CO, from the air; dissolves in water to form a caustic alkaline colution (v. Banum,

HYDROXIDE OF).

Reactions .- I. Reduced by heating with polassium .- 2. Decomposed to Ba + O by electrolysis .- 3. Heated in chlorine, BaCl, and O are formed .- 4. Heated with sulphur, BaS and BaSO, are produced .- 5. Heated in carbon disulphide vapour, the products are BaS along with BaCO₂...6. Heated in phosphorus vapour, in presence of H, barium phosphide BaP₂ (q. v.) and Ba₂P₂O₂ are formed (Dumas, A. Ch. [2] 32, 364) .- 7. Heated with arsenic vapour barium arsenite (q. v. under Arsenites) is eaid to be produced.

Combinations .- 1. Combines with water to form BaO₂H₂ (q. v.) with production of much heat, and increase of volume: [BaO, HO] = 22, 260; [BaO, 9HO] = 49, 730 (Th. 3, 266). 2. With carbonic anhydride, to form BaCO, (dry BaO has no action on CO₂; Scheibler, B. 19, 1973); with sulphuric anhydride to form BaSO,; [BaO, CO²] = 62, 220; [BaO, SO²] = 110, 590 (Th. 3, 266).—3. Hented in air or crygen to about 450°, forms BaO, (q. v);—4. With methylic or ethylic alcohol, forms BaO.2CH,O or BaO.2C,H,O.

II. Bantum Dioxide. BaO₂. Mol. w. unknown. S.G. 4-96 (Playfair a. Joule, C. S. Mem. 3, 84). Discovered by Thenard (A. Ch. 8, 308).

Formation .- 1. BaO, or a mixtare of BaO,H, and CaO or MgO, is heated in nearly dry air, or O, to dull reduces in a glass or porcelain

First to reduces? the RCI formed is disadved out by cold water (Liebig a. Wöhler, P. 96, 173); the impure hydrated BaO, containing BaO (Herthelot, A. Ch. [5] 6, 207, says the residue is nearly BaO.BaO.) is rubbed in a mortar with water, and added little by little to very dilute HClAq, bat not in quantity sufficient to noutralise the acid; tha colution (which contains H.O.) is filtered, made elightly alkaline by a ldition of dilute BaOAq, whereby alumina and iron oxide are ppd., the liquid is again filtared through linen, and an excess of BaOAq is udded; lustroue plates of BaO .. 8H O are ppd. : (the filtrate must contain H2O2, proved by the production of a blue colour in other when shaken with other after acidifying and adding diluta K.Cr.O.Aq); the pp. is washed with cold water, pressed between filter paper, and placed over H₂SO₄ until all water is removed and BaO₄ remaine (Berthelot, A. Ch. [2] 6, 207). Or, H O.Aq is added to BaOAq, the pp. of BaO, 8H,O is washed with cold water, pressed between filter paper, and heated in dry air free from CO, to 100°-120° (Schone, B. 6, 1172). - 2. Pure BaO is heated to low redness in a stream of O (Brodie, T. 1850, 775).

Properties. - A white powder, resembling MgO; insoluble in, and combines with water; melts at full red heat with evolution of O.

Reactions .- 1. Decomposed by heat to BaO and O; at slightly reduced pressures (750 to 730 mm.) decomposition begins at about 450°; at ordinary pressure at a higher temperature than this; if the BaO produced is allowed tocool to 450° in presence of O under reduced pressure BaO, is va-t rined (Bonssingault, A. Ch. [5] 19, 464) - Ection each very slowly by cold, quickl, by hot, water, forming BaO, H, O [BaO, H, HO] = 2,760 (giving BaO, H₂ + O, Borthelot, A. Ch. [5] 14, 433) .- 3. Conc. sulphuric acid forme BaSO, and evolves O at temperatures above 602-702, but O mixed with ozone at lower temperatures (Houzenu, C. R. 40, 949) .-4. Heated in dry carbonic anhydride, BaCO, and O are produced. 5. Heated with carbon monoxide, or sulphurous anhydride, light and heat are produced, and BaCO, or BaSO, is formed (Wöhler, A. 78, 125).—6. Acts as a powerful oxidiser towards carbon, phospherus, No. &c. (comp. Slater, J. pr. 65, 253; and Brodie, T. 1862, 837].—7. With dilute acids forms salts of Ba, and ll,O, or O.

Combinations. - 1. With water, combines to form BaO.8H.O (produced also by stion of BaOAq on H.O.Aq; v. Prepara.com? prismatic dimetric crystals, which lose 8H.O in vacuo, or by heating in absence of CO, to 100°-120°. Berthelot gives the formula BaO, 10H,O to the hydrate (A. Ch. [5] 21, :57); he also describes another hydrate with 711,0 (l.c. [5] 6, 207); he gives the data (BaO', 10H2O) - 9,100 (Lc. [5] 14, 438).—2. With hydrogen peroxide forms very nustable, monoclinio, crystale, BaO, H.O. produced by adding excess of H.O.Aq to BaOA or by adding NH, Aq to the bound of the mixed with H₂O₂ (Schöne, A. 192, 257).

BARIUM, or by adding NH, Aq to the solution of a Ba sale

tabe.

Sulphines of; monosulphine, Reactions.

Preparation.—A mixture of 4 parts finely powdered KClO, and 1 part BaO is thrown little by little into a porcelain crucible heated produced by passing H charged with P vapour

ever hot BaO; decomposed by H.O. giving PH. and BaHPO, (Dumas, A. Ch. 32, 364).

Barium, salts of. Salts produced by replacing H of scids by Bs; they form one series belonging to the form BaX, where X=Cl, &. O, SO, CO, &c. As none of these salts has been gasified we do not know the molecular weight of any of them; the spec, heat of Ba is undetermined; the formula are, therefore, based on analogies between these salts and those of similar metals which form gasifiable compounds, especially Zn and Cd, and also on analogics between the salts of Ba and Ca, the atomic weight of the latter metal having been settled by the spec, heat method. Barium forms salts with most, if not all, the acids; very few basic salts are known, and those which have been prepared are generally salts of the weaker acids, e.g. borie, tungstic, molybdic, &c. Tho haloid salts are very stable; the carbonate, nitrate, iodate, chlorate, &c., are decomposed by heat; Ba salts of the oxyacids are reduced by heating with C, II, or CS₂. Most Ba salts are isomorphous with the corresponding salts of Ca and Sr; many with the corresponding salts of Pb. A few Ba salts are soluble in water; the greater number are slightly soluble only, or insoluble (r. Bohates, Carbonates, Phosphates, Sulphates, &c., &e.).

Bariam, ssleaids of. BaSo. Mol. w. nuknown. White solid, changing in air, obtained by heating BaSeO, in H to dull redness (Fabre, C. R. 102, 1469).

Barlam, ssisacovanids ef. BaSe₂(CN)₂(?). Prepared by Crookes (J. pr. 53, 161). Data very meagre.

Bariam, sllicoflaorlde ef. BaS. v. Barium

Barlum, salphidss and hydrosalphids (or sulphydrate) of. Three sulphides of Ba are known; a fourth probably exists in solution. The monosulphide BaS is obtained by ficating BaO in a stream of II.S; by heating BaS + 28 to 360°, the trisulphide BaS; is formed; by boiling BaSAq with 3S and crystallising, BaS, may be prepared; and if BaSAq is boiled with considerable excess of S the solution reacts as if it contained a pentasulphide BaS. Only one hydrosulphide or sulphydrate, BaS.II., la known. The sulphides and the hydrosulphide are fairly stable compounds; they are solublo in, and partly decomposed by water; they resemble the sulphides of the alkali metals in their reactions, and BaI.I.Aq reacts with As.S₃ to form barium thio arsente Ba.Aa.S₃ (q. v.). Sabaticr (A. Ch.) [5] 22, 1) gives the thermal data:—[BaO, II.S] = 22,100; [BaS, O] = 236,500; [BaS, Aq] = 7,000.

I. Mosesulrupr. BaS.

Formation.—1. BaO is heated in a stream of H₂S.—2. BaSO₄ is reduced by heating in H or coal gas.

Preparation.—1. A stream of CO₂ is passed through CS₂ and then over red-hot BaCO₂; CS₂ must be in excess as BaS is decomposed by CO₂; the product is freed from higher sulphides by heating in If (Schöne, P. 112, 193).—2. BaO.II.—H₄O (prepared by heating BaO.H.—8II.O 60° in H) is acted on by dry H.S; the groducts are BaS and II₂O (Veley, C. J.

Properties.—Awhiteamorphoussolid; soluble in water; exposed to sunlight and then placed in the dark, it gives off light; oxidised in moist air.

Reactions .- 1. In moist air decomposes to BaCO, and BaS,O, with evolution of H.S .- 2. Heated in air is slowly oxidised .- 3. Heated in steam, BaSO, is formed and H evolved (Lauth, C. C. 1863. 880).-4. Chlorine, bromine, and iodine, decompose BaS, forming BaX, (X=Cl, Br, or 1) and S.—5. Dilute acids form Ba salts and ovolve H₂S.—6. Water brings about partial decomposition into BaS,H, BaO,H, polysulphides and oxysulphides of Ba (v. Veley, C. J. 49, 369). The action of water on crude BaS has been examined in detail by H. Rose (P. 55, 415). If hot water is added in quantity just sufficient for solution, the liquid gives a pp. of MnS, without evolution of H2S, on addition of an aqueous solution of a neutral manganous salt; the solution, therefore, contains either BaS or hydroxide and hydrosulphide in the ratio BaO.H.:BaS.H.;

(7BaO.H.Aq+BaS.H.Aq+2MnCl.Aq=2MnS+2BaCl.Aq+211.0). If cold water is added to crude BaS in an open vessel, in quantity rather less than sufficient for complete solution, and the liquid is evaporated, BaO.H. separates out, then various oxysulphides (v. infra), then, on evaporating the mother liquor in a retort, crystals of BaS.611.0 (v. infra, Combinations) separate, and finally on evaporating to dryness BaS.H. remains.

The oxysulphides prepared as above described, or by cooling the solution obtained by acting on crude BaS with boiling water in a closed vessel, seem to be three:

Ba,O,S.,58II,O[=4(BaO,H.,9H,O).3(BaS,6H,O)], Ba,O,S.,58II,O[=(BaO,H.,9H,O)(BaS,H.O)], and Ba,S,O.28H,O[=(BaO,H.,9H,O).3(BaS,6H,O)], and Ba,S,O.28H,O[=(BaO,H.,9H,O).3(BaS,6H,O)], The compositions of these bodies are, however, far from settled; the compounds are very unstable and are separated by recrystallisation into BaO,H., and BaS,H., If successive quantities of cold water, each less than sufficient for complete solution, are shaken with crude BaS in a closed vessel for some hours, the first solution contains BgS,II., along with a little of the higher sulphides of Ba (the solution gives MnS and also H,S on reddition of MnCl,Aq); the next solution contains either BaS or BaO,II. and BaS,H, in the ratio BaO,II,BaS,H, (with MnCl, it gives MnS without evolving H,S); the following solutions contain BaO,H., as they give more and more MnO,II., on addition of MnCl,Aq and less and less MnS.

BaO.H. H.O (prepared by heating BaO.H., 8H.O by 60° in H) is acted on by dry H.S; the products are BaS and H.O (Veley, C. J.

by evaporating Ball. Eq. (g. a.) accoming Ball. GH.O crystallises in white six-sided places; slightly soluble in cold, easily in hot, water; insoluble in alcohol; loses GH₂O between 100° and 850° with partial decomposition

with partial decomposition.

II. Tarsulphime. BaS, Prepared by heating 2 parts BaS with 1 part S, and removing excess of S by distilling it off at 350° (Schoue,

ing 2 parts BaS with 1 part S, and removing excess of S by distilling it off at 350° (Schoue, P. 112, 193). Forms a yellowish-green mass, soluble in hot water; heated to rednsss in absence of air gives BaS+2S. A solution of BaS, in much boiling water evaporated in vacuo deposita (1) BaS.511.0 (v. supra), then (2) a mixture of BaS, H.O (v. infra) and orange dichroio monoclinio prisms of 3(BuS.6H.O.(BaS, H.O.(6H.O.(Schoue, L.C.)).

HI. TETRASCEPHIDE. Known only in combination with H₂O as BaS₂H₂O. By evaporating a solution of BaS₃ in hot water in racuo, or by evaporating BaSAq with 3S, trimetrie, dichroic, needles separato; yellow by transmitted, red by reflected, light; soluble in water, may be reorystallised from hot water; insoluble in alcohol; at 300° loso H₂O with decomposition into H₂S, S, and BaS₃ (Schöne, l.c.). A more hydrated salt, probably BaS₂-2I₂O, was obtained by Velcy by dissolving S in BaS₂H₂Aq (C. J. 49 378).

IV. PENTABULPHIDE. BaS., Not known in definite form. BaSAq or BaS.H.Aq boiled with excess of S. yields a yellow alkaline solution, from which on cooling S and BaS, separate out; the mother liquor contains Ba and S in ratio BaS, on evaporation crystals of S separate out, and BaS, remains in solution (Schöne, Lc. confirming older observations of Berzelins).

V. Hydrosulphide of Sulphydrate, BaS.H., Formation, — By action of 11.0 on BaS (v. Monosulphide; Reactions, No. 6).

I'rrparation.—BaOAq (saturated at 100) is saturated with ILS at 60°-70°; the liquid is decanted in absence of air, and is cooled to about 10°; the crystals of BaS₂H₋₄II₋O which separate are dried between paper out of contact with air, and then heated in a stream of H (v. Veley, C. J. 49, 369).

Properties and Reactions.—With 4H O forms white acicular cryetals, which effloresce in air, and gradually absorb O, forming BaS.O, and BaS.O,; these crystals are soluble in water but insoluble in alcohol; aqueous solution evolves H₂S when boiled; heated to redness out of contact with air, H₂S is removed and BaS remains (for details, v. Veley, Lc.). BaS.H₂ is strongly basio in its reactions; e.g. with As.S₂ it forms Ba thie-arsenite.

Barium, snlphocyanide of. Ba(SCN)₂. Obtained by decomposing NH, SCNAq by BaOAq (v. SULPHOCYANDES, under CYANDES).

Barinm, thio-antimonate of. Ba₂(SbS₂)₂. Obtained by the action of BaCl₂Aq on Na₂SbS₂Aq (compare Thio-ANTIMONATES under ANTIMONY, THIO-ACIDS of).

Barium, thic arsenite of. Bs. As. S., Obtained by digesting Bs. S. H. Aq with As. S.; and Barium thic-arsenates Ba. (As. S.), and Ba(As. S.), obtained by the action of H. S. on Ballaso, Aq (b. THIC-ABSENITES and THIC-ARSENATES nuder ARSENIC, THIC-ACTES OF).

M. M. P. M.

DARTIA. Colds of Barton, v. Bikitis,

The characteristic reaction of an BARE. a id is that the whole, or a portion of the hy-drogen of an acid can be displaced by a metal, with production of a now body, called a salt, composed of the mstal and the elements of the acid, excepting the displaced hydrogen (v. Acids). If the oxide of a metal reacts with an acid to form a salt, the hydrogen displaced from the seid combines with the oxygen of the oxide to form water; the products of the reaction are a salt and water. The salt is not characterized by the properties either of the acid, the metal, or the metallic oxido; it has been built upon the metal or metallic oxide by combining this with the scid. The name base was given by Rouelle in 1744 to those bodies which reacted with acids to form salts. The name has some times been applied to metals, as well as to exides and hydroxides of metuls; at other times it has been confined to compounds of metals with H and O; at all times the conception underlying the name has been that of a substance which, while chemically very unlike an acid, reacts with acids to form salts. The dual origin of a salt is implied in the statement that for its production there is required the interaction of an acid and a base. A definition of any one of the terms, acid, base, salt, implies a definition of the other two. The chemical reaction oharacteristic of bases, as the term is now used, is the production of salts by the mutual reaction of a base and an acid; in some cases water is also formed, in other cases the salt is the sole product. Typical reactions are as follows:

K,OAq + H,OAq - K,SO,Aq + H,O 2KOH q + H,SO,Aq - K,SO,Aq + 2H,O 2KOH Aq + H,SO,Aq - (NH) SO,Aq

2NII Aq + II SO Aq (NII) SO Aq.

A base may then be (i.) a metallic oxide, (ii.) a metallic hydroxide or an allied compound such as NEt. OH, PMc, OH, SEt. OH, &c., (iii.) ammonia or a derivativo thereof, e.g. NILEt, NMe, &c. The terms strong and uvak may be applied to bases with meanings similar to those given to the terms when applied to acids; a strong base, in this sense, is one which, when it reacts in aqueous solution with another base and an acid-all being present in equivalent quantities, and all possible products being soluble in water—combines with a large proportion of the seid and leaves only a small proportion for the other base to combine with. The hydroxides MOH, SEt,OH are noarly as strong bases as the alkalis (v. Arrinity). By the term a strong base is sometimes meant a base which reacts with various acids to form very stable salts; e.g. salts which are not changed by water, hot or cold. In this meaning of the term BaO.H. is a strong base, but BiO.II, or SnO.H. is a weak base. The oxides and bydroxides of polyvalent metals appear to be weaker bases than the corresponding compounds of the monovalent metals. The more positive a metal is, the more basic are its oxides and hydroxides. Sometimes a metallio oxide, or hydrated oxide, may react towards strong acids as a base, and towards strong bases as an acidic oxide; thus

Al.O. BEO. Teache with H.SO. As to form Al.O. BEO. (= Al., SEO.), but Al.O. 2H.O reagts with much KOHAq to form K.O.Al.O. (= K.Al.O.). In some cases the basic and acid o functions of a compound may be nearly equal; thus amido acetic acid (? CII, NII, COOH) forms salts by its reactions with bases as other acids do, but it also combines with acids, as NH, does, to form salts. The hydroxides of certain metals which in some of their reactions behave as non-metals react as bases towards most soids, but if oxygen is added to these hydroxides compounds are formed which react as bases only towards the stronger soids and at the same time react as acids towards the stronger bases; such compounds are SnO2II2, and SnO.O2II2, respectively. Bases are sometimes divided into mono-acid, di-acid, tri-acid, &c., according as one reacting weight interacts with one, two, three, co., reacting weights, of a monobasic acid, to form a salt. The poly-acid bases are weaker than the mono-acid bases. As examples of mono-soid bases may begiven KOH, NH, NILEt, &o.; of di-acid bases, CaO,H., BaO,H., ZuO,H., NH,C,H., &c.; of tri-acid bases, FeO,H., C,H,(NH,),OH,&c.; of tetra-acid bases, ZrO,H., do. (comp. Acids and Salts). M. M. P. M. BASES, ORGANIC, v. Alkaloids, Amines,

AMIDES, AZINES, PYRIDINE, QUINOLINE, &c. The nomenclature of bases containing earbon

and nitrogen in one ring is as follows: N:CII Pyridine Pyrazine (Ketiver) Purimidine. HC:N'CH HC:CH.CH lic:n.ch

HC:N.CII HĊ:N.ČH • N:CH,CH Isoquinolene Quinoline CII:CII ,CII:CH N : ĊII CII:N

Quinazolina Quinoralino Cinnoline CH. CH:N N:CIL CH:CH

BASIC OXIDES. Oxides which react with acids to produce salts. The greater number of tho metallic oxides are basic; oxides of well-marked non-metals are never basic. The correlative term is acidic oxides (v. Bases, Acros, Salts). M. M. P. M.

BASICITY OF ACIDS v. Acids, pasicity or, BASILICUM, OIL OF. The ossential oil obmined by distilling the leaves of Ocymum basilisum with water contains C10 II 10 3II O which crystallises in prisms (Dumae a. l'eligot, A. 14, 75).

BASSIA LATIFOLIA. The seeds of this

Himalayan plant yield by pressure a buttery Formed by the action of alcoholic potash ca

(Hardwick, U. J. S. 201). BASSORIS. The insoluble gum said, pro-

bably meta-arabic acid, or at least a meta- acidallied thereto (v. Arabin) of gummi bassorm, G. Toritonense, or G. Kutera. These gume consist of a part (the meta- acid) that swells up to a jelly when they are treated with water, and of a soluble part, the alkaline or earthy salt of the soid.

BASTOSE v. CELLULOSE.

BASYLOUS. A name sometimes applied to the more positive, usually oxygen-containing, radieles, or groups of atoms, which combine with more negative, or chlorous, groups to form

salts; e.g. K2O.SO3, K2O.CrO2, Cr2O3.3SO3, &c. The name is sometimes also applied to the elements which displace H from acids with formation of salts. The correlative term is M. M. P. M. chlorous.

BDELLIUM. A gnm-resin (Johnston, J. pr.

26, 145) BEBEERINE C, H2 NO. Bebirine. [180°].

Occurs, together with a resin (sepirin) and an acid (bebirio acid) in the bark of the bebeeru tree of Demerara (Rodie; Maclagan, A. 48, 106; Muclagan a. Tilley, P. M. 27, 186; v. Planta, A. 77, 333). It is an amorphous powdor, v. sl. sol. water, v. e. sol. alcohol, v. sol. ether. - B', H, PtCl4: orange amorphous pp. Buxine has been considered to be identical with bebeerine (Walz,

N. J. P. 14, 15). BEE'8 WAX v. WAX. BEHENIC ACID C.H.O. Benic acid. [76°]. Occurs as glyceride in oil of bon and in Benic acid. the fatty oil of black mustard seed. Needles, resombling stearie acid. - NaA'. - BaA'. PhA',-EtA' [49°] (Voolcker; Strecker, A. 64, 346)

BEHENOLIC ACID Cz.H.O. Benolic acid. [57.5°]. Formed by the action of alcoholio potash on di-bromo-behenic acid (Haussknecht, A. 143, 41). White needles (from alcohol); v. sol. alcohol and other, insol. water. Not reduced by sodian-amalgam, but combines with Br., forming C.,11,,Br.O., [47°], and with Br., forming C.,14,Br.O., [78°].

Salts.—MgA', 3aq.—AgA'.—BaA',

BELLADONNINE. An alkaloid occurring in

the mother-liquor from which sulphate of atropine (q. v.) has been crystallised (Hübschmann, Schweiz. Z. Pharm. 1858, 128; Krant, A. 148, 236; B. 13, 165; Ladenburg a. Routh, B. 17, 152; Merling, B. 17, 381). Amorphous, v. sl. sol. water, v. sol. alcohol, other, and chloreform. It is but slightly attacked by boiling barytawater, but is split up by alkalis into tropio acid and oxy-tropine C.H., NO. This would indi-cate that belladonnine is oxy-atropine C., H., NO. (L.). According to Merling, belladonnine is C₁, If NO, and gives tropine, atropic acid, and iso-atropio acid when boiled with baryta-water.

Salts: B',H,PtCl,...-B'HAuCl,... BEN, OIL OF. A fatty oil expressed from the fruits or Moringa nux behen. It contains glyceryl palmitate, stcarate, cleate, and behenate. (v. Behenie acid).

BENIC ACID v. BEDENIC ACID. BENYLENE C, H = (223°-228°).

4. 147, 184. BENEAUIN C.H., N.O. [150°]. A neutral crystalline substance obtained by extracting

with alcohol the product of the action of ZuEt, on phenyl-acctonitrile (Frankland a. Tompkins, C. J. 37, 569).

BENZAL v. BENZYLIDENE.

BENZALDEHYDE e. BENZOIC ALBEHYDE.

BENZALDOXIM C.H. CH:N.OH. Oxim of benzoic aldehyde. (c. 220° with decomposition). Colourless oil, formed by the action of hydroxyl. amine on benzoic aldehyde (Petraczek, B. 15, 2735). Formed also by reducing benzamidoxim with sodium amalgam (Tiemann a. Kriiger, B. 17, 1692). By heating with HCl it is split up into hydroxylnmino and benzaldehyds. Ac.O converts it into benzonitrile (Lach, B. 17, 1571).

Sodium salt C.H.: N(ONa) nq : white oasily aduble plates, formed by the action of sodium ethylate on benzaldoxim in alcoholic solution; gives characteristic pps. with the sults of the heavy metals.

Hydrochloride C,H.: N(OH), HCl; white glistening scales, rotates on water.

Methyl other C.H.: N(OMe): (191° uncorr.); colourless oil, lighter than water and slightly soluble; formed by the action of methyl iodide and sodium ethylate on benzaldoxim; by HCl it is split up into bonzaldeliydo and methyl-hydroxylamine.

Ethylether C.11 .: N(OLt): (208° uncorr.); colourless oil, split up by HCl into ethyl-

hydroxylumine and henzaldehyde. Propyl ether C,H,:N(OC,H,): (225° un-

ecrr.), colourless oil. Iso-butyl ether C,Ha:N(OC,Ha): (238°

uncorr.), colourless oil. Amyl ether C.II,:N(OC,H,1): (161° un.

corr.), colourless oil (Petraczek, B. 16, 823). BENZAMIDE C.II, NO i.e. C.H., CO.NII, Amide of benzoic acid. Mol. w. 121. [130] (Ciamician n. Magnaghi, B. 18, 1828). S.G. 21-34 (Schröder, B. 12, 1612).

Formation.—I. From BzCl and NII,—2. Together with NII,0Bz, by the action of Bz,0 on NII, -3. From EtOBz and NII, -4. By boiling hippuric acid with water and PbO, or by heating hippuric acid in a current of dry HCl.

Properties.—Monoclinic tables: a:b:c= 228:1:1068; β=89° 22' (Klein, A. 166, 184); v. sl. sol. cold water, m. sol. hot water, especially if it contain NH2; v. sol. alcohol and ether.

Reactions .- I. It splits up into water and benzonitrile when heated with dehydrating agents (P2Os, P2Ss, or H SOs) and to some extent when heated alone at 290° .- 2. Boiling aqueous polash forms KOBz. -3. Boiling acids form benzoic acid.-4. BzCl or Bz,O form benzomtrile and benzoic acid. - 5. Reduced in acid solution by sodium-amalgam to benzyl alcohol, -6. Boiling phenol gives benzoyl-phenol (PhOBz) and NII, - 7. PCl, forms an unstable substance which rapidly splits up into HCl and benzonitrile (Wallach, A. 184, 19).—8. COCl, gives beuzo-

mitrile, cyaphenine, and di-benzoyl-drea (Schmidt, J. pr. [2] 5, 35).—9. CSCl, gives benzenitrile, COS, and HCl (Rathke a. Schäfer, A. 169, 107).— 10. With chloral it combines forming C.H.Cl.NO 343.41 3. 31

bromine in alkaline bolusion into saille (Hofmann, B. 18, 2787).—13. A solution contamide in bromine deposits crystals of

anstable BzNH Br. Combinations. BzNH HCl: long prisms formed by saturating a mixture of bonzamid formed by saturating a mixture of benzaming and HClAq with HCl (Dessaignes, A. Ch. [8 34, 146; Pinner a. Klein, B. 10, 1897). Whet exposed to the air it gives off all its HCl.—BzNH, 3HCl. [178°] (E. v. Meyer, J. pr. [2] 30, 122).

Salta-(BaNII),Hg [2240]. Formed by boiling benzamide with water and HgO. Lamina (from alcohol); v. sol. alcohol and ether .-BzNHTl: slonder neodles (Church a. Crookes, C. J. 17, 151).

Additional References .- Liebig a. Wöhler. A. 3, 268; Fehling, A. 28, 48; Schwarz, A. 75, 195; Laurent, Revue Scient. 16, 391; Henry, Z. [2] 5, 446; Brauns, Ar. Ph. [2] 126, 214; Oppenheim a. Czarnemsky, 6, 1392; Guaresohi, G. 4, 465; A. 171, 141; Kekulé, B. 6, 113; Sohiff a. Tassinari, B. 10, 1785; Friedburg, A. 158, 26.

Bons-chloro-amide C.H., CO.NHCL. Prepared by gradually adding a cone, solution of chloride of lime to a cold cone solution of benzamide acidified with AcOII, the product being shaken out by ether as it is formed. Long colourless prisms (from water) (Bendsr, B. 19, 2274).

Dibenzamide C, H, NO, i.e. NIBz, [148°]. S. 12 at 15°. Formed, together with benzamids, by the action of KNII, on BzCl dissolved in ether (Banmert a. Landolt, A. 111, I); and from benzonitrile (10 g.) and fuming 11,80, (7 g.) (Barth a. Amoler, B. 9, 975; Gumpert, J. pr. [2] 30, 67). Trimetrio crystals, a:b:c= 931:1:1 969. Sl. sol. boiling water, v. sol. alcohol, ether, and benzene.

Salts. - NaNBz, 2aq: small prisms, sol. ether. - AgNBz

BENZAMIDINE C, H, N, i.e. C, H, C(NH).NH, [75%-80%].

Preparation .- Benzonitrile is converted by treatment with isobutyl alcohol and IICl into the hydrochloride of Call, C(N11), OC, II, whonce ammonia produces benzamidino (Piunor a. Klein,

B. 10, 1880; 11, 4).
Properties.— M. sol. water, sl. sol. ether, v. c. sol. alcohol; deliquescent; very volatile. De-composed by heat into NH, and cymphenine.

Salts. Bilcl: flat needles. B', H, PtCl, AgC H.N...

BENZAMIDO. v. Benzoyl-Amido. BENZAMIDO-ACETIC ACID v. Hippung

BENZ AMIDOVIM C.H.N.O i.e.

PhC(NIL),NOII. Benzenyl-oramidine. Benzenyl-amidorim. I.omitroso-benzylamine. [807].
Formation.—1. By the action of hydroxyl-

amine on an alcoholic solution of benzonitrile (Tiemann, B. 17, 128).-2. By the action of hydroxylamine on the hydrochloride of bense amidine.—3. As a by product in the action of hydroxylamine hydrochloride upon benzimide. ethyl-ether.—4. By digesting thio benzamide with an alcoholic solution of hydroxylamine [151] (Wallach, B. 5, 251). -11. With ctivil (Tiennann, B. 19, 1608). Long flat monosymmetric it forms N., benzoic ether, and H.O metrical prisms a:bic -2:502;1:1077. Volatilises

indecomposed. V. sol. alcohol, ether, bensene, and hos water, al. sol. cold water. It is poisonous. It dissolves both in seids and alkalis. Fe.Cl. gives a red colouration. The ammo-Fe.Cl. gives a red colouration. The ammodniacal solution gives white crystalline pps. with BaCl, AgNO, Pb(OAc), and ZnSO. The silver pp. on heating in the solution in which it is formed gives a splendid silver mirror.

Reactions .- 1. Gives the carbnmine reaction with chloroform and alcoholic potash .-2. Nitrous acid forms benzumide. - 3. Sodium amaigam reduces it to benzaldoxim and NH, (Tiennan a. Nägeli, B. 18, 1086).—4. When quickly heated at 170° it splits up into benzonitrile and NII, 5. By heating with actic anhydride it yields benzenyl-azoxim-ethenyl $C_{\bullet}\Pi_{\bullet}C \stackrel{NO}{<}_{N} > C.CH_{\bullet}$

Salts. A'Nn: white crystalline solid, decomposed by water. A'K: crystals. A'Ag: unstable white crystalline pp. A'(CuOH): amorphous dark green pp. -A'H, HCl: large that plates or concentric needles. -A'II,ILSO,: large prisms.—(A'II)_zII_oSO_z: maorphous solid.

Mothyl ether.—A'Me: [57°]; (230° mi-

corr.); prisms; v. sol. alcohol, ether, and benzenc, sl. sol, water. By HCl and NaNO, it is ennverted into benzenyl methoxim chlorido

Call, CCI(NOMe).

Ethyl cthar. - A'Et: [67]; trinietric plates. By ditute H.SO, and sodium nitrite it is converted into benzhydroximie ethyl . other (ben-oyl hydroxytamine ethyl ether) C, II, C(OII) NOEt. With IICI and NaNO, it yields benzenyl-ethoxim-chloride.

Benzyl other. A'C,H,: [91°]; scales.

Benzoyl derivative.
C.H. C(NII):NOBz: [140"]; slever white needles; v. sel. nlcohol, ether, and nqueons acids, insol, water. On heating it readily splits off H₂O, giving rise to benzenyl-nzoxim-benzenyl $C_{\epsilon}H_{s}$, $C < \frac{N.O}{N} > C.C_{\epsilon}H_{s}$.

Acetyl derivative Call, C(NH,):NOAc. [90°]; thin plates or that prisms, sol. alcohol, sl. sol. ether, v. sl. sol. water. By boiling with water it loses ILO and is converted into benzenyl-nzoxim ethenyl.

Butyryl derivative

O.H. C.N.II.):NO.CO.C.II., [91°]: fine needles.

Ethylane ether (Ph.4:(NII):NO).C.H.,
[156°]. White plates. Sol. alcohol, ether, benzene, and ligroon, insol, water. Formed by heating an alcoholic solution of benz-umidoxim (2 mols.) and said in cthylate (2 mols.) with ethylene bronnide (1 mol.).

Gompound with chloral C,H,N,Cl,O, [135] Obtained by mixing the constituents. White crystalline powder. V. sol. alcohol and ether, insol, water. By treatment with H.SO, or by long boiling with water it is resolved into its components (Falck, B. 19, 1485).

References .- Pinner, B. 17, 184; Tiemann a. Krüger, B. 17, 1685; 18, 731, 1053; Tiemann, B. 19, 1479, 1668; Schulz, B. 18, 1080; Falck, B. 19, 1484 (c. also Azoxins).

Benz amidexim carbonic other

C.H. C(NII.):N.O.CO,Et. [127°]. Formed by the action of chloroformic other apon benz-amidoxim (Falck, B. 18, 2407). Long glisten-ing needles. V. sol. alcohol, ether and benzene, less in lignoin. On heating it splits off alcohol, forming benzenyl-azoxim-carbinol-C.H.O NO COH).

Carbonyl-di-benz-amidoxim C13H14O2N4, 6.4. C.H., C(NH.):NO CO. [129°]. Formed by the action of carbonyl chloride upon benz-amidoxim dissolved in benzene (Falck, B. 18, 2470). White plates. Sol. alcohol and ether, v. sl. sol. benzene, insol. water.

BENZ-AMIDOXIM-m-CARBOXYLIC ACID C.H.,N.O. i.e. [3:1] C.H.(CO.H).C(NOH)NH., [200°]. Crystalline solid. Sol. hot water and alcohol, sl. sol. ether, nearly insol. obloroform and benzenc.

Formation .- 1. By saponification of the ethyl other which is obtained by combination of m cyano benzoic other with hydroxylamine.-2. By digesting a mixture of equivalent quantities of m-cyano-bonzoic acid, hydroxylamine hydrochlorido, und sodium-earbonate, in dilute alcoholic selution for 12 hours at 80°-100°.

Reactions. - The aqueous solution of the munonium salt gives sparingly pps. with CuSO, Pb(OAc)., AgNO, and ZnSO, Heated with neetic auhydride it is converted into m-carboxy. benzenyl-nzoxim-ethenyl

 $C_{\bullet}H_{\bullet}(CO_{\bullet}H).C \leqslant N > C.CH_{\bullet}.$

Ethyl ether A'Et [118°]; needles; v.sol. alcohol, sl. sol. water (Müller, B. 19, 1455).

Beaz-nmidoxim-p-carboxylic acid [4:1] C.H.(CO.H).C(NOH)NH. [above 330°]. Formed by digesting p-cyanobenzaic acid (1 mot.) hydroxylamine hydrochloride (1 mol.), nnd sodium carbonato (1 mol.) in dilute alcoholio solution for 18 hours. Sol. dilute alcohol, sl sol. water, nearly insol. absol. alcohol, ether, and benzene. A dilute aqueous solution of the ammonium salt gives pps. with CuSO, and with AgNO₃. By boiling with acetic nuhydride it is converted into p-carboxy-benzenyl-nzoxim-ethenyl $C_a \Pi_i(CO_a\Pi).C \leqslant NO_N > C.CH_a$ (Müller, B. 19,

Ethyl ether A'Et:[135°]; obtained by licating the othyl ether of p-cyanobenzoic acid with hydroxylamine in alcoholic solution (Müller, B. 18, 2185). Colourless crystals; sol. boiling wnter.

BENZAM-MALONIC ACID v. CARBOXY-PHENYL-MALONAMIO ACID

BENZAM-OXALIC ACID v. CARBOXY-PHENYL-OXAMIC ACID

BENZAM-SEBACIC ACID v. CARBOXY-PHENTL-SERACAMIC ACID.

BENZAM-SUCCINIC ACID v. CARROXY-PHENTI.-SUCCINAMIC ACID.

BENZ-ANHYDRO. v. Benzentl- or as derivatives of Benzamidine.

BENZARSEN- v. ARSENIC, ORGANIC DERIVA-TIVES OF.

BENZARSENIC ACID v. ARSENIC, ORGANIC DERIVATIVES OF.

BENZ-BROMO-QUINOLINE v. (B.)-Bromo-QUINOLINE.

BEAZ-CHLORO-AMIDE v. BENZAMIDE. BANZ-CHLORO-QUINOLINE v. (B.)-CHLORO-QUINDLINE.

REEL DERATES

Bens. (a) Swelly L. Grossymbia.

NH, C(NH), NMc.C, H., OO, H. From bensglycocyamine, cono. ROH, methyl alcohol, and
MeI (Griess, B. 8, 324). Narrow lamine (consiming 11/20) all and reference of the concyamine of the control o taining 1 aq), sl. sol. water and alcohol. De-composed by baryla into urea and methylamido-benzole acid. - B'HClaq. -B',H,PtCl, 2aq.

Bens-(8)-methyl-glycocyamiae NHMe.C(NH).NH.C.H.,CO.H. From ethoxy-carbimidamido benzoic acid' and cold conc. NMeH, solution. Lumine, v. sl. sol. cold water. Boiling baryta forms methyl-urea and amidobenzoic acid. -B'HCl.-B', P.PtCl. 2aq. BENZCREATININE

o-Benz (a)-methyl-glycocyamidiae C,H,N,O

N(CII,) = C, H, . Prepared by the NH-- co action of McI on o-benzglycocyamidine (Griess,

B. 13, 978). White needles. Sol. alcohol, sl. sol, ether and hot water. Insol. caustic alkalis. Salts. - B'HClay: soluble plates. (B'HCl) PtCl,

o-Benz-(B)-methyl-glycocyamidine

C₀H₀N₁O i.e. IIN = C NII = C H.

NICH₁)=CO

Prepared by the action of methylamine on 'etlioxy - cyan-amido - benzoyl' (C, H, N, O,) B. 13, 978). White needles. Sol. lkalis. Weak base. (Grices, caustic ulkalis.

Salts.-BHCl: small tables or prisms, decomposed by water. -(B'HCl), PtCl4: sparingly soluble vellow plates.
BENZCYANIDINE v. BENZOYL CYANIDE.

BENZEINS. These hodies, which much resemble the phthalems, are hydroxylated aromatic earbinols, such as di-oxy-tri-phenyl carbinol C.H. C(C.H.OH), (OH). Prepared by heating phenois with benzotrichloride. The compound from resorein is yellow, those from cresois, pyrocatechin, hydroquinone, orcin, and (B). naphthol are yellow or yellowish-red, pyrogallol gives a blue, and (a)-naphthol a green dye. On reduction they give the corresponding derivatives of methano (Doebner, B. 13, 610; A. 217, 227).

BENZENE C.H. Mol. w. 78, [6°]. (80·1°) (R. Schiff, A. 220, 91). S.G. 163 8839 (S.); 30 8799 (Brühl). V.D. 274 (calc. 270) (S.). S.H. 3834 + 0010 13t at to (R. Schiff, A. 231, 320). H.C.p. (liquid) 776,000 (Berthelot, A. Ch. [6] 23, 193); 779,530 (Stohmann, Rodatz a. Herzberg, J. pr. [2] 33, 25*); (gaseons) 799,350 at 18° (Th.); 787,488 (S. R. a. H.). H.F.p. (asvapour) -12,510 (Thomsen, Th. 4,61); -13,670 (Th.). C.E. (14-2 to 80-1) 001:05 H.F.v. (S.). S.V. 95 91 (S.); 95 8 (Ramsay, C. J. 357 463). μ_s 1·5134 (B.); μ_o 1·5062 at 14 (Negreano, C. R. 104, 423); 1·5050 iGladstone, C. J. [2] 7, 101). R 2 42 16 (B.); 44 02 (Kanonnikoff, J. pr. [3] 31, 352). Dielectric constant 2 292 at 14° (N.).

Formation .- 1. By heating benzoic acid with lime (Mitscherlich, A. 9, 39; Peligot, A. 9, 48, 257; 11, 277; 12, 39), or by passing the vapour of benzoio acid over red hot iron (Darcet, A. Ch. [2] 66, 99).—2. By heating phthalic acid with lime (Marignac, A. 42, 217).—3. By dry Matillation of quinic acid (Wohler, A. 51, 146). You L

4. By passing oil of bergames over red-hos line (Ohms, A. 31, 318).—5. Together with naphthalene, styrene, retene, do., by heating acctylane to a red heat (Berthelot, A. Ch. [4] 0, 460). Benzene is among the products obtained by passing tolueno, xylene, 4-cumene, or styrene through a red-hot tube. -6. Metallio succinates give on dry distillation a liquid (hydroquiuone dihydride?) which yields benzene when distilled with zine-dust (v. Richter, J. pr. [2] 20, 206),-7. From benzene sulphonic acid by passing steam through its solution in diluted H.SO, at 175° (Armstrong a. Miller, C. J. 15, 118), -8, From phenol (70 g.) by distilling with P.S. The yield is small $(16 \text{ g.}): 8C_s 1 | 011 + P_s S_s = 2(C_s H_s)_1 | O_s + 3H_s S + 2C_s H_s$ (A. Genther, A, 221, 55). 9. The homologues of benzene when treated at their boiling points with a current of HCl gas in presence of Al₂Cl₈ evolve McCl and yield lower homologues, but at the same time part of the McCl attacks other portions of the hydrecurbon with production of higher homologues. Thus it HCl is passed into boiling mxylene containing Al₄Cl, bouzene, toluene, psendocumene, mesitylene, and durene are obtained (Jacobsen, B. 18, 338; Anschütz a. Immenderif, B. 18, 657). - 10. Benzene is produced when benzylideno chloride, PhCHCl, or benzotrichloride, PhCCl, is heated with sodalime (Limpricht, A. 159, 303). -11. From diazobenzene nitrate or sulphate by heating with ulcoliol or alcoholic KOIL

Preparation .- 1. Coal tar is distilled and the fraction boiling below 150° is freed from phenois by shaking with NaOHAq and from lasses by shaking with H.SO. It is then rocti-It is convenient to keep the still-hand at 100°; toly as is then condensed while benzene pa ses over (Faraday, T. 1825, 440; Hofmann a Mansfield, C. J. 1, 2(1). On the large scale the distribution is performed in an i.on boiler, to which is attached a tall vertical column divided into compartments; the fire is regulated so that pure benzene passes over, while its homologues condense and run buck to the boiler. Benzene is completely freed from its homologues by cooling with ice and salt, when it solidifies and the homologues which remain fluid may be pressed out. It is freed from thiophene by shaking with cone. H280c 2. A mixture of benzoic acid (1 pt.) and slaked lime (3 pts.) is distilled from an iron tube: the product is slaken with KOHAq, distilled with steam, dried with CaCl₂ and Na und rectified. Diphenyl and benzophenous ere by products. The benzoic as id used must not be prepared from tolucie, otherwise it will contain thiophenic acid, and the benzene will contain thiophene.

Properties. - Colourless, mabile, strongly retracting liquid. Volatile with steam. Scarcoly soluble in water, v. c. sol. ulcohol, ether, glacial acctic acid, acctone, and chloroform. Crystallises in trimetrio pyramids, a:b:c = 891:1: 779 (Groth, Z. [2] 6, 553). It dissolves S, P, I, fats, resins, oils, and many alkaloids. It burns with a luminous flame; Ig. burnt with excess of hydrogen giving out a light equal to 58g, of spermaceti. When burnt with CO, and CH, the light equals that of 6-1g., and 7-8g., of spermaceti respectively (Frankland a. Thorna, C. J. 33, 93). There are four bands in the

Chartey, C. J. 39, 162; 47, 594).

Detection.—Benzene is converted by fuming HNO, into nitro-benzene; this is washed with

HNO, into nitro-benzene; this is washed with water and reduced by tin and HCI to aniline; sanstic soda is added and the aniline extracted with ether; the ether ie evaporated and the aniline dissolved in much water; the aqueous solution gives a violet colour with bleachingpowder. A mixture of HNO, and H.SO, forms

anime dissolved in much water; the aqueous solution gives a violet colour with bleaching-powder. A mixture of HNO, and H.SO, forms di-nitro-benzeno, which, after crystallising from dilute alcohol, mells nt 89°.

Detection of Thiophene in Benzene.—Thiophene, which is usually present in small quanti-

tiss in commercial benzene, ie indicated by the blue colour produced by shaking with conc. H₂SO₄ and isatin (V. Meyer, B. 16, 1465; Baeyer, B. 12, 1309).

Impurities.—Crude benzene may contain traces of toluene, xylene, thiophene, CS₂, amyleno, crotonylene, alcohol, and ucctonitrile.

Reactions.—1. Benzene when passed through a rea hot tube forms hydrogen, a little acetylene, diphonyl, benzerythrene C₂, H₁₀, p-di-phenyl-benzene, iso-di-phenyl-benzene, and triphenyl-ene (Berthelet, Bl. §2 6, 272, 273; G. Schultz, A. 174, 201; H. Schmidt a. G. Schultz, A. 203, 118).—2. A mixture of benzene vapour and

ethylene passed through a red-hot tube gives

diphenyl and small quantities of anthracene, styrene, and phenanthrene (Berthelot, Bl. [2] 7, 113, 274; Ferko, B. 20, 660).—3. A mixture of equivalents of benzene and toluene dropped at the rate of 80 g. per hour from n tap finnel into the turned-up end of an iron tube kept in low red hent is converted, to the extent of about 10 per cent., into gases, naphthalone, diphenyl, p.tolyl-di-phenyl, o.p.di-tho, y.h. and (3) diphenyleu-methane, phenanthrene, anthracene, p.di-phenyl-benzene, a hydrocarbon C. H. a

p.di-phenyl-benzene, a hydrocarbon $C_{21}I_{18}$, a hydrocarbon (13°) (293° 316°) and two liquid hydrocarbons (359° 383°) and (404° 427°) (Carcolley, C. J. 37, 701).—4. Induction sparks passing through liquid benzene produce a gas that contains 42° pc. acetylene and 57° p.o. hydrogen (Hestrem, Bl. 12) 42, 267).—5. Aluminium chloride (c. p. 147) acting upon a mixture of bonzene and an alkyl chloride causes 11Cl to escapo with the resulting formation of nn alkyl-bonzene (l'cicelo In. Crafts, C. R. 84, 1392, 1450; 85, 74; A. Ch. [6] 1, 449). (a) Thus methyl chloride passed into a mixture of bonzene and Al₂Ol₈ gives (oluene, c., m., and p.xylone, y-cunene, mosityleno, durene, isodurene, pentamethyl-benzene, and hexa-methyl-benzene (Ador

a. Rilliet, B. 12, 329; Jacobsen, B. 14, 2621).

(b) Chivral is converted by benzene in presence of Al,Cl_e into CPbCl_eCH(OH)Cl, the hydro-chloride of aa.-di-chloro-phenyl-neetio aldehyde (Combes, C. R. 98, 678; Bl. [2] 41, 382).

(c) Methylene chloride in presence of Al,Cl_e gives di-phenyl-methanne, anthracene, and toluene (Friedel a. Crafts, Bl. (2] 41, 322).

(d) Chloropicrin, Al,Cl_e, and benzene form CHPh, and Ph,COH (Elbs, B. 16, 1274). (e) Allui

Chloropierin, Al, Cip, and benzene torin Chiri, and Ph. COH (Elbs, B. 16, 1274). (c) Allyi chloride, Al, Cle, and benzene give di-phenyl-propane and n-propyl-benzene (Vispok a. Zuber, A. 218, 374). (f) Vinyl bromide, Al, Cle, and benzene give etilyl-benzene, u-di-phenyl-ethane, and di-mothyl-authracons dihydride (Angeblie a. Anschütz, B. 17, 167). Vinyl tribromide,

McLoughlin, B. 20, 195). (j) Acetylene in pu scnce of Al₂Cl₄ forms styrene, di-phonyl-ethan and di-tolyle (Varet a. Vienne, Bl. (2) 47, 917). 6. Heated with Al₂Cl₄ (2 pts.) at 200° in seelat tubes benzene givee tolucne, ethyl-benzene, aldiphenyl (Friedel a. Crafte, C. R. 100, 692). 7. Phenol is among the products of oxidatic of benzene by H₁O₂ (Leeds, Ph. [3] 11, 1066 cf. Kingzett, C. N. 44, 229). Phenol is all formed when benzene is digested for some data 40° with supress blevide and divergent.

at 40° with cuprous obloride and dilute HC atmospheric oxygen attacking benzene an Cu₂Cl₂ simultaneously (Noncki a. Siober, J. p. [2] 26, 25). In the animal body it is exidee to hydroquinone and pyrocatechin (Nencki Giacosn, H. 4, 325; cf. Schultzen a. Naunyı C. C. 1867, 705). Oxidation with MnO₂ an dilute H₂SO₂ produces formic, benzoic, an phthalic acids (Cariue, Z. 4, 505; A. 141

dilute H₂SO, produces formic, benzoio, an phthalic acids (Cariue, Z. 4, 505; A. 14: 50). The formation of benzoic acid is per haps preceded by that of diphenyl (Kekulé PbO, and H₂SO, give benzoic acid; PbO, an boiling dilute HNO, give only oxalic acid; CrC gives only CO₂ (Holder, Am. 7, 114).—8. PCl₂ a a red heat forms PhPCl₂, diphenyl, and 1 (Michaelis, A. 181, 265; Köhler, B. 18, 1623).—

9. S.C., at 250° forms chloro benzene, HOl, and S (Schmidt, B. 11, 1168).—10. Italia acid and H.S.O. on heating slowly form iodo-benzene (Peltzer, A. 136, 194).—11. SO,Cl, at 150° give chloro-benzene (Dubois, Z. [2] 2, 705).—12 CLSO,COII forms PhSO-Rh, PhSO-Cl, and PhSO,II (Knapp, Z. [2] 5, 41).—13. 11ClO forms C, H.,(OII),Cl, the trichler hydrin of phenose (Carius, A. 136, 323).—14. Aqueons HClO, form trichloro-phenomalio acid, chloro-benzene, and dichloro-quinone (Cariue, A. 142, 123).—15. CrO,Cl, acting upon benzene diluted with HOAc (1 vol.) gives trichloro-quinone (Carstanjen, J. pr. 107, 331). When benzene is heated with CrO,Cl, there is formed a brown pp. of

C.H.(CrO.Cl), which is converted into quinone by water (Etard, A. Ch. [5] 22, 269).—16. Condenses with sulphuric acid and aldehydes, XCHO to XCHPh₂. Thus chloral forms COL.CHPh₂; bromal forms CBr,CHPh₂; chloro-aldehyde forms CIL,Cl.CHPh₂; fermic aldehyde forms CH,Ph₂ (Goldschmiedt, B. 6, 986; Hepp, B. 6, 1489).—17. Benzene is not attacked by HIAq and P at 250°, but at 280° it gives hexahydro-benzene (Wreden a. Znatowicz, A. 187, 163; cf. Berthelot, A. Ch. [3] 15, 150).—18. When chloring

is passed into benzene containing thiophens Hol is evolved and the benzene then no longer gives the indophenine reaction (Willgerodt, pr. [2] 33, 480). Pure benzene is not attacked by chlorine in the cold and in the dark, but as 80° or in smallght benzene hexachloride is formed. In presence of carriers, i.e. substances capable of combining with oblorine in more than one proportion, chloro-benzenes are produced. 18. Nitric acid forme nitro- and di-nitro-benzenes.

Be B gives C.Br. (Gustavaon, B. 11, 2151).— (C.H.), AlBr.—(C.H.), SabCl.; monoclinic tables, corned by warming SbCl, with bensene (Watson Smith a. Davis, C. J. 41, 411).

Petassium-benzene C.H.K mixed C.H.E. Formed by heating benzene with K at 250° (Abeljaux, B. 5, 1027; 9, 10). Blue-bleck erystalline mass, insol. benzene. Takes fire in erystalline mass, insol. benzene. sir. Converted by water into di-phenyl-benz-ene, hydrogen, and di-phenyl.

Constitution of Benzew. - That the molecular formula of benzeno is Call, and not any multiple or submultiple of this, is settled, not only by its vapour density, but also by the existence of the following series: C,H,Cl, C,H,Cl, C,H,Cl,, C,H,Cl,, C,H,Cl,, C,HCl,, C,Cl,. We may call the siz atoms of hydrogen in the molecule of benzeno, n, b, c, d, c, f. The first question is: are these six atoms of equal value, or could we, by displacing a by an element or radicle li, get a product different from that which would be produced by displacing b by R?

Proposition I .- Four, at least, of the hydrogen

atoms are of equal value.

Ordinary phenol contains hydroxyl in place of one hydrogen; call this hydrogen a. Beomine and phosphorus convert phenol into C.H.Br. Sodium and carbonic ucid convert this bromophenol inte sodie benzoate, C.H., CO.Na. Hence the carboxyl of benzoic acid has taken the place of the hydrogen atom n. Now, there exist three oxy-benzoic acids, C. II, (OII) (CO.II), and since in these the carboxyl is in position a, the three hydroxyls must have displaced three other atoms of hydrogen, say b, c, and d. When distilled with lime, these three acids, instead of giving three phenols, the hydrozyl being in places b, c, d, give the same phenol which is identical with the original phenol. Hence, the four hydrogen atome which we have called n, b, c, and d, are of equal value (Ladenburg, B. 7, 1681).

Proposition 11. - To every hydrogen atom in the molecule of benzene there are two pairs of hydrogen atoms similarly related. Benzoic acid,

C.H.(CO.H) gives bromo-benzoic acid, which

ws may call C. II, Br(CO, II). This, when seted upon by nitric acid, produces two isomeric nitrobromo benzoic acids. We may call these

But by reduction these lose their bromine, and give rise to amido benzoic acids :

C.H.(NH.)(CO.H), C.H.(NH.)(CO.H). are found to be identical, being anthranilic acid.

Hence, b and f are symmetrically related with regard to a (Hubner a. Petermann, A. 149, 129).
Again, ordinary nitro-benzoio acid may be converted into the above bromo-benzoic acid by the

diazo- reaction, hence it is C.H.(NO,)(CO,H). On nitration it gives a di-nitro-benzoio acid which we may call C.H.(NO.)(NO.)(CO.H), which may be reduced to C.H.(NO.)(NH.)(CO.H), CH,(NO)O(CO,H), CH,(NH)O(CO,H), and

O.H.Cl(CO.H). The last acid is found to be identical with the chloro-benzolo acid

C.H.Cl(CO.II) obtained by the diazo- reaction

from C.H.(NO./(CO.H). Hence c and s are similarly related with regard to a. Therefore we have a second pair of hydrogen atoms similarly related with regard to a (Hübner, A. 222, 94, cf. Wroblewsky, A. 192, 206).
Proposition III.—The six atoms of hydrogen

in the molecule of benzene are of equal value. Since a, b, c, and d are of equal value, and the situations of f end e are similar to those of b and c respectively, all six atoms of hydrogen ure similarly placed and of equal value. conclusion might also be deduced from the fact that no instance of isomerism among the monosubstitution products of benzene has been

Isomerism among di-substitution products. Since two pairs of hydrogen atoms are symmetrical to any fifth, it follows, that only three diderivatives of benzene can exist with a given formula. Using our former notation, these are nb = af, nc = ac, and ad.

This is confirmed by experiment.

Structural formula.-Since the atom of carbon is assumed to be tetravalent, all the hydrogen atoms of benzene cannot be attached to the same atom of carbon, and symmetry requires that they must be either each attached to one carbon, or else three must be attached to one, and three to another; or, finally, two must be attached so one carbon, two to another, and the rem ming two to a third. The fwe latter hypotheses do not account for more than two di- substitution products; hence the former is established. The earbon atoms must be united amongst themselves in a symmetrical fashlon. Each atem of carbon must be united with at least two other atoms, or the group would not hold together; but it may be united with three other ctoms. The former hypothesis results in the formula:

This is the ring-formula of Kekule, which is one of the two formula originally put forward by him (A. 137, 160). If we number the posttions occupied by the atoms of hydrogen thus,

we see that the formula shows the possible existence of four di-substitution products—vis. 1:4, 1:3, 1:2 and 1:6. To get over this difficulty, Kskulé resorts to a peculiar mechanical hypothesis. Hs supposes that what ws rspresent by straight lines in a formula really indicates that two atoms vibrate with reference to sach other so that the above formula would mean that, in a given unit of tims, 1 approaches 6 twice as often as it approaches 2; and so for the other atoms. Now, if this were the ease, the di-substitution product 1:2 would differ from 1:6; but he assumes that the motions of 1 are as follows: first, it approaches 6 twice; then it approaches 2 oncs; next it approaches 6 once; then it appronches 2 twice; then 6 twice; 2 once; and so on. This is equivalent to saying that the above formula for benzene is true for one instant, after which it changes to

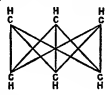


and the next instant it changes back again, and so on. This assumption leads to the deduction that only three di-derivatives can exiet, and, if we could dovise no other formula for benzene, we should be obliged to accept it. As a matter of fact, it is now almost universally adopted; not so much on its intrinsic merits, as on account of the enormous service where it has rendered to chemistry.

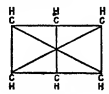
There remains, however, a second hypothesis possible, which is that every stom of carbon is united to three other atoms. The following mechanical construction may help to elucidate this hypothesis.

Let three rods be driven into the ground at the angles of an equilateral triangle, and let the top of each rod be joined by elastic string with the bottom of each of the two adjacent rods. A figure somewhat resembling a coronet is obtained, and we may suppose the six atoms of earbon in the benzeno molecule situated at the two extremities of each of the three reds. At first sight it might appear that this reprecentation of the benzene melecule would indicate the existence of three di-derivatives-namely, (1) when the substituted hydrogons are attached to two carbon atoms at opposite ende of one rod; (2) when they are attached to earbon atoms which are both on the upper or both on the lower ends of two different rods; (3) when one carbon atom is on the upper end of one rod and the other is on the lower end of another rod. But if we assume that formulæ and other mechanical symbols represent net actual position in apace, but merely modes of combination of atoms, especially showing which atoms are directly and which indirectly united, (3) is identical with (1), for it can be converted into (1) by simply bolding the string, at the opposite ends of which the carbou atoms have been placed, upright, and doing the same with the two corre-

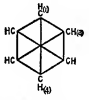
sponding range. The rods will now take positions formerly occupied by the strings now held upright, and the figure will be the same as before. Hence this figure for benzene gives only two di-dsrivations, and accordingly it must be discarded. When the figure we have just considered is projected on a plans it assumes the form:



This figure, by simply twisting the osntral rod, is converted into



and if the central rod be new elongated we get:

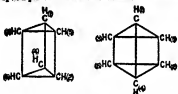


This symbol, which has been a favorite with some chemists, must, of course, be abandened along with the selid figure from which it is derived; but it is also very easy to see that 1:2 and 1:4 di-derivatives, are identical, aince if we pick up the carbon atom (4) and place it upon (2), and then take up (2) and place it where (4) was, supposing all the while that the connections, which ws may imagine to be elastic, are not broken, tha figure will be whelly unaltered.

Thers remains one other benzene formula: it is obtained by joining the snds of the three rods placed vertically by six strings as before, but with this difference, that whereas in the previous formula the top of one rod is joined to the bottom of the others, in this formula the top of each rod is joined with the top of each of the others, and the bottom of each rod is joined with the bottom of each of the others. We thus obtain a right-angled prism on a triangular base. This formula, defended by Ladenburg (Theorie der aromatischen Verbindungen, Brunswick, 1876), is capable of explaining most of the reactions of benzene, and the objections that have been brought against it are shieffer the

secult of misconception of its nature. Projected apon a plane this formula becomes:

3 33.14



If the upper triangle be rotated through 180°, and then the figure be projected upon a horisontal plane, we obtain a figure which resembles a star:



The two former figures are not symmetrical, but the latter is clearly so, and it has this advantage over the prism formula, that, if the atoms of carbon are numbered consecutively, they correspond to the atoms of carbon in Kekulé's formula, also numbered enneceutively; whereas this would not be the case with the second of the three formula here given, which is that used by Ladenburg.

For most purposes it will not be necessary

For most purposes it will not be necessary to decide which formula we adopt, for both the star-formula and the formula of Kekulé:

may be represented by the simple hexagon:

The numbering of the carbon atome here given is used throughout this diotionary. Thus, the expression C.H.Br., [1:5] must be taken to mean that one bromine atom has displaced the hydrogen atom numbered (1) and the other that hydrogen atom numbered (5).

hydrogen atom unmbered (5).

Physicists have tried to decide between the formule of Kekulé and Ladenburg. Thomsen

(Th. iv. 273; A. 205, 123) considers that thermachemical data favour Ladenburg's formula, but the assumptions he makes in the course of his argumant lead him in other cases to impossible conclusions. The specific volume of benzene is H; whereas that calculated on the assumption that the S.V. of C-11, and that of H-5:5 is 99: this would nearly show that the relation between the carbon atoms in the benzene molecule is different from that in saturated paraflins. If we compare the specific volumes of hexane, diallyl, and benzene, we find that:

Hexane, C.H., has a S.V. 140 0 Diallyl, C.H., "125-7 Benzene, C.H. "95-9

Benzene, C.H., ", 95.9.

We see that the difference between the first and second (14.3) is less than that between the second and third (29.8) and that when hexana is converted into benzene by the removal of 811 the S.V. is lowered by 8 × 5.5, while tha removal of 411, in converting bexame into diallyl, lowers S.V. by only 4 × 3.6. That is to say the want of saturation of diallyl is accompanied by an unusually large specific vulume, whereas this is not observed in the case of benzene. This would indicate that henzen is not unsaturated in the ordinary sense, and can be used as an argument in favour of Ladenburg's formula for benzene (Lossen, A. 214, 129; It. Schiff, A. 220, 303),

On the other hand, the S.V. of hexalydrotolene (147-2) differs from that of tolene (148) 23-8. This is about three times the difference (7-2) between the S.V. of pentane (117-2) and amplene (110). Hence it would appear that the change in the state of saturation in passing from hexalydra "henc to toluche is of a similar charrier to the change in passing from pentane to amylene: the removal of H₂ in both cases producing a diminution of between 7 and 8 units in the S.V. This supports Kekulé's formula for benzone (Lossen, A. 225, 119; Horstmann, B. 20, 766).

The refractive power of benzene is about amal to that calculated on the assumption that Kekulé's formula is correct, provided that eering assumptions are made regarding the connection between the refractive powers of compound molecules and the refractive powers of the constituent atoms (Brühl, A. 200, 228; Kanonnikoff, J. R. 18, 473).

Passing from physical to chemical considerations, we note first that the behaviour of benzene towards halogens is, on the whole, more like that of a saturated chan an unsaturated compound. The following special arguments have also been omployed. Sodium acting upon succinic ether gives succinity-succinio other, which loses H, on or lation, changing to di-exyterephthalic ether. The formula of succinities succinic ether may be written in one of the following ways:

(I.) CO.Et.CH.CO.CH.

CH.CO.CH.CO.Et.

(II.) CO.Et.CH.C(OH):CH

CH.C(OH):C.CO.Et.

(III.) CO.Et.CH.C(OH):CH.

CH.C(OH):CH.CO.Et.

(IV.) CO,BLO : O(OH) OH,

(Y.) CO,Et.CH.CO—CH,

CO.T.L.CH.CO. CH...

If the first corroctly represents succinyl-snoc nio sther, it must be supposed to change into (II.), (III.), or (IV.) during the oxidation. The third formula would naturally lead to the formula

 $CO_1Et.C$ C(OII) - CH $C.CO_2Et$ and therefore

to the benzene formula HC

proposed by Wislioenus, a formula which would indicate the existence of two chloro-benzenes. The second formula

CH = CH

 CO_2 Et.CH< C(OH) = CH $> C.CO_2$ Et, would give CO_2 Et.< C(OH) = CH $> C.CO_2$ Et on oxidation,

CO₂Et.C CH—C(OH) O.CO₂Et on exidation, a formula based upon Kekulé's ring. The fourth formula, like the third, leads to the benzeue ring of Wislicenus. The fifth formula would lead to Kekulé's or Wislicenus' ring, but with the carboxyls in the ortho-position, whereas in terophthalic acid they are in the para-position.

Ladenburg's formula for dioxy-terephthalicacid is

CO₂Et.O CII — CH

CCO₂Et, the forma-

tion of which from formule I., II., III. or IV. requires the improbable assumption of a wandering of hydroxyl such as takes place when salicylic acid changes to p-oxy-benefit acid. Ladenburg's formula can be derived from V, but only by assuming a roarrangement of the unsaturated uniens.

By the action of sodium upon malenic ether a tricarboxylic ether,

is found to be phloregluein tri-earboxylic ether,

HOO O(OH), and this undonbtedly CO_Et. O O.CO_Et

favonrs Kekulé's hypothesis, especially when it is remembered that phlorogluoin, C(OH) forms a tri-oxim C(NOH)

C(OH) forms a tri-oxim C(NOH)

CH CH H,O CH,

C(OH) C(OH) (HON)O C(NOH)

number of researches curried out on the beaucon derivatives, the constitution of beaucon itself still remains unsettled.

Recent discussions on the Bensene formula.

Ladenburg, B. 19, 971; 20, 62; Baeyer, B. 19, 1797; A. K. Miller, C. J. 51, 208; Thomsen, B. 19, 2944; Claus, B. 20, 1422.

Orientation.

Benzene gives rise to only one mono-substitution product. It gives three di-substitution products, and those, assuming either Keknlê's or Ladenburg's formula, are named as follows:

1, 2 = 1, 6 is called ortho. 1, 3 = 1, 5 , , meta. 1, 4 , , para.

When we come to tri-substitution products we must distinguish several cases.—1. Compounds of the formula $C_*H_2A_3$, that is to say, where the three substituting elements or radicles are all alike. There are three such compounds.

1, 2, 3 is called consecutive.
1, 3, 5 ,, , , symmetrical.
1, 2, 4 ,, , irregular.

1, 2, 4 , , irregular.
2. Compounds of the formula C, H₁AB₂. There are six such compounds.—3. There are ten compounds of the formula C₂H₁ABC.

In the case of tetra-derivatives of benzene;

1. There are three compounds of the formula C₄H₂A₄;

1, 2, 3, 4 is called consecutive. 1, 2, 4, 5 ,, ,, symmetrical. 1, 2, 3, 5 ,, ,, irregular.

1, 2, 3, 5, ,, ,, irregular.

2. There are seven compounds of the formula $C_aH_AB_3$,—3. There are thirteen compounds of the formula $C_aH_AB_3$,—4. There are sixteen compounds of the formula $C_aH_ABC_a$.—5. There are thirty compounds of the formula $C_aH_ABC_a$.

There is only one penta-derivative of the formula C.HA,, and only one compound of the formula O.A.

The next question is how to determine, in a given ease, the position of substituting radiots in the benzene ring. In isolated cases it is frequently found that this may be settled by special considerations, but the only general method known is that which was thoroughly worked out, by Kokulé's pupil Körner, in a most laborious research, in the course of which he discovered no loss than 126 new compounds (G. 4, 305). This research has done more than anything else towards establishing the ring formula for benzene.

Suppose we convert C.H.Br. into C.H.Br.; by reference to a figure it will be found that we can introduce a bromine atom in place of an atom of hydrogen in ortho-dibromobenzene in such a way as to produce either a consecutive or an irregular tribromobenzene, but not so as to produce a symmetrical product.

Meta-dibromobenzene can give rise to consecutive, irregular, or symmetrical, tribromobenzene, while para-dibromobenzene can cally give rise to an irregular tribromobenzene.

give rise to an irregular tribromobenzene.

An unknown dibrombenzene is therefore para, ortho- or meta-, according as we can get one, two, or three tribromobenzenes by treating it with bromine. Thus the dibromobenzene from dibromoaniline gives rise to three tribromose.

mine the chief product of the action of two solicules of bromine upon benzene gives rise to only one tribromobenzene on further treatment with bromine; hence this product is

gives rise to two, and only two, tribromo-bensenes; therefore it is ortho-dibromobenzene. In order to investigate the constitution of a

given tribromobenzene, two methods may be followed: either introduce another atom of bromine in place of hydrogen and see how many tetrabromobenzenes result, or displace an atom of bromine by hydrogen and carefully oxamino bow many dibromobenzenes are formed. It can easily be seen by reference to the formula that consecutive tribromobenzene produces two dibromobenzenes and also two tetrabromobenzenes, while synanctrical tribromobenzeno produces one dibronio and one tetrabromo benzene, and irregular tribromobenzene gives riso to all three dibromobenzenes and all three tetrabromobenzenes.

The following are the melting and boiling points of the bodies described;

Dibromobenzenes.									
Ortho .			,	[-10] (2240)					
Meta .				fiquid (220°)					
Para .				[89°] (219°)					

Tribromobenzenes.

Consecutive		[87°]
Symmetrical		[120°] (278°)
Irregular .		4401 (2760)

Tetrabromobeuzenes.

[160°] [137°-140°] Consecutive (1, 2, 3, 4) Symmetrical (1, 2, 4, 5) Irregular (1, 2, 3, 5) am. [99°] (329)

It will be observed that the isomerides differ widely in melting-points, but very slightly in boiling-points, and this is usually the case where isomerism is due to difference of position of substituents in the benzene nucleus.

The orientation of any given benzene derivasive must be determined either by preparing it from one of the three bremobenzenes, or elso by preparing a bromobenzene from it.

Examples.

Para-dibromobenzene when treated with sodium and methyl iodide gives a dimethyl-benzene or xylene: C₂ll₄Br₂+2Cll₄I+4Na= 2NaI+2NaBr+C₄ll₄(CH₂)₂. By oxidation this is converted first into toluic acid, C.H.(CH.)CO.H. and next into terephthalic acid, C.H.(CO.II). It is therefore evident that the xylene, the toluic seid, and terephthalic seid, are all para compounds. Also since a certain bromotoluche, O.H.Br(C11,), whon treated with sodium and methyl iodido gives the above para-xyleno, it must be the para-bromotoluene, and the bromo-benzoic acid derived from it by oxidation— C₈H₆BrCH₅ + O₅ = C₈H₁Br.CO₂H + H₂O—must be para-bromobenzoio scid.

As another example we may take the orthoseries. A certain bromoaniline, C.H.Br.NH., is known to be ortho- because when the amidogen is displaced by bromine the product is orthodibeomobenzene. Now, this ortho-bromoaniline

may be get by reducing a betweentrobeniesh C.H.Br(NO.J., which may be breased by diamered in from a nitroaniline C.H.(NH.J)(NO.J. and this may be got by acting on a nitroaniso C.H.(OCH.)(NO.) by ammonia, and this nitropara-dibromobenzene. On the other hand, the artisol may be obtained from a nitrophenol minor product of the dibromination of benzene C. (Oll)(NO₂), and this nitrophenol may be itself reduced to an amidophenol C.H.(OH)(NH.). and this amidophenol may be converted by diazo-reaction into a chlorophenol C, II, (OH)CI, and this chloropheuol may be converted by cautious fusion with potash into a dioxybenzeno Call (Oll), All the compounds here enumerated are clearly ortho- compounds, and as the dioxybenzene is found to be pyrocatechin, we have proved that pyrocatechin is ortho-dioxybenzene.

In the para- series we may trace, in the same way, the connection between p-dibromelona-ene and p-nitroanisel C_sH₄(OCH_s)NO_s. Thence we proceed by the following steps: reduce C.H.(OCH.)(NII.), convert this into C, II, (OCII,)(OII) by nitrous acid, and treat with hydrio iodido. In this way we get a second dioxylenzene, which is found to be hydroquinone, and this body is therefore a paracompound. The remaining dioxybenzene is resorcin, which must be the meta- compound.

The rules governing substitution in the bennone molecule are discussed in the article Ano-MATIC SERIES. Derivatives of benzene are described, as Aniling, Phenol, Bromo., Bromo. KITRO-, CHLORO-, CHLORO-NITRO-, IODO-, METHYL-, NITHO, OXY. BENZENE, etc.

BINZENE HEXABROMIDE Bromine is dropped into boiling benzene exposed to direct sunlight; the hexabrouide crystallises out on cooling; it is separated from tri-bromobenzene by sublimation, and finally crystallised from a nixture of alcohol and henzene (Mitscherlich, P. 35, 374; Mennier, C. R., 101, 378; A. Ch. [6] 10, 269). Prisms, isomorphous with the (a) hexachlorido; not attacked by HNO. or II,SO. Alcoholic KOH splits it up into 11Br and u-tri brome benzeue. If the product of the action of ZnEt, upon benzenu hexabromide dissolved in benzeno be oxidised with ohromiomixture, benzoic, isophtbalic, terephthalio, and di bromobenzoio acids are formed (Ador a. Rilliet, Bl. 12124, 485).
BENZENE CARBOXYLIC ACIDS v. BENZOIC,

PHTHALIC, Tri-MELLITIC, Tri-MESIC, Hemi-MELLITIC, PREUNITIC, PYROMELLITIC, MELLOPHANIC. and MELLITIC ACIDS.

Benzeno penta-carboxylic acid C, Il.O. f.s. C.11(CO.11), Formed by oxidation of pentamethyl bearene (Friedel a. Crafts, A. Ch. [6] 1, 174). Amorphous (containing 6aq). The K salt forms small deliquescent prisms; the salts of Ag, I'b, Ba, Fe, Cu, and Al form insoluble pps.

BENZENE (a) HEXACHLORIDE C.H.Cl.

Moi. w. 291. [157]. S.G. 187. Prepared by chlorinating benzene in direct sunlight (Faraday, A. Ch. [2] 30, 275; Mitscherlich, P. 35, 870; Lesimple, Bl. [2] 6, 161): 850 g. may be got from 600 g. benzens (Leeds a. Everhart, A. C. J. 2, 205). It may be freed from C.HCl, and C.H.Cl, by treatment with H.SO, or HNO. (Meunier, A. Ch. [0] 10, 223). Monoelluse orystals; may be sublimed. At 286° it boils, splitting up into HCl and (1, 2, 4)-tri-chlorobenzone. The same decomposition is effected by heating with alcoholic KOH.

Reactions.-1. Zinc reduces it, in alcoholic solution, to benzene (Zinin, Z. 1871, 284).— 2. Fumling nitric acid has no action.—3. Silver

acetate forms crystalline C.H.Cl.(OAc), C.H.Cl.
Bengene (d) hexachloride C.H.Cl. [310].
V.D. 9 28. Formed at the same time as the (a)compound; when the mixture is sublimed, the (8)- compound sublimes last. If the mixture (4 pts.) be boiled with KCN (3 pts.) and alcohol, the (B)-compound is left while the (a)-compound la decomposed. Regular octaliedra, cubes, tetrahedra, or tetrakis tetrahedra. Alcoholic potush splits it up into 11Cl and (1, 2, 4)-tri-chlorobenzene, but more slowly than the (a)-compound

(J. Meunier, C. R. 98, 436; 100, 358).

BENZENE-HYDRAZIMIDO- v. pp. 369, 370.

BENZENE - PHENYL - HYDRAZIMIDO -NAPHTHALENE v. Benzene-AZO-phenyl-(6)naphthylamine.

BENZENE.PYROOALLOL.PHTHALEIN Tri - OXY - TRI - PHENYL - CARBINOL - CARBOXYLIC AMELYDRIDE

BENZENE-TRI-QUINONE C.O. 4aq. ealled 'oxy carboxylic acid' of hereli. [c. 95°].

Formation. - 1. By the action of HNO, upon the hydrochloride of tetra-oxy di amido-benzene; the yield is 65 p.c. -2. By the action of HNO, npou di-imido-di-oxy-quinone Ca(N11)2(O11) Og-8. By oxidation of hexa-oxy-benzene C. (OII) ..

Properties .- Colourless microscopic needles. Nearly insoluble in cold water, alcohol, and ether.

Reactions. By roducing agents it is converted successively into di-oxy-benzene-diquinone C₀(OII)₂C₀, tetra-oxy-benzene-quinone C₀(OII)₄C₀, and finally hoxa oxy-benzene C₀(OII)_n. On heating to 100° or on boiling hoxa oxy-benzeno with water it evolves CO, and yields erocome aoid O.H.O. (Nietzki a. Benekiser, B. 18, 504). BENZENE-RESORGIN PHTHALEIN v. Di-

OXY - TRIPHENYL - CARBINOL - CARBOXYLIC HYDRIDI

BENZENE - SULPH - AMIDO - ANILIDE v. BENZENE-SULPHONIC ACID.

BENZENE - SULPH - AMIDO - TOLUIDE v. BENZENE-SULPHONIC ACID.

BENZENE SULPHINIC ACID C.II.SO, i.e. O.H.SO.H. [84°].

Formation .- 1. By adding zinc-dust to a cooled alcoholic solution of the chloride of benzene sulphonie acid; the resulting zine salt is very slightly solublo in water; it is treated with Na.CO.; the filtrate is concentrated and the acid ppd. by 11Cl (Schiller a. Otto, B. 9, 1584).—2. Frank the phenyl-hydrazide of benzene sulphonic acid 178O,N,11,Ph, called also di-phenyl-sulphazide, by boiling with baryta-water (Limpricht, B. 20, 1239). - 3. By passing SO, into a warm mixture of benzeno and Al₂Cl₄ (Friedcl a. Crafts, C. R. 86, 1368; Adrianowsky, B. 12, 853).—4. By the action of ZnEt, on C.H.SO.Cl (Kalle, A. 119, 156).— 5. From C.H.SO.Cl and Pb(SEt), thus:

2PhSO,Cl + 2Pb(SEt), =

(PhSO₂), Pb + PbCl₂ + S.Et₄ {Schiller a. Otto, B. 9, 1636)...6. From diphenyl disniphide and alcohol potash: 2Ph₂S₂ + 4KOl1 = PhSO₂K + 3PhSK + 2H₂O (S. a. O.).

Properties.-Long radiating prisms. Sl. sol. sold, v. sol. het, water; v. sol. alcohol and

other; said to test-paper. Above 1000 it decomposes.

Reactions .- 1. Water at 180° gives benzene sulphonie acid and phenyl benzene-thiosul-phonate (Otto, A. 145, 317); the same reaction takes place slowly in the cold, especially in presence of HCl (Pauly a. Otto, B. 10, 2181) .-2. Ethyl mercaptan at 100° gives di-ethyl disulphide and phenyl-ethyl di-eulphide (Otto a. Rössing, B. 19, 3136). - 8. Phosphorus penta-PhSO,H+PCL thus; chloride reacts $= PCl_1 + IICl + PhSO_1Cl. - 4.$ Potash fusion gives benzene and K2SO3.-5. Sodium chloroacctate gives phenyl sulphonyl acctio acid, Ph.SO, CH, CO, H .- 6. Sodium di-chloro-acetale gives phenyl chloro methyl sulphone Ph.SO, CH, Cl. -7. Methylene iodide resets gives thus: Cll, I, + Ph.SO.Na = NaI + Ph.SO.CH,I.-8. Sodium aa-di-chloro-propionate acting upon sodium benzene sulphinato gives di-phenyl othyleno di-sulphone, Ph.SO, C, H, SO, Ph. 9. Phenyl-hydrazine in presence of IICIAq forms phenyl benzene-thiosulphonate and the phenyl hydrazide of benzene sulphonio acid (q. v.).-10. When H.SO, is added to a solution of Ph.SO, Na and NO, Na a pp. is got which may be crystallised from alcohol. It weter, CS₂ or ligroin, but v. sol. alcohol and ether. At 100° it evolves nitrons acid gas. Boiling water, alkalis, or acids, decompose it into PhSO,11 and nitrous acid (Königs, B. 111, 615).—11. Funing niltric acid forms C₁₄11₁,NS₂O, which may be (Ph.SO₂)₂NO. It forms erystals, [98·5°], insol. alkalis, sl. sol. alcohol, m. sol. benzene (Otto a. Gruber, A. 141, 370; Königs, B. 11, 615, 1590).

Salts .- BaA'2: clumps .- ZnA'2: tablets, sl. sol. alcohol and other, sl. sol. water (Kalle) .-ZuA', 2uq: insol. cold water (S. a. O.). -AgA'.

Ethyl ether EtA' .- Formed by means of EtOII and 11Cl, or, together with CO, by heating Ph.SO.Na with Cl.CO.Et. Non-volatile oil. KMnO, in acetic acid solution exidiscs it to Ph.SO₂Et (Otto a. Rössing, B. 18, 2495; 19,

Benzene di-sulphinio acid C.H. (SO.II), [1:3]. From [1:3] C.II. (SO.CI), and zino-dust (Panly, B. 9, 1595). Oil.—BaA

BENZENE-SULPH-NITR-ANILIDE v. BENZ-ENE-SULPHONIC ACID.

BENZENE SULPHONE v. DI-PHENYL-SUL-

BENZENE SULPHONIC ACID C.H.SO. i.e. C.II. SO.H. Phenyl-sulphurous acid. Sulphobenzolic acid. [42].

Formation.—I. From benzeus and fuming

H.SO. (Mitschorlioh, P. 31, 283, 634; Stenhouse, Pr. 14, 351; Wurtz, C. B. 64, 749). 2. By the exidation of benzene sulphinio acid (Otto a. Ostrop, A. 141, 369) .- 3. By the oxidation of phenyl mercaptan: PhSII + O, = PhSO,H (Vogt, A. 119, Ib1). -4. By boiling p-diazo-benzone sulphonio acid with alcohol under pressure (R. Schmitt, A. 120, 129) .- 5. Together with phenyl benzene-thiosulphonate by heating benzene eulphinio acid with water at 180° (Otto, A. 145, 817):

8Ph.SO.H - PhSO.II + Ph.SO.SPh + H.O. Preparation.—Benzene (2 pts.) is shaken with fuming H.SO. (3 pts.) with gentle warming. The seid is separated from undissolved benzene, diluted, and neutralised with BaCO. or lead carbonate. In the filtrate the Ba, or Pb, salt is decomposed by H,SO, or H,S respectively.

Properties .- Small, four-sided, deliquescent

plates (containing 1 aq).

Reactions.-1. By fusion with potash, soda, or a mixture of the two, it is converted into phenol. The percentage of phenol obtained increases with the amount of alkali and with the temperature of the fusion. The percentage of phenol is given in this table; one equivalent of acid being used :

кон	NaOH	Temperature	Phenol
2		253°	23
3		210°	7
8		267°	79
-	3	2090	1
	3	280°	26
8	3	211°	2
3	3	277°	89
3	8	360°	64
7		252°	96

(P. Degener, J. pr. 125, 401). 2. The potassium salt distilled with KCN or K, FeCy, gives benzomitrile (Merz, Z. [2] 5, 33). 3. Dry distillation gives 11.80, benzene, 80, and di-phenyl-sul-phone.—4. Dry distillation of the ammonium salt gives benzene and small quantities of beuzene sulphamide, diphenyl, di-phenyl sulphone, phenyl mercaptan, and (traces ol) quinoline (Egli, B. 18, 575) .- 5. The potassium salt distilled with NaNII, gives aniline (Jackson a. Wing, Am. 9, 75) .- 6. Distillation of the Na salt gives di-phenyl sulphide, di-phenyl di-sulphide, phenyl mercaptan, CO,, and SO, (Stenhouse). -7. A mixture of H.SO, and water boiling at 175° converts it into benzene and 11,80, (Armstrong, C. J. 45, 151) .- 8. Fusion with potassium formate gives potassium benzoate.

Salts (Freund, A. 120, 76; Kallo, A. 119, 161).—BaA, aq: pearly plates, sl. sol. alcohol. -CuA'₄Gaq: large blue tables, sol. alcohol. -AgA'8aq: tables. - ZnA'₄Gaq: six-sided tables. Methyl ether MoA'. S.G. 41 1-27. Formed

y action of NaOMe upon Ph.SO,Cl in ether (It.

Hühner, A. 223, 225). Oil.

Ethyl ether Eth'. S.G. 12 1-22. From NaOEt and PhSO, Cl in ether. Formed also by oxidising PhSO Et (Otto a. Rössing, B. 19, 1225). Oil, miseible with alcohol, ether, and benzene. Saponified by boiling water.

Propyl ether PrA'. S.G. 11 1-179 (II.). Phenyl ether PhA'. [35°]. Formed by

acting upon Ph.SO,Cl dissolved in benzene with sodium phenol. Formed also by action of zincdust on a mixture of phenol and PhSQ.Cl. Trimetric crystals, a:b:c = 6847;1; :8576. V.sol. benzene, ether, and alcohol, insol. water. Slowly saponified by boiling aqueons KOH; alcoholic NII, even at 200° does not affect it. On nitration it gives the nitro-phenyl ether, and also a tri-nitro- derivativo [116°] (Schiaparelli, G. 11, 66; R. Otto, B. 19, 1832).

p-Nitro-phenylether C.H., SO, C.H. (NO.). [82]. Formed by nitrating the preceding; or from p-nitro-phenol, ZnCl., and PhSO.Cl (Schia-parelli, G. 11, 70). Sl. sol. cold alcohol. Chlorida Ph.SO.Cl.—Benzene sulphochlo-

section of PCI, on a sait of bensene sulphonic acid (Gerhardt a. Chancel, C. R. 35, 690), or by passing chlorine into an aqueous solution of PhSO₂H (Otto a. Ostrop). Oil; v. sol. alcohol and ther. Slowly solidifies at 0° forming large pombio crystals. May be distilled in vaoue, but is much decomposed on boiling under a tmospheric prossuro. Hardly attacked by water. Reactions ... 1. Tin and IICl form phenyl

mercaptan.—2. Sodiumumalgam or ZuEt, forms a benzene sulphinate.—3. PCl, at 210 gives a benzene sulphinate.—3. PCl, at 210³ gives C.H.Cl, phosphorus oxychloride, and SO₂Cl₄ (Kekulé a, Barbaglia, B. 5, 876).—4. PbO₂ gives at 180° PbSO, and Call Cl (Wallach, A. 214, 219).-5. Phenol (1 mol.) and zinculust gives Ph.SO,Ph; phenol (1 mol.) and ZuCl, gives Ph.SO,C,H,O.SO,Ph (2) [123] (Schimparelli, G. 11, 66).

Bromide Ph.SO.Br. From PhSO, II and Br (Otto, A. 141, 372). Oil.

Amide Ph.SO,NII, Benzene sulphamids. Benzene sulphonamide. [156'] (Hybbeneth, A. 221, 206). S. 43 at 16°. Formed by the action of NII, on the chloride or bromide (Otto a. Ostrop, A. 141, 365), or, in small quantities, by heating the anunonium salt at 200" (Stouhouse,

Pr. 11, 351). Needles (from water) or plates (from alcohol). V. al. sol. water, sol. hot NILAq, v. sol. alcohol and ether. Anunoniacal AgNO.

PhSO,N:(C,O.)C,H., [160°], whence cone. NH,Aq produces Ph.SO,N.CO.C,H.,CO.NH., [166°] (Gerhardt a. Chancel, C. R. 35, 690; Gerhardt a. Chiozza, A. Ch. [3] 47, 129).

Benzoylderivative Ph.SO, NIBz. [147°]. From beazene suphonomide and BzClat 145° (Gerhardt, A. 108, 214; Wallach, A. 214, 210). Prisms (from alcohol). Salt. PhSO NNaBz. Silky needles (from alcohol). Iteactions. -1. PCI, gives the imide-chloride Ph.SO, N.CCI.Ph This forms triclinic plates (from benzo-1809 line) a:b:c = '862:1: 7, a = 87' 59', B 91' 31', y=

63° 24'. When this imido-chloride is heated it splits up into benzonitrile and benzone sulpho - eliloride (Wallach a. Gossmann, A. 214, 210). Aniline converts the imide-chloride into phenyl-sulphonyl-phenyl-benzamidlne (PhSO,N):C(NHBH).Ph.—2. The Pb and Ag derivatives are converted by Etl at 100° into the

original amide (Remsen a. Palmer, Am. 8, 235), Dibenzoylderivativel'bSO, NBz, [1059], An oil. Methylamide Ph.SO, NUMe. formed by treating the chloride with aquoous

methylamine (Itomburgh, R. 3, 16). Methyl-nitro-amide Ph.SO, NMe(NO).
4°]. From the preceding and HNO, (S.G. 148).

Ethylamide Ph.SO, NEtll. [58°]. From the chloride and NEtH, (Romburgh, R. 3, 13). Ethyl-nitro-amide Ph.SO, NEt(NO.). [44°]. Formed by the action of HNO, on the preceding or on the succeeding compound.

Needles (from alcohol); volatile with steam.

Di methylamide C.H. SO NMe. [48] Di-ethylamide Ph.SO, NEt. [42°].
Anilide Ph.SO, NHPh. [102°] (Wallach,
A. 214, 221). S. 4.3 at 16°. From the chloride and aniline (Biffi, A. 91, 107; Gericke, A. 109. 217: Meyer a. Ascher R 4 8961

o-Nitro-anilide Ph.SO, NH.C, H.(NO,) [1:2]. [104°]. From e-nitro-aniline and PliSO, (Lellman, A. 221, 16; B. 16, 591). Yello

plates, sol. alcohol, glacial HOAc, and CHCl. m-Nitro-anilide Ph.SO, NH.C.H. (NO. [1:8]. [132]. From m-nitre-anilins and PhSO,Cl. Flat yellow needles (L.).

p-Nitro anilids Ph.SO .. NH.C. H. (NO..) [1:4]. [139°]. From p-nitro-aniline and Ph.SO.Cl. Yellow crystals. o-Amido-anilide Ph.SO., NH.C. H. (NII.)

[1:2]. [168°]. From the c-n-tro-amilide by tin and HCl (L.). Needles (from 50 p.c. alcehol). V. sol. alcehol, sl. sol. ligroin. - B'HCl.

p-Toluide Ph.SO, NH.C, H, Me [1:4]. [120]. From the chloride and p-toluidins (Wallach a. Huth, B. 9, 427).

m-Nitro-p-teluide Ph.SO, NILC, II, Me(NO.) [1:4:3]. [99°]. From the preceding by nitration; or from Ph.SO.Cl and nitro-p-toluidine (Lollmann, A. 221, 18). Cubss (from alcohol). Not attacked by alcoholic KOII.

Di-nstro-p-teluids
Ph.SO_{2.NII.C.11.Me(NO₂)₂. [178°]. Fermed by nitration of the p-toluide. Yellow prisms, sl. sol. cold alcehol (Lellmann, B. 16, 595). Not} attacked by alcoholic KOII.

m. Amido-p-toluide

Ph.SO₂,NII.C₆H₃Me(NH₂) [1:4:3]. [146·5°]. From the nitro- compound by tin and HCl. Colourlese needlee (from dilute alcohol); sl. sol. water,

di phenyl amide Ph.SO.NPh. [124°]. From PhSO.Cl and NHPh, at 200° (Wallach, A. 214, 220). Silk-like needles (from alcohol). Sol. alcohol, ether, or benzene, insol. water. Cone. H.SO, forms a blus solution. Insol. HCIAq.

Phonyl-hydrazide C₁₂H₁₂N₂O₂S Phenyl-benzene-sulphaside. Ph.SO, N.H.Ph. bi-phenyl sulphazide. [148°-150°]. Forma-tion.—1. By the action of SO, upon diazo-ben-zene (Kamigs, B. 10, 1531; Wiesinger, B. 10, -2. From benzene sulphonic chloride and phenyl-hydrazine (Fischer, A. 190, 132).—3. From benzene-sulphinic acid (q. v.) and phenylhydrazine hydrochlorido (Escales, B. 18, 893),-4. By reduction of C. II. N., SO, C. II, with zing-dust and acetic acid. Preparation. - Aniline is dissolved in alcohol saturated with SO2; ths solution is cooke-1 Cow 0° and a cone, selution of about double the theoretical -nantity of KNO. is slowly added; after standing for 24-36 hours it is precipitated by water; the yield is 80 p.o. Properties. - White felted needles (from alcohol). With NaOEt it gives a very unstable crystalline sodium compound C₁₂H₁₁NaN_SO. Reaction.— By boiling with aqueous alkalis (i.s. barytawater) it is decomposed into benzane sulphinic acid, benzene, and N_s; Ph.N_sH_s.NO_s.Ph. = PhH + PhSO_sH + N_s (E., B. 18, 893; Limpricht, B. 20, 1288).

BENZENE-o-DISULPHONIC ACID C.H.S.O. te. C.H.(SO.H), [1:2]. From amido-benzene meniphonic acid by sulphonation, diszotisation, and boiling with alcohol (Drebes, B. 9, \$48).

No.

Amide C.H.(SO,NH.). [2889]. Benzene-m-disulphonic soid C C.H.(80,H).

Formation. - From amido-benzens p-sulphonic acid by sulphonation, diazotisation, and heating with alcohol (Zander, A. 198, 8).

Preparation.-When beuzene or benzene eulphonic acid is treated with funning H,SO, both m and p disulphonic acids are formed. The m acid is formed chiefly when the temperature is low or when this mixture is kept at a high temperature for a short time only (Buckton a. llofmann, C. J. 9, 255: Barth a. Senhofer, B. 8, 754, 1477; 9, 969; Limpricht, B. 9, 550; Körner a. Mouselise, B. 9, 583). Benzeno (1 pt.) is freed from thiophens by shaking with conc. H.SO., and is then dissolved in furning (70 p.c.) sulphurie acid (4 pts.) at 40°. The solution is heated for 2 hours at 275°; cooled; peursd into water, and neutralised with lime; CaSO, is remeved by filtration, and the lime ealts of the m and p acid may be separated by crystallisation, the former separating first (Binschedler a. Busch, Monit. Scient. 1878, 1169; cf. Egli, B. 8, 817; Heinzslmann, A. 188, 159).

Properties.—Very deliquescent crystals (containing 2 aq). The alkalins salts are v. sol. water.

Salts.—Na,A"4aq.—K,A"aq. S. 66.6 at 100°.—BaA"2aq. S. 44.2 at 100°. Large prisms.—CuA"6aq: v. sol. water.—CuA"1\aq.—ZnA"4aq.—PbA"2uq. S. (of PbA') 86.2 at 26°.—Ar.A" Λg.A".

Reaction.—1. By fusion with potash or soda it is converted into resorcin. The acid is first converted into m-phenol sulphonic acid (at 180°). The amount of resorcin formed by fusing this body (I mol.) with potash (21 mels.) at 270° for 10 minutes is 2.7 p.c., at 270° for 20 minutes it is 21 p.c., and at 250° for 30 minutes it is 26 p.c. When soda, or a mixture of potash and coda, is used, the yield is rather less (Degener, J. pr. 128, 318). - 2. By fusion with potassium cyanids the potassium salt is converted into Call, (CN), which, when boiled with potash, gives isophthalic acid (Wislicenus a. Brunner, B. 4, 981; Ross-

acid (Wisheenus a. Brunner, B. 4, 1001; Mona-Garrick, Z. 5, 549; Barth a. Senholer, A. 174, 238; B. 8, 754; V. Meyer a. Miehler, B. 8, 672).

Chleride C.H.(SO.Cl), [63°]. From addium benzene disulphonate and PCl, or S.O.Cl. (Heumann a. Köchlin, B. 16, 483). Monosymmetrical crystais, a:b:c=1·1991:1:0·8688, \$=85° 44' (Otto, B. 19, 2424).

Amids C.H. (SO.NH.). [229°]. Needles. Benzene-p-disulphenicacid C.H. (SO.H.). [1:4]. Prepared as abovs. The potassium salt distilled with KCN gives di-cyano-benzene, which, on saponification, gives terephthalic acid (Wislicenus

a. Brunner, B. 4, 984).
Salts. -K,A"aq: thin plates. S. (of K,A")
66-6at100°.—BaA"aq. S. (of BaA") 7-19 at 100°.— Chlorids C.H.(SO,CI), [1319]. Needles. S. (of PhA")

Amide C.H. (SO,NH.). [288°]. Thin scalse (from water).

s-Bensone-tri-sulphonic soid C.H. (SO.H) [1:3:5]. Prepared by heating 5 pts. of the potassitus sait of the mono- or m. disaphhonia said with a

the of Mining H. 100, in an open disk till H. 100, colabilises; this ready sulphonation is probably due to the presence of KHSO, or of potassium pyrosulphate. By heating the potassium salt with KCN the nitrile of trimesio acid C.H. (CO.H), is obtained. NaOlI fusion yields phlorogluciu The Ba salt is eparingly soluble in water.

Salts.-K.A"3aq.-Ba,A",-Ba,A", 6aq.-A," 4aq: slonder needles, v. sol. water.-Pb_A_"4aq: slender needles, v. Ag_A" 8aq (Senhofer, A. 174, 243).

Ohloride: [184°], Amide: [306°] (Jackson a. Wing, B. 19, 898).

Benzoyl-amide Calla(SO,NHBz)3. [285] AnilideC,11,(SO,NPhH),[237-](J.Am.9,346) Derivatives of the sulphonic acids of benz-

sne are described as - Diazonenzene schemonio ACID (p. 405), Sulpho-henzene-azoxy-bynzene видиноми асто (р. 428), Амиро-, Вкомо-, Вкомо-AMIDO., BROMO-NITRO., CHLORO., IODO., METHYL., NITRO, PROPYL-, RENZERS SULPHONIC ACIDS.

BENZENE DI-SULPHOXIDE v. PHENTL

BENZENE-THIOSULPHONATE.

BENZENE SULPHYDRATE C. PHENYL MER-CAPTAN.

BENZENE THIOSULPHONIC ACID

C,H,SO,SH.

Preparation .- 1. By the action of KIIS on benzene-sulphonic chloride. -2. By beating a solution of a salt of benzene-sulphinic acid with sulphur.

Salts .- KA': v. sol. hot alcohol and water, NaA' 1 aq.

Ethyl ether C.ll.SO.SEt. phenyl-di-sulphoride. From the potassium salt and EtBr. Colourless heavy oil. Insol. water, miscible with alcohol and ether. Slowly volatilises with steam. On reduction with zinc and 11,80, in alcoholic solution it gives phenyl mercaptan and ethyl mercaptan. KOllAq gives benzene sulphinic acid and diethyl di-sulphide (Otto, B. 15, 127).

Ethylene ether A', C. II, : [85°]. Formed by heating an alcoholic solution of the Na or K salt (2 mols.) with cthylene bromide (1 mol.). Small thin silky needles. Withent taste or smell. V. sol. benzene and hot alcohol, far less in cold alcohol. By warming with alcoholic KOH it gives benzene sulphinic acid, ethane disulphinio acid C₂H₁(SO₂H)₂₁ and di-ethylene tetra sulphide (C₂H₁)₂S₄. By warming with alcoholic KHS it gives the potassium salt and ethylene mercaptan C,II,(HS), If the alco-holic solution is warmed with ethylene mercaptan, benzene sulphinic acid and di-ethyleno tetra sulphide are produced. On reduction it gives primarily benzene sulphinic acid and ethylene mercaptan. By warming with alcoholic H,S it is converted into phenyl tetrasulphide, ethylene sulphydrate, &o. (Otto a. Bossing, B. 20, 2079, 2090).

Phenyl ether C.H. SO, SC. II. Benzene di-sulphoxide. Diphenyl di-sulphoxide. [45]. Formed by the decomposition of benzene sulphinic soid (q. v.) by boiling water, or even by apontaneous decomposition (Panly a. Otto, B. 9, 1639; 10, 2181; 11, 2070). Monosymmetrical prisms, a:b:c=1446:1:14709 (Otto, B. 15, 131). Reactions.-1. By saponification with alkalis it is decomposed into benzene sulphinic acid and di-phenyl-di-sulphide:

ACH_BOLE+ (CH.)

place in two stages:

(a) 20 H., 80 Sc.H., + 2H., 0 =

30 H., 80 H + C.H., 8H.

(b) C.H., 80 S.G.H., + C.H., 8H =

C.H., 80 H + (C.H.), S.

The benzene thiosulphouic-phenyl-ether reacts

in the cold with sodium phenyl mercaptide according to the last equation (b) (Otto a. Rössing, B. 19, 1235). 2. Zinc-dust added to an alcoholic solution gives zinc phenyl mercaptide and zine benzene sulphinate. - 8. H.S. forms benzene sulphinic seid, phenyl mercaptan, phenyl tetra sulphide, &c. -4. Zine phenyl mercaptide added to an alcoholio solution gives diphenyl di-sulphido and zinc benzone sulphinate .- 5. Phenyl mercaptan on heating forms di-phenyl di-sulphide and benzene sulphinie acid ... 6. Ethyl mercaptan at 115 gives di-ethyl di sulphide, phouyl-ethyl disulphide, di-phenyl di-sulphide, and other products (Otto a. Russing, B. 19, 3137).

BENZENYL ALCOHOL v. Ortho-BENEOIS ACID.

BENZENYL-AMIDINE v. BENZAMIDINE. BENZENYL-TRI-AMIDO-BENZENE

C,H,N, i.e. Ph.C N11 C, 11, NII, 1240%. Amido-phenylene-benzamidine. From benzonylnitro-phenylene-diamine, tin, and HCl (Hübner, A. 208, 809). Needles, v. sl. col. water, v. sol. alcohol.-B"211Cl. B"211NO, B"H SO, 2aq. Bensoyl derivative

Ph.C NII C. H.NIIIz. [125°-214°]. di-benzoyl-nitro-phenylene-diamine dissolved in HOAc by boiling with tin and 11Cl (Ruhemann, B. 14, 2653). Plates (containing aq. from dilute alcoh...). B'HCl: needles.

BHNZENYL-(β)-AMIDO-a-NAPHTHOL

 $C_{ii}H_{ii}NO \text{ i.s. } C_{ie}H_{e} < \stackrel{O}{N} > C.C_{e}11_{s}, [122]^{\circ}]. \text{ Prepared}$ by reduction of the benzuyl-derivative of enitroso (a) mphthel (Wurms, B. 15, 1816). Colouriess needles. Sublimable. Sol. alcohol and acctic acid, sl. sol, water.

Benssayl-(a)-amido-(B)-naphthol

Formation .- 1. As a by-product in the reduction of (a)-nitro (B)-naphthyl-benzoata.

2. By heating benzoyl-anido-(B)-naphthol.
(Böttcher, B. 16, 1936; C. C. 1884, 898).

Properties. - Long colourless needles. Sublimable. V. c. sol. alcorol ether, and benzen sl. sol. petroleum ether, mscl. water. Dissolver in streng acids. Its solutions bave a beauti ful blue fluorescence. B',ll,Cl,PtCl,: yellow

Benzenyl-(a)-amido-naphthyl mai Captan $C_{i_0}H_i < \stackrel{S}{N} > C.C.H_i$, [103°]. Formal by heating benzoyl-(a)-naphthylamine (2 pte with sulpbur (1 pt.) for two hours (yield 10 pt. (Hofmann, B. 20, 1798). Obtained also be oxidation of the thiobenzoyl derivative of an aphthylamine 0, H., N.G(SiI), C.H., with possium ferricyanide (Jacobsen, B. 20, 1806). sium ferricyanide (Jacobsen, B. 20, 18 Glistening colcurless needles (from alcol V. sol. ether, benzene, and hot alcohol.

Caprage MU,H,[NO],OH; [1819]; small | SERRERIL ARIDO TOLYL MERCAPIAN orange needles.

Benzehyl - (6) - amide - naphthyl mercaptan C,H,<8>0.0,H, [107°]. Prepared by Felted needles. -B',H,Cl,PtCl, (Hofmann, B. 20, 1803).

BENZENYL - AMIDO - PHENANTHROL PRENANTURENE.

BENZENYL-o-AMIDO-PHENOL C13H, NO i.e. o-amido-phenol and BzCl or phthalic anhydride (Ladenburg, B. 9, 1526). Plates (from dilute alcohol). Insol. water, sol. dilute H.SO. Split up by HCl at 130° into henzoic acid and e-amidophenol. Its salts are decomposed by water .-B',H,PtCl,

BENZENYL-DIAMIDO-DIPHENYL C,,H,,N, 4.6. C_eH₃.C≪N₁₁>C_eH₄.C_eH₄. [198°]. From benzoyl-nitro-p amido-diphenyl, tin, and glacial HOAo (Hübner, A. 209, 347). Plates (frem alcohol). - B'fICl, -B'211, l'tCl, -B'2H2SO4.

BENZENYL . AMIDO . PHENYLENE DIAMINE v. BENZENYL-TRI-AMIDO-BENZENE.

BENZENYL-AMIDO-PHENYL MERCAPTAN C₁₁H₂NS i.e. [1:2] C₂H₂<S CPh. [115°]. (c. 360°).

Formation. - 1. By heating amido phenyl mercapian with BzCl, with benzoic aldehyde (probably benzyl alcohel is also formed), or with benzenitrile (NII, being evolved). -2. By heating phenyl anilido acctenitrile with sulphur (H.S and HCN being evolved).—3. In small quantity by the action of benzoyl chloride on methenylamido-phenyl mercaptan .- 4. From thiconzoic anilide and alkuline K.FeCy. (Jacobsen B. 19,

Freparation. Benzanilide (2 pts.) is heated with S. (I pt.) for a few hours to boiling. The yield is 50 to 60 p.e. of the benzanilide used. Properties. - Noedles.

Weak base. aloohol, ether, CS, and HClAq. Has a pleasant smell of tea-roses and geraniums. It remains almost unaltered on boiling with aqueous acids or alkalis. On fusion with KOH it is decomposed into benzoic acid and o amido-phenyl merceptan. Salt .- B'HAuCl.

References .- Hofmann, B. 12, 2359; 13, 7, 1286; Tiemann a. Piebst, B. 15, 2033.

BENZENYL . AMIDO . THIO . CRESOL BENZZNYL-AMIDO-TOLKI CIERCAPTAN.

BENZENYL . AMIDO . THIO . PHENOL v. BENZENTL-AMIDO-PHENYL MERCAPTAN.

BENZENYL . TRI . AMIDO . TOLUENE $C_{14}H_{12}N_{2}$ i.e. $Ph.C \leqslant \frac{N11}{N} > C_{4}H_{2}Me.NH_{2}$. $\begin{bmatrix} 4 \\ 6 \end{bmatrix} : 1:3$ [188°]. From benzoyl-di-nitro-toluidine, tin, and HCl (Kelbe, B. 8, 877). Needles, insol. water.—B HCl.—B H.SO, aq

BENZENYL-TRI-AMIDO-FOLUENE Ph.O NH O.H.Me.NH. [1:1 6] (?).

Bensoyl derivative CnH, N,Oaq. (196-218). From di benzoyl-aitro-m-tolylene-diamine, tin, and HCl (Ruhamsan, B. I4, 2656). Needles (from alcehol).

C. H. NS i.s. C. H. (CH.) \ N C-O. H. [125] Prepared by heating amido-tolyl mercaptan C_sH_sMe(SH)NH_s [1:3:4] with benzoyl ohloride. heating benzoyl (8) naphthylamine with sul- Siender needles.—B'HCl: tables, decomposed by water.—(B'HCl),PtCl, aq (Hoss, B. 14, 493).
BENZENYL-AMIDOXIM v. BENZ-AMIDOXIM.

DI-BENZENYL TRI-AMINE C14H12N, i.e. {Ph.C(NH)}2NH. [109°]. Formed by boiling benzamidine with Ao.O. Needles (Pinner a. Klein, B. 11, 8).

BENZENYL - AMYL - PHENYLENE - DI-AMINE v. BENZENYL-PHENYLENE-DIAMINE.

BENZENYL - BROMO - PHENYLENE - DI-AMINE

C₁₃H₉BrN₂ i.e. C₃H₃C NH C₈H₃Br [1/2:4]. [200°]. From benzoyl-bromo-nitro-aniline by reduction (Hübner, B. 8, 564; 10, 1710). Small needles, insol. water. - B'HCl. - B'HNO.-B'H SO

BENZENYL-DICINNAMYLENE-DI-AMINE

PhCH:C11.C1F-NH, C23H22N, i.e. CPh [207°]. PhCH:CH.CH_N /

Phenyl - di - styryl - glyoraline di-hydride. Formed by heating di-benzoyl-dicinnamylene-diamine with petash in a sealed tube (Japp a. Wynne, C. J. 49, 470). Faint yellow crystals. Beiled with HCl it forms a hydrochloride, sel. Elflo, which yields a platinochloride (C_nH_n,N_n,licl),Ptcl.

BENZENYL-ETHOXIM-CHLOBIDE

 $C_{\rm s}H_{\rm s}CCl:{\rm NOEt}$ (125°) at 45 mm.; (230°) at 760 mm. V.D. (to H) = 9.26 (ebs.). Colourless oil. V. sol. alcohel and ether, insol. water. It is very stable towards water, acids, and alkalis. Formed by the action of NaNO, and HCl upon the ethyl other of benzenyl-amidoxim C.H.C(NH.):NOEt. By heating with alcoholic NH, the parent substance is reproduced. By the patent shostance is reproduced. By heating with sodium ethylate it yields benzenylethoxim - ethyl ether C.H.G(OEt):NOEt (Tiemann a. Kriiger, B. 18, 727; 1057).

BENZENYL - ETHOXIM - ETHYL ETHER

C.II. C(OEt): NOEt. a. Ethyl-bers-hydroxamic-cthyl ether. (128°) at 40 nm.; (238°) at 760 mm. V.D. (to II) = 96·75 (obs.). Celourless oil. Insol. water. Formed by heating benzenylethexim-chloride C.H. C(NOEt)Cl with sedium ctlylate. By IICl it is decomposed into benzoic ether and the ethyl-ether of hydrexylamine

H₂N,OEt (Tiemann a. Krüger, B. 18, 742).

BENZENYL-ETHYL-AMIDINE C₂H₁₂N₂ i.e.

PhC(NEt).NH₂. From benzamidine and EII

(Pinner a. Klein, B. II, 7). Oil.—(B'HCl)₂PtCl₄.

DI-BENZENYL-IMIDO-AMIDE v. DI-BENZ

ENYL-TRIAMINE BENZENYL - ETHYL - PHENYLENE - DI -AMINE v. Benzenyl-phenylene-diamini

BENZENYL-METHOXIM-CHLORIDE C,H, CCl:NOMe. (225° uneorr.). V.D. 74-95. Formed by the action of HCl and NaNO, npon the methyl ether of benzenyl-amidoxim C,H,C(NH,):NOMe (Krüger, B. 17, 1689; 18, 1057). Oily fluid. Very volatile with eteam. Sol. alcohol, ether, benzene, and ligroin, insol.

BENZENYL-METHYL-PHENYLENE - DI -AMINE v. Beneentl-phenylene-diamine.

BEEKEYL BAPHTHYS. NAPHTHYL-BENEAMIDINE.

BENZENYL-NAPHTHYLENE-DIAMINE

O11H11N2 i.e. PhC NH C11H. [210°]. From benzoyl-nitro-(\$)-naphthylamine, tin, and HCl long needles, v. e. sol. water. — AgA': white in presence of alcohol (Ebell, A. 208, 328) gelatinous pp. Yollowish crystals, sl. sol. water, v. sol. alcohol. Salts. - B'HCl. - B'HNO, - B', H, SO, -

B'C, H, 1: small needles; NaOliAq or hot water removes the isoamyl jodide. BENZENYL-OXAMIDINE is BENZENYL-AMID-

OXIM (q. v.). BENZENYL-OXIMIDAMIDE is BENZENYL. AMIDOXIM (q. v.).

BENZENYL. PHENYL-AMIDINE C. PHENYL-BENZAMIDINE.

BENZENYL-DI-PHENYL-DI-AMINE v. Di-PHENYL-BENZ-AMIDINE.

BENZENYL-o.PHENYLENE.DI.AMINE

 $C_{13}H_{16}N_2$ i.e. C_6H_3 . $C \le \frac{NH}{N} > C_6H_4$. [280°]. Anhydro benzoyl di amido-benzene. From benzoyl-o-nitro aniline, tin, and HCl (Hubner, A. 208, 302; 210, 328). Plates (from glacial HOAc). M. sol. alcohol, sl. sol. benzeno and water.

Salts .- B'HCl. -- B',H,PtCl. -- B'HI su .--

B'HNO, B', H, SO, I had.

Reactions. - 1. Benzoyl chloride at 200° has no action. - 2. Heated with cyanogen iodide and benzenc, palo yellow crystals of Ph.C \(\sigma_N(CN)\) > C.H. [106°] are formed (Howe, Am. 5, 416). - 3. Methyl iodide forms C₁₃H,MeN₂Mel, [141°], whence potash forms C₁₃H₃MeN₂MeOH [152°], insol. hot water, sl. sol. cold alcohol. It forms the follow ing crystallino salts: C₁, H₂MeN₂MeCl a₁, - (C₁, H₂MeN₂MeCl₂(2s0°), (C₁, H₂MeN₂MeSO₂) - C₁, H₂MeN₂MeSO₃H aq. -4. Ethyl iodide forms at 180° Call Etha ni. sol, water. It forms soluble crystalline salts, B'HCl and B',II,SO, -5. Ethyl todide at 210° forms C,H,EtN,EtI, [155°] whence hot potash liberates liberates the crystalline mumonium base C₁₄H₂EtN₄EtOH [132], insol, water and alkalis, si. soi. cold alcohol, v. e soi. benzene and ligroin. Its salts are crystalline and not dehgrom. Its saits are crystamine and not accomposed by NH, Na₂CO₃, or cold KOH, but hot potash liberates the base. Saits.—
C₁₃H₂EtN,EtCl 2aq. — (C₁₃H₂EtN,EtCl),PtCl₃—
C₁₄H₂EtN,EtN, — C₁₃H₃EtN,EtSO, if aq.—
6. Isocomyl boldide at 170° gives C₁₄H₃C(H₁₃)N,
Calta = 1810° — 18110° — 18110° — 18110° — Salts. - BHCl. - BHL -B'II,SO, 2aq.—7. Isoamyl indido at 165° for 24 hours gives C₁₃II₂(C₃I₁₁)N₂(C₃I₁₁)I₁ (112'). The free ammonium base, C₁₃II₂(C₄I₁₁)N₂C₅I₁₁OII [81° and 92°] crystallises from alcohol, and is insol. water. Salts .- B'C, H, Clay (and Jag) .-B'₂(C₁H₁₁)₂PtCl₂. - B'C₂H₁₁NO₂HNO₂. [90°]; - B'C₂H₁₁N. -8. Conc. nitric acid forms a nitrocompound, C12II, (NO2)N2, [196°].-9. II, SO4 forms an unstable sulphonic acid.

BENZENYL - PHENYLENE DIAMINE - p-CARBOXYLIC ACID $C_rH_i < N > C.C_rH_i$.CO.H.

Slender needles (containing I aq). Almost insol. in cold, v. sol. in hot, water. Prepared by the exidation of toluenyl-phenylene-diamine CH. NH CO.H. OH. H,80,

Salts.—BaA's 6aq: small needles, al. sol. water. - CaA', baq: slender needles. - KA' 7aq:

gelatinous pp.

Reaction. The silver salt on distillation gives a ketone 'Anhydro-tolyl-ketamine C,H,NO or

 $c_n H < \frac{N}{NH} > c_n C_n H \cdot c_n C_n H$ (277°) which forms salts II"2HCl and B"H PtCl_

Ethyl ether EtA'.—(213°); oolourless needles (Stoddard, B. 11, 293; Brückner, A. 205, 113; Hålmer, A. 210, 337).

BENZENYL - PHENYL - THIURAMIDOXIM C4H2.C(NH.CS.NHC4H2):NOH. [163°]. Formed by the combination of phenyl-thio-carblinde with benz-amidoxim (Krilger, B. 18, 1060). Plates. V. c. sol. alcohol, other, and benzeno.

BENZENYL-PHENYL-URAMIDOXIM

Cally C(NH.CO.NHC, Ha): NOH. [115°]. Formed by the combination of phenyl cyanate with benz-amidoxim (Krüger, B. 18, 1059). White plates. V. c. sol. alcohol, ether and benzsue, insol, cold water.

BENZENYL-TOLYLENE-DIAMINE

 $C_{14}H_{12}N_2$ i.e. $C_{\alpha}H_{\alpha}.C\leqslant \frac{N11}{N}>C_{\alpha}H_{\alpha}.CH_{\alpha}$ [14].

(210°). From benzoyl-nitro-p-toluidino, tin, and HCl (Hübner, A. 208, 316). Also by heating acetophenone with o-tolylene-diamine (Ladenburg a. Ragheimer, B. 12, 951). Crystalline, insol. water, v. sol. hot alcohol; may be dis-

Salts. -- B'HCl; needles; al. sol, cold water and alcohol.—B',H,SO,: needles; v. s. sol. cold water.

Reactions. - I. Methyl indide at 210° gives C₁₁H₁₁MeN₂MeI₁ [106°] whence KOH forms C₁₁H₁₁MeN₂MeOH [144°], insol. water, of which base the following salts are crystalline, viz.:

 $\begin{array}{c} C_{14}H_{11}Me_2N_2Cl\,2aq, \quad (C_{14}H_{11}Me_2N_2), PtCl_4, \cdots \\ C_{14}H_{11}Me_2N_2L + C_{14}H_{11}Me_2N_2SO_1L, \quad Nitrous \ acid. \end{array}$ produces a nitro-compound C14H10 (NO,) Me, N,OIL [165°] (Hübner, A. 210, 371) which forms & platinochloride, (C₁, H₁₀(NO₂)Mo₂N₂Cl) PtCl₄, and reduces to an amido-compound. 2. Ethyl todds at 300 gives $C_{14}H_{11}EL_N_2I_4$ [129°] whence $C_{14}H_{11}EL_N_2OH$ [163°] and its salts

 $C_{11}H_{11}Et_2N_2CHCL = (C_{14}H_{11}Et_2N_2)_2PtCl_4$,— $C_{14}H_{11}Et_2N_2L = C_{14}H_{11}Et_2N_2SO_4H$ aq.

BENZENYL - p - TOLYL - TOLYLENE - DI .

AMINE C,H. N C,C,H, [166°]. Needles of prisms. Formed by reduction of benzoyl-nitrodi-tolyl amine with tin and HCl.

Salts. -- B'H'Claq. -- B', H, SO, *; prisms (Lellmanu, B. 15, 832).

BENZENYL-XYLYLENE.DIAMINE

C,,H,,N, i.e. C,H,.C NH>C,H,Me, [414] [195°]. From benzoyl-nitro-xylidino [185°], tin. and glacial HOAc (Hubner, A. 208, 320). Needles, insol. water. Boiling fuming HNO, forms a compound, crystallising in yellow needles [2027].

Salts.-B'HCl.-B'HNO, -B',H,SO, -B'H,C,O,.

C,H,C NH O,H,Me, (\$18°). From bensoylnitro-zylidine [178°]. Needles. - B'HCl Saq (Habner, B. 10, 1711)

BENZ-ERYTHRENE Cz.H. [307°-808°] Formed in leading benzene through a red-hot tabe (Berthelot, J. 1860, 541; 1867, 599, 695; Schultz, B. 11, 95). Small leaflets. Almost insoluble in slechol, difficultly soluble in hot acetie acid and in cold benzene.

BENZ. FURIL C, H,O, i.e. Ph.CO.CO.C,H2O. [41°]. From benz-furoin and Fchling's solution at 50° (Fischer, A. 211, 229). Yellow needles. V. sol. alcohol or ether (unlike furil). Volatile. Tetra-bromide C₁₂H₂O₂Br₄. [127 -128°].

Yellow noedles.

BENZFURILIC ACID C, H, O, i.e. C,H,O.CPh(OH).CO,H. From benz-furil and squeous KOH at 60'. Prisms (from a mixturo of ether with light petroleum). Turns brown at 108°. V. sol. ether, alcohol, or chloroform; d. sol. light petroleum. Its aqueous solution gradually decomposes. The crystals in conc. LSO, give a blood-red colour, turning brown. The oily residue got on evaporating an etherenl solution of benzfurilie acid gives with cone. H.SO, a reddish violet colour, water then gives a blackish-blue pp., which dissolves in conc. H.SO, giving a blue colour (E. Fischer, A. 211, 281).

BENZ FUROIN C₁₂H₁₀O₃ i.e. 15thor Ph.CH(OH).CO.C₄H₃O or C₄H₃O.CH(OH).CO.Ph. [1870-1390]. From furfurol (18 g.), benzoic aldehyde (20 g.), alcohol (60 g.), water (80 g.), and KCN (4g.); by boiling for 20 minutes (E. Fischer, A. 211, 228, B. 13, 1339). Slender prisms (from alcohol). May be distilled. V. priams (from alcohol, biay no distinct. v. sol, hot alcohol, ohloroform, or benzene, sl. sol, water or light petrolemn. Alcoholic KOL forms a dark-red solution with hluish-green reflex.

BENZGLYCOCYAMINE C.II, N.O.. Formed

by boiling the dioyanide of m-amido-benzoic acid (v. p. 157) with KOHAq (Griess, B. I, 191; 3, 708; Z.4, 725; 6, 728) or by the netion of cyanamide on an alcoholio solution of m-amidobenzole acid containing NII, (Griess, B. 7, 575), Thin white four-sided plates (containing aq). Si. sol. hot water, v. sl. sol. ether, insol. NH, Aq, sol. aqueous mineral acids. Boiling baryta converts it into m-amido-benzoic acid, urea, NH, and uramido-benzoic seid.

Salts.—B'HCl.—B'.H.PtCl.. BENZGLYCOCYAMIDINE C.11,N.O, i.s. HN:C NH C.11 Formed by the prolonged

heating of 'ethoxy cyanamido benzoyl' (v. p. 166) with alcoholio NII, at 100°; or by the action of oyanamide on o-amido-benzoio acid Griess, B. 1, 191; 2, 415; Z. [2] 5, 574; B. 7, 574; B. 322; 13, 977). Nacrecus lamine, al. sol. water, sl. sol. holling alcohol. Its mitrate forms narrow lamine, v. sl. sol. alcohol.

Platinochloride: B'gH,PtCl. m - Carboxy - phanyl - bons - glycccyamidina IIN.C.H.,.CO

Formed from CO.H.C.H.NH.C-N

mixture of 'dicyanamido-benzoyl-'

and m-amido-benzoic acid by

heedles of plates. In Tolorably strong said.

Salts.-A"Ag,: white amorphous pp. A",H.Bs 10aq (Griess, B. 18, 2420) Imido-phenyl-henz-glycocyamidine NH.C,H,.CO

NH.O,H,NH [1:3]. Formed by balled. HN:C

heating cyancarhimidamidobenzoio acid with o-phenyleno diamine (Gricss, B. 18, 2414). Sixsided tables. Nearly insol. water, alcohol, and ether. It has both acid and basic properties.

Salts: A'Ba .- A'HIICl: white six-sided plates.

BENZHYDRAMIDE v. BENZOIC ALDERYDE. BENZHYDRAZOIN v. TRI-PHENYL-HYDRAZOIN. BENZHYDROL v. DIPHENYL CARBINOL

BENZHYDROLENE C, 11, [210°]. Got by distilling succinate or benzoate of benzhydrol (Linnemann, A. 133, I). is Identical with tetra-phenyi-ethanc (Zagumenny, J. R. 12,

BENZHYDROXAMIC ACID v. 11yproxyl-AMINE.

BENZHYDRYL ACETATE v. Acetyl-BENZ-TIVENEST.

BENZHYDRYLAMINE v. DI-PRENYL-CAR-DINYLAMINE

BENZHYDRYL-BENZOIC ACID v. Excoxy-DENZYL-BENZOIC ACID.

BENZHYDRYL CARBOXYLIC ACID v. Exo-OXY-DENZYL-DENZOIC ACID.

BENZHYDRYL PHENOL v. DI-OXY-DI-PHENYL-METHANE

BENZHYDRYL-iso-PHTHALIC ACID v. Oxy-DENZYL-130-PHTHALIC ACID.

BENZHYDRYL-PROPIONIC ACID v. v-oxy-

7-PRENTL-BUTTRIC ACID.
BENZHYDRYL-PROPIO-CARBOXYLIC ACID

OXY-CARBOXY-PHENYL-PROPIONIC ACID. BENZIDINE and derivatives v. DI-AMIDO-

DIPHENYL and derivatives.

C.H. CO.CO.C.H. + 2HCl (Laurent, A. Ch. [2] 59, 402) .- 2. By oxidising benzoin with nitric acid (Zinin, A. 34, 188), and in small quantity by the air-oxidation of benzoin dissolved in alcoholic potash $C_{11}H_{12}O_2 + O = C_{12}H_{10}O_2 + H_2O$ (Limpricht a. Schwanert, B. 4, 335).—3. Together with stilbene by heating stilbene dibromide with water to 150°: 3C,H,.CHBr.CHBr C,H, + 2H,O = C.H., CO.CO.C.H., + 2C.H., CH:CII.C.H., + 6HBr (I. a. S., A. 145, 338).—4. Together with tolane by heating tolane-dibromide with water to 2000:

2C,H,,CBr:CBr.C,H, +2H,O =
C₁₄H₁₅O₂+C₄H₃.C:C.C₄H₃+4HBr
(L. a. S., B. 4, 380).—5. From tolane-tetrachloride by heating it with glacial acetic acid

chloride by heating it with gladial acetic acrd or with cone. H.SO, to 165°:

C₁H.,CCl.,CCl.,C₂H., +2H.SO. =

C₁E.,O., +2SO., +4HCl

(Liebermann a. Homeyer, B. 12, 1975).—6. By heating benzoyl chloride with sodium amalgama;

2C.H.,CCCl. +Na₂=C₁H.₁₆O₂ +2NaCl

(Klinges, P. 18, 2031).

B. 16, 996). Preparation.-Benzoin is heated with twice

Properties .- Crystallises from hot alcohol n transparent yellow needles; from ether, by pontaneous evaporation, in large six-sided risms. Insol. water, v. sol. hot alcohol and ether. The crystals are optically active, showing sometimes dextro-rotation, sometimes lavo-rotation, but having no hemiliedral faces. Inactive in a fused state or in solution (Descloiseaux, C. C. 1870, 418).

Reactions .- 1. Benzil is readily attacked by reducing agents. Iron filings and acetic acid, or sine and hydrochloric acid, acting upon the alcoholic solution, convert it into benzoin (Zinin, A. 119, 177). When heated with alcoholic potas-sium hydroculphide at 120° it yields a mixture of deoxybenzom (C, II , CII, CO, C, II,) and benzom (Jens, A. 155, 87). Sodium amalgam and water convert it into hydrolonzom (Zincke a. Forst, B. 8, 797). When a solution of benzil in moist ether is exposed in a scaled tube to simlight the benzil is reduced to benzil-benzoin C, II , O, [131 '-135"], which separates in resettes of white or pule yellow crystals on the sides of the tube, whilst the ether is converted into aldehyde. On melting benzil-benzoin, or on attempting to recrystallise it from any of the usual solvents, it hreaks up into benzil and benzoin : C, Il, O, w $2\mathbf{C}_{11}\mathbf{H}_{10}\mathbf{O}_{2} + \mathbf{C}_{11}\mathbf{H}_{11}\mathbf{O}_{2}$ (Klinger, B. 19, 1864). 2. Benzil is very stable towards accls. It may be boiled with furning hydrochloric acid without undergoing change; cone, sulphuric acid dissolves it, and the addition of water precipitates it unaltered; ordinary strong nitric acid does not attack it. On boiling it with fuening nitric acid, however, it yields a mixture of two dinitrebenzils, $C_{14}H_1(NO_d)O_2$, which may be separated by crystallisation; octahedcal crystals [1312], sl. sol. alcohol; and lamine (1170), less soluble than the foregoing (Zagumenny, J. R. 4, 278). An isodinitrobenzil [205°] has been indirectly prepared by the exidation of aor y- dinitro-deoxybenzoin; v. el. sol. alcohol, m. sol. boiling benzene or glacial acctic acid m. sol. both of the control of partial accept and (Goldbeff, J. R. 13, 29). A mononitobenzil, C.,H.(NO.)O. [110°), has also been indirectly obtained by boiling I part of deoxy-benzoin with 8 parts of nitric acid (S.G. 1-2), or by adding benzoin to nitric acid (S.G. 1-2) cooled to 0°. Yellow lamine or flat needles; ni. sol. alcohol, mora readily in other. Hot alcoholic

potash decomposes it into potassium mazo-potash decomposes it into potassium mazo-potassium mazo-enzoate and potassium maxybenzoate: 2C₁,H₁,NO₂|Q₁+4KII) = N₂(C,H₂,CO₂Kl₁+2C,H₂KO₂+2II,O. Tin and hydrochloric acid convert it into another deoxybenzoin, C14H11(NII2)O (Zinin, A. Suppl. 3, 153).-8. When heated with soda-lime, benzil yields benzene and benzophenone, CO(C,H,), (Jens. A. 155, 87); distilled over heated litharge it yields benzophenone (Wittenberg a. V. Meyer, B. 16, 501).—4. When benzil is heated with an squal weight of caustic potash or caustic soda dissolved in 20 parts of water, the benzil slowly dissolves, forming a benzilate (v. BENZILIC ACID): Q.H., CO.CO.C.H. + KOH = (C.H.).C(OH).COOK. amall quantity of diphenyl-carbinol is formed

heating, the colour is discharged and the solution contains potassium benzilate. According to Klinger (loc. cit.) aqueous potash gives this colour-Ation only with benzoin, and then only when air admitted. When benzil is dissolved, without beating in personal linear transfer or the second ating, in very dilute alcoholic potash (4 grams KOH to the litre) and allowed to stand, with exclusion of air, for about a fortnight, it is converted into a mixture of two compounds, C. 11, 0 [200° 201°] (Limprieht a. Schwanert's so-called chipt-dibensets) and C₄.H₄,O₄ (232). The alco-hol takes part in the formation of these compounds: 2C₁H₁₀O₂+C₁H₂O₃-C₂₀H₂O₄+H₁O₇+H₁O₇ and 3C₁₄H₁₀O₇+2C₂H₂O C₁₆H₂O₇+4H₁O. Thase two substances can be separated by recrystallisation from alcohol, in which the compound CasH24O, is the more soluble. The compound C. II.O, is sl. sol. boiling alcohol, sl. sol. boiling benzene, v. sol. boiling glacial acetic acid; deposited from alcohol in colourless granular crystals with 1 mol. of alcohol of crystallisation which is expelled at 120° but not at 100° crystallises also with 1 mol. of acetic acid and with benzene of crystallisation. pound Calls,O, is finally purified by dirsolving in boiling phenol and precipitating with alcohol; yellow crystalline powder, v. sl. sol. alcohol, n. sol. boiling pheno (Japp a. Owens, C. J. 47, 90; cf. also Limpricht a. Schwanert, B. 4. 385). Benzil also reacts with isopropyl alcohol: when it is dissolved in a dilute volution of oaustic potash in isopropyl alcohol (2:100) and allowed to stand for some months in a stoppered vessel a compound $C_{\rm M}H_{20}$ ($[147^2-148^2]$ is formed; $2C_{\rm c}H_{16}O_{\rm c}+C_{\rm B}H_{20}O_{\rm c}+O_{\rm c}$ a reduction taking place, $C_{\rm M}H_{20}O_{\rm c}$ forms very lustrous erystals, of piombohedral liabit, v. sl. sol. builing alcohol (Japp a. Raschen, C. J. 49, 832). -5. Renzil reacts with acctone in presence of caustic potash, yielding the compounds a cetoue-banzil, C₁₁H₁O₂; anhydracctone beuzil, C₂H₁O₂; and anhydracetone dibenzil, C₂₁H₂O₄ (v. Acetone-denzil, p. 32). With methyl ethyl ketone it yields methyl anhydracetons hanzil, C., H., O.: colourless thin prisms [1799]; with diethyl ketone, dimothyl-anhydrace tone-banzil, Calla O,: rhomboidal platet [150°]; with methyl propyl ketone, a thyl anbydracotone-benzil, Call, Oz: needles [156]; with methyl heryl ketone, umyl-anby dracetone benzil, C,H,O,: silky needlei [150.5°]; all sol. alcohol (Japp a. Burton, C. J. 5], 431),—6. Benzil reacter in ammonic, and also with aldehydes and ammonia jointly, to form a number of well-characterised conclenas. tion-compounds (r. Benzil, ammonia-deriva tives or). Heated with ariline in a scaled tube at 200 it yields benzil aniiule, C.H., C(N.C.H.).CO. C.H. [105°]. Yellow prisms, sol. alcohol (Volgt, J pr. [2] 84, 23).—7. Warmed with (1,2,4)-tolylene diamine in alcoholic solution it forms diphenel toluquinoraline: C.H.CO H.N.

C.H.CH. C.H.C.N.C.H.CH.+#H.O. CH CO HA (Hinsberg, B. 17, 322). Benzil also reacts with ethylene-diamins to yield a crystalline compound (Mason, B. 19, 113). -- 8. Forms compounds Benefit with free laydroxylamine in aqueousalcoholic solution at the ordinary temperature yields beneil-mono-owim C.H., C(N.OH).CO.C.H.,

[130°-131°], even when an excess of hydroxylamine is employed. Small square white leaf-lets, v. sol. alcohol and ether, el. eol. water

(Wittenberg a. V. Meyer, B. 16, 503). Boiled with hydroxylamine hydrochloride in methy sloohol, with the addition of one drop of hydrochloric acid, a dioxim, (a) diphenyl-glyoxim C.H., C(N.OH).C(N.OH).C.H., [237°] separates.

Forms lustrous white lamine, el. sol. methyl algohol, alcohol, and other. Dissolves in conc. caustic sods and is reppd. by acids. Sol. with difficulty in ammonia, the solution giving with silver nitrate a yellow pp. (Goldschmidt a. V. Meyer, B. 16, 1617). It benzil, hydroxylamine hydrochloride, and alcohol, acidulated with hy-

drochlorio acid, are heated in a scaled tube for several hours at 170°, an isomeric (B)-diphenylglyoxim [206"] is formed. White needles, v. sol. boiling alcohol, sl. sol. other and boiling water. Sol. caustic soda and ammonia. The (a)-compound can be converted into the (B)-com-

pound by heating it with alcohol in a sealed tube at 180°. The nature of this isomerism is not understood; but analogous cases of isomeriem have been observed in the benzil and benzoin groups - thus benzil and isobenzil, hydrobenzoin and isohydrobenzoin (Goldschmidt, B. 16, 2177). By heating benzil with phenylhydrazine on the water bath, benzil-di-phenyl-

hydrazide, C.H. C(N,H.C.H.).C(N,H.C.H.).C. [225°], separates. Faintly yollow needles; v. sol, hot chloroform and benzene, al. sol, alcohol and ether. Gives a dark-violet colouration with cone, sulphuric acid. Does not regenerate phonylhydrazine when heated with etrong hydrochloric acid (Pickel, A. 232, 230).-. When

benzil in alcoholic solution is mixed with an excess of nearly anhydrous hydrocyanic acid and allowed to stand, large, colourless, tabular, rhombic orystale of benzit dihydrocyavide C.H..C(OH).CN are deposited (Zinin, A. 34,

C.H. C(OH). CN
189). The same compound is formed when an ethereal solution of benzil is mixed with powdered potassium oyanide and cone, hydrochloric acid is added drop by drop, cooling during the

aporation. Melts at 1329, with decomposition into benzil and hydrocyanic acid. Insol. water and benzene, v. sol. ether and light petroleum. When dissolved in alcohol it is decomposed, even in the cold-time benzil and hydroganic acid (Jacoby, B. 19, 1519). When finely powdered benzil dihydrocyanide is mixed with a large excess of a concentrated solution of IIBr in glacial acctic acid and left for some weeks, a solution is obtained which by enontaneous evapora-

tion deposits lustroue crystals of di-phenyl-tartra-O_aH_a.C(OH).CONH_a.HBr mide hydrobromide C.H.C(OH).CONIL [186°] and these, when decomposed by ammo-nia, yield the free di-phenyl-tartramide. V. sol.

anhydrous hydrosympis and one been added, is saturated with gaseons HOI and allowed to stand for some weeks, an acid, O_1H_1 ; NO_4 (colourless priams, sol. boiling water, melting at 196°), and a compound, $O_{18}H_{12}N_2O$ [196°–197°], are formed. The compound $O_{18}H_{12}N_2O$ forms lustrous pale yellow lamings or flat needles, sl. eol. boiling

water and benzene, v. sol. boiling alcohol. It is feebly basic and forms with hydrochlorio acid a colourless salt which is stable only in presence

of excess of acid (Japp a. Miller, C. J. 51, 29). When benzil is heated with absolute alcohol and hydrocyanic acid at 200° for four hours, it is decomposed into benzoic aldehyde and ethylic benzoate: C_aH_a .CO.CO. $C_aH_a + C_aH_a$.OH = Calla, CHO + CaHa, CO2, C2Ha, the hydrocyanic acid apparently undergoing no change. Benzoio acid ie also formed (Michael a. Palmer, Am. 7. 191). Benzil and alcohol may also be made to

react to form benzoio aldehyde and ethylic henzoate by triturating benzil and potassium cyanide with alcohol; but in this case a portion of the benzaldchyde is further changed into benzoin by the action of the potassium ovanide. fly neutralieing with eodium carbonate the small

quantity of benzoic acid which is also formed in thie reaction, and which would otherwise decompose the potassium cyanide, 1 part of the cyanide may be made to induce the foregoing change in 50 parts of benzil (Jourdan, B. 16, 658). -10. Benzil unites with nitriles in presence of conc. sulphurio aoid, taking up the elements of water. Thus whou powdered benzil (1 mol.) is suspended in conc. sulphuric acid, and propionitrile 22 mols.) is gradually added, cooling during the process, a compound $C_{20}H_{\perp}N_{\perp}O_{\parallel}\{197^3\}$ is formed: $C_{11}H_{10}O_{\perp}+2C_{\parallel}H_{2}N_{\perp}+H_{\parallel}O=C_{20}H_{\perp}N_{\perp}O_{\parallel}$. Lus-

trous, colourless needles; v. sol. hot alcohol. Boiling with dilute solphuric acid decomposes it into benzil, propionic acid, and ammonia.~ With benzonitrile and sulphuric acid benzil yields a mixture of two compounds which can be readily separated by means of their very different solubilities in hot alcohol: Cz.H.zN2Oz [168°], analogous to the propionitrile derivative, v. sol. boiling alcohol, orystallises from alcohol in efflorescent oblique prisms of the formula

(C. II., N.O.), EtOII; and a compound C., II., NO., [225], almost insol. alcohol, v. col. boiling phenol, cl. col. boiling benzene, which crystalliecs from benzene in microscopie, flat rectangular prisms: $2C_1H_{10}O_2 + C_1H_1N + H_1O = C_2H_2NO_3 + C_4H_4CO_2H$ (Japp a. Tresidder, B. 16, 2652). —11. Heated with comowhat more than the equivalent quantity of phospherus penta-chloride, benzil yielde dichloroleoxybenzoin (chlorobenzil) C.H., CCl., CO.C., H., (71°). Short, thick rhombio prisms, insol. water, v. sol. ctber, not so readily in alcohol. When heated with alcohol or water to 180 it yields henzil and hydroohloric acid. Zine and hydrochloric acid, acting

duce it first to deoxybenzoin, and finally to etil-bene. PCl, at 200° replaces the remaining oxygen atom by Cl., yielding tolane tetrachloride, C1, 11, Cl. (Zinin, A. 119, 177; J. 1880, 614; A. 149, 374). Ischeaul.-1. A substance having the same composition as benzil is obtained by acting with holls solution of benzil, to which an excess of sodium amalgam on a solution of benzoyl

on an alcoholic solution, convert the compound into deoxybenzoin; acetic acid and zinc-dust re-

alcohol, v. sl. sol. ether (Burton, B. 16, 2232). Pure benzil dihydrocyanide ie not hydrolysed by conc. hydrochlorio acid, but when an alcochloride in anhydrous either (Brigel, A. 185, 173). Ordinary benzil is formed at the same time (Klinger, B. 16, 995). Isobenzil forms colourless tabular monoclinic crystals, also Instrous lamine or needles [145°-150°], sol. alcohol, ether and CS... It is probably a polymeride of benzil, for when treated with bromiue it yields benzil (I mol.) and benzoyl bromide (2 mols.): $C_{\mu}H_{\mu}O_{+}+Hr_{\nu}=C_{1}H_{\mu}O_{+}+2C_{1}H_{\nu}COBr$. Heated with alcoholic potash it gives the violet colouration of benzil and is converted into benzilio acid together with a small quantity of benzoic acid (Klinger, loc. cit.; also B. 19, 1862).—2. By heating benzoic allehyde with sodimu amalgam in an atmosphere of CO_{μ} Alexéet (A. 129, 317) obtained an oil (314° spprox.) to which he assigned the formula $C_{1}H_{\mu}O_{\mu}$ —S.G. 12—1104 (approx.).

BENZIL, AMMONIA DERIVATIVES OF.
There are four general reactions known, according to which compounds containing the dicarbonyl-group—CO.CO—form condensation-compounds with aldehydes and ammonia jointly. As a knowledge of these reactions is necessary to an understanding not only of the behaviour of benzil with aldehydes and ammonia, but also if that of benzil with aumonia alone, the general equations for these reactions will be introduced at this stage. In this following equations X stands for the monad hydrocarbon-radials of the dicarbonyl-compound, and R' for the monad hydrocarbon-radicle of the slidehyde:

Here the dicarbonyl-compound is broken up into two halves, whilst the two aldehydo-groups become directly united. Lastly:

The constitution of the compounds of the last-montioned type is unknown (Japp a. Streatfeild, C. J. 49, 165; Japp a. Hooker, C. J. 45, 673; Japp a. Wynne, C. J. 49, 464).

It will be shown later on, that in the reactions of benzil with ammonia a part of the benzil is first broken up with formation of benzoic acid and benzoic aldebyde, which latter then takes part, together with benzil and ammonia, in the final reaction. The benzilammonia reactions are therefore in reality Tok.

benzil-aldebyde-ammonia relations; and will be more readily understood if the reactions of the latter class are described first.

REACTIONS OF BENEIL WITH ALDERTOES AND AMONIA:—

Equation I. (vide supra).—Bensil gives no pactions according to this equation so long as free aldehydes are used, but with massent benzoic aldehyde, produced by the decomposition of a portion of the benzil, it reacts according to this equation, yielding benzilam, Callano (vide infra). In the case of some other disarbonylcompounds, however, such as phenanthraquinons (q. v.), this reaction occurs with various free aldehydes.

Equation II.—Benzil reacts according to this equation with formic allehyde, acetic aldehyde, isovaleric aldehyde, glyoxal, benzuic aldehyde (free), and p-oxybenzoic aldehyde. The compounds formed are derivatives of gly-

oxaline | CH - NH CH.

C,II,.CO

Thus when bouzil and formic ablehyde are warmed with ammonia in alcoholic solution at about 40° diphenylglyoxaline is formed;

hol and ammonis is expelled by heating; the base is extracted with dilute hydrochloric acid, precipitated with ammonia, and crystalliced from hot alcohol. It separates from hot alcohol am cooling in long oblique crystals (monosymmetric), and from cold alcohol by evaporation in short restrous crystals (also monosymmetric, but not referable to the same parameters) (227°). Monucid base, (0,111,10,11Cl),pRlc1,: pale-yellow, amorphous precipitate, epeedily changing into microscopic flat needles (Japp, C. J. 51, 558)]. When acetic aldehyde is substituted for formic aldehyde in the foregoing reaction, methyldiphenylglyoxaline, C,11, C-NH

C.CH., is formed. [Orthorhombia C.H., C. N. orystals [235°], v. sol. ether and hot slochel. (C.H., N., HCl.), PtCl. (2aq. yellow microscopic needlos (Japp a. Wynne)]. The bare forms a molecular compound with diphenylglyoxaline (J.)]. Benzii, isovaleric ablehyde, and ammonia yield isobutylliphenysyncaline,

Call. C—NH

C.CH. CII(CII). [Needles

Call. C—N

[223]; sol. hot benzene and alcohol.

(Call. N,HCl). PtCl.: amorphous yellowish

brown precipitate, or small crystals (J. a. W.).]

Glyozal, as a dialdehyde, reacts with twice the

proportion of benzil and ammonia, forming

tetraphenylolycosine,

O.H. C—NH

NII—C.C.H.

O.H. C.—N. S.—C.C.H. alcohol in tuits of silky needles of the formula C_{2s}H₂N_{*}, C.H₂O. Melts above 300°. Feebly basic (Japp a. Cleminshaw, C. J. 51, 553)]. Benzil, benzoic aldehyde, and ammonia yield

HH

0.0 H. (Radziszowski) 1498). In a similar manner p bensoic aldehyde gives poxylophine, C.H.O-NH. C.C.H.(OH). [Tuits of colour-

O.H. O--n/ less needles (254°-255°), v. sol. hot alcohol, eol. caustio soda, forming a sodium compound. Heated with acetic anhydride, it forms a monacetyl - derivative C2,H1,3(C2H2O)N2O [229°]; needlee, with a faint eatiny lustro, eol. hot alcohol. Dietilled with zinc dust, p-oxylophino is converted into lophine (Japp a. Robinson, C. J. 41, 826)1.

Equation III. - Illustrations of thle equation are to be found in the reactions of salicylic aldshyde and furfuraldshyde (pyromucio aldebyde) with benzil and ammonia. Cinnamic aldshyde also reacts according to this equation ; but in this case another reaction, according to Equation IV., occurs simultaneously.

Thus when equal weights of ealicylic aldehyde and bonzil are dissolved in warm aloohol and the liquid is saturated with gaseous ammonia, the condonsation compound dibenzoyldioxystilbene diamine separates as a crystalline powder:

20,H4(OH).CHO+ | CO.C.H. +2NH,= C.H.(OH).CH.NH.CO.C.II, + 2H₂O.

C.H.(CH).CH.NH.CO.C.H. It is purified by dissolving it in boiling phonol and procipitating with alcohol. Microscopio plates, molting with decomposition above 300°, insol. in the ordinary organic solvents, sol. boiling phenol, sol. oaustic soda, forming a sodium compound. By fusion with caustic soda it yields sodium benzoate and sodium salicylate. Heated with dilute hydrochloric acid at 210° it is hydralysed into benzoio acid and dioxystilbene-

C.II.(CII).CII.NIf. diamine ; small lustrous

C.H. (OH).CH.NII. base: the Pt salt, C, H, N, C, 2HCl, PtCl, 4sq forms thick, orango-coloured, rhomboidal plates with bevelled edges, anbydrous at 100°. This base is, however, more readily obtained from its acetyl-derivative (in/ra). When the condonantion-compound is boiled with acetic anhydride antil it dissoftes with acetic antilone. C.H.(C.C.H.O).CH.NH(C.H.O)

diamine

C.H.(O.C.H.O).OH.NH(C.II,O) brmed (rhomboidal lamine [225 -227°], sol. sectic soid); and by boiling this compound for light hours with acetic anhydride the bensoyl-groups are replaced by acetyl-groups rielding diacetyl-diacetoxystilbens-diamine rielding diacetyl diacetoxysssocial C.H.(O.O.H.O).CH.NH(O.H.O)

C.H.(O.C.H.O),OH.NH(C.H.O)

\$16°-219°] sol. glacial acetic acid and alcohol, eposited from latter solvent with 1 mol. of constallisation. This compound is settle-acetyl derivative of the above-mentioned are. By the action of caustic alkali or conc.

Broatstand bein on this teles-acetyl someon the four sactyl-groups may be removed in suc-cessive pairs, yielding first diacetyl-diacey C,H,(OH).CH.NH(C,H,O)

stilbene-diamine C.H.(OH).CH.NH(C.H.O)

talline powder melting above 300°, sol. bet plienel, sol. caustic alkalie), and finally dioxystilbene diamine. The latter base ie most conveniently prepared by heating the tetra-acetyl compound with cone. hydrochlario acid at 120°. By heating the condensation-compound with benzoic anhydride a dibenzoyl derivative, corresponding with the diacetyl derivative is obtained; it is a tetra-benzoyl derivative C,H,(C.C,H,C).CH.NH(C,H,O)

of the base, thus:

C.H.(C.C.H.C).CH.NH(C,H.O) Microscopio platee [246°-248°], sol. acetic acid; sol. dilute caustio potash on long boiling, regenerating the condensation-compound. These various acetyl and benzeyl derivatives may alsn be synthesised from dioxystilbene diamine by treatment with acetic anhydride and benzoic anhydride. The dibonzoyl derivative thus prepared is identical with the original condensationcompound (Japp a. Hooker)].

Benzil, furfuraldehyde, and ammonia also

react according to Equation III. : $C_{t_1}H_{t_2}C_{t_2}+2C_1H_tO_1+2NH_3=C_2H_{t_2}N_1O_4+2H_2O_1$ forming, however, two isomoric compounds of the formula $O_{t_1}H_{t_2}N_2O_4$. One of these is an analogue of the salicylio aldohydo compound; it is separated by means of its insolubility in alcohol, and purified by dissolving in boiling phonol and precipitating with alcohol. Cryetalline powder, v. sol. het phenel, sl. sol. glacial acetic acid. The isomeric compound orystallises from alcohol in tufts of silky needles [246°] (J. a. H.).

When cinnamic aldehyde, benzil, and ammonin are allowed to react in alcoholic solution. a mixture of two compounds is obtained. Boiling alcohol extracts one of these, and the remaining compound, which is insoluble in alcohol, is purified by dissolving in hot phenol and precipitating with alcohol. The compound soluble in alcohol is cinnimabenzil, C₃,H₃₀N₃C₃ (infra). The compound insoluble in alcohol is dibenzoyl-dicinnamylene-diamine and is formed according to Equation III.:

C,II,.CO 2C,II,.CII:CII.CHO+ + 2NH. = C_11,.CO C.H.CH.CH.CH.NII.CO.C.II, + 2H.O.

C.H., CH:CH.CH.NH.CO.C.H. Crystalline powder, consisting of short microecopio prisms [264°], insol, in the nrdinary organio menstrua, sol, hot plucnol. When heated with a solution of potash in methyl aloobol at 150 it parts with the elements of benzoic acid and convorted into bensenyl-dicinnamylene-diamine C.H.,CH:CH.CH-NH

C.C.H. Silky needles, _ N// C.H.,CH:CH.CH. [223°], sol. benzene and alcohol. Monacid base; Pt salt, (C,H,N,HCl),PtCl, 2aq, forms ailki needles.

Equation IV.—The above-mentioned soles product of the reaction of cinnamic algebra

C.H. CO

C.H., CO. C., H., N.O. + 2NH, = C.H., CO. C., H., N.O. + 2H., O. Crystallises from hot alcohel in two forms—stender needles and minnts short prisus [188], sol. benzens. A selotion of potash in methyl alcohol, when allowed to act upon it in the cold, forms potassium benzeate and cinnidimatersil, C., H., N.O.;

C₁,H₂,N₁O₂ + KOII = C₁,H₂,N₁O₂ + O₄H₃,COOK.

Crystalline powder [283°], sol. het plenel, precipitated by alcohol. By beiling einnimabenzil with dilete sulphuric ecid it is hydrolyaed, yielding benzilimide (infra), einnamie aldohyde, benzole acid, and ammonia: C₂,H₂,N₂O₂ + 21L₂O = O₁,H₃,N_O₃ + C₄H₄O + C₅H₄O₅ + NH₅ (Japp and Benzilimide Wynne).

RECTIONS OF BENZIL WITH AMMONIA ALONE,— Lannant, Revue Scient 10, 122; 19, 440; Zinin, A. 34, 190; Zincke, B. 16, 890; Japp, B. 16, 2638; Henius, A. 228, 339; Japp a. Wynne, G. J. 49, 473.

By heating benzil with alcehelie ammonia the following compounds are obtained: ionabunzil. C., fi.[1,N,0.; benziliamide, O., II, N.O.; benziliam, C., II, N.O.; and lophine, C., II, N.O. At the same time benzele acid, cthylic benzeato, and benzemide are fermed. This reaction was first studied by Laurent, whe prepared tha first three of the above moetiened compounds, ascribing to them, however, incorrect formula.

Benzil is dissolved in alcohol so that the solution is saterated at 40°; gaseous ammonia is passed into the warm liquid to saturation, and tha whola is allowed to stand for 24 hours. Prismatic crystals of inabenzil are deposited, whilst benzilinide, benzilam, and tha ether compounds above onumerated remain in selution. If slender acicular crystals of benzilimide should separate they may be removed by warming with alcohol, in which imabenzil is seluble only with difficulty. The formation of imabenzil may be arpressed by the equation: 3C₁₄II₁₆O₂ + 2NH₂ = C₃₄H_{2N}O₂ + C₄II₄COOH + II₄O. Imabenzil forms small instrous ortherhombic prisms [194°], al. sol. het alcohol, decomposing on long boiling, and yielding ameng other products benzilinide; the best solvent is het methyl alcohol. Beiled with dilute sulphuric acid (1 vel. acid: 2 vels. water) it is converted into benzilinide, benzil, and ammonia:

C_mH_mN₂O₃ + H₂O = O₃|H₁, NO₂ + C₁, H₁₀O₅ + NH₂; whilst cold conc. sulphuric acid dissolves it, experting it into benzilam, benzaldahyda, benzols acid, and ammonia:

C₁,H₂,N,O₂ + H₂O₃

C₂,H₃,NO + C,H₂O₄ + NH₃

(Japp a. Wynna). The fermatien of bensilam in this reaction was first observed by Laurent. Boiling with alcoholic polash converts imabenzil into benzilimide (Laurent). Acetic acid and acetic arhydrids act like dilute su'phuric acid, decomposing it en boiling with formation of assistimide and benzil (Henius). Hented for some time to 140° it decomposes and melts, haming benzilimide, benzilam, and lophine, whilst an odour of benzols aldshyde is percepticated.

The steakelle transcripted mether liquer from the preparation of imabonall yields, when concentrated, a mixture of benefitmide and besigner. A similar mixture is obtained by heating behall with alcoholic ammenia for some hours at 100°, the imabenzil which is first formed being converted into benzilmide and benzilam; at 130° lophine is also formed (Henius). The fermation of lephine occurs according to the equation 2C, H., O. + 2Nii. =

equation 2C₁,H₁₀O₂ + 2Nli, =
C₁,H₁,N₂ + C₁I₁,COOH + 2H₂O.

Benzilimida and benzilam are best separated by beiling the mizture with light petroleum, which extracts the whole of the benzilam, depositing it en cooling, in resettes of prisms, and hardly dissolves the benzilimide, which may be purified by crystallisation from hot alcohol. Benzilimide is formed from benzil and ammonia according to the equation—

20, H₁₀O₂ + NI₁ = C₂₁H₁,NO₂ + O₂I₁,COOH.

Tufts of silky needlos [137°-189°] (1f.), sol. het alcohol. Concentrated sulphuric acid dissolves it in the cell, abstracting the elements of water, and converting it into benzilam (L₁)_w, C₁(I₁,NO₂ - I₁O = O₂I₁,NO. Heating with acctic anhydride produces the same effect (H₂). Chromic mixture exidises it to benzelo acid (If.). Benzilimide may also be prepared from imabenzil by Laurent's method of boiling it with alcoholic potash.

Beneilam (C.,11, NO) may be obtained as above, along with benzilimide, and separated from it as already described, or it may be obtained either from imabenzil or from bonsilimide, by the action of cold cone, sulphurle acid. The solution of imabenzil in the cold acid is poured into water, when benziliam separates and may be purified by recrystallisation from aicohol. Mestrea ily obtained by heating benzil with fased ammonium acotate in a flask ever a flame until the ammonium salt is relatilised (Jappa, Wilson, C. J. 49, 829, footnete); but as some lophina is formed at the same time, the benzilam must be extracted by means of hot light petroleum, in which tha lephina is practically inscluble. The formation of benzilam from benzil and ammonia may be expressed thus:

2C, 11, 02 + NH2 = C, 11, NO + C, 11, COOH + H, O. Well-daveleped rhombic prisms (from a mixture)

2C₁H₁O₂ + NII₂ = C₁H₁NO + C₂I₁COOII + H₂O. Well-daveleped rhombic prisms (from a mixture of ether and alcohol by spontaneous evaporation (H.)); colourless when pure; thin lustrous lamine, sometimes iridescent, (from het aloohol); rosettes of prisms (from het light petroleum). [113"-114"]. Distils at a high temperature without decompositir (L.,. V.D. (air 1) 10-23; calculated 10-28 (J.). With nitric acidit yields a mono-nitro derivative (needles: [178"-182"] from lenzene) and a dinitro-dark vative (II.). Chromic mixturs oxidises it to benzoie acid (H.).

The above reactions of benzil with ammonias may be explained as follows. In the first place, a pertion of the benzil is broken up according to the following equations:

(a) O.II..CO.CO.C.H., + H.O = C.H..COOH + C.H..CHO;
Bensoio scid Bensoio sidehyde

(b) C,H,CO.CO.C,H,+EtOH= C,H,COOEt+C,H,CHO; Ethylic beasonte

(a) C.H., CO.CO.O.H. + NH. = O.H., CO.NH. + O.H., CHO. Bensamide

These three compounds-benzoic acid, ethylio benzeats, and benzamide-are always for fred in the reaction of benzil with alcoholic ammonia. (The benzamido may also be regarded as having been formed in a secondary reaction, from ethylic benzoate and ammonia.) benzoic aldohyde, which is the by-product in every case, then reacts, in the nascent state, with benzil and ammonia according to one or other of the following equations:

$$\begin{array}{c} C_{0}H_{s}.CO \\ (d) & | & + C_{0}H_{s}.CHO + NH_{s} = \\ C_{s}H_{s}.C - O \\ | & \\ C_{s}H_{s}.C - O | \\ C_{s}H_{s}.C - NH_{s} + H_{s}O. \end{array}$$

$$\begin{array}{c} C_{s}H_{s}.C + O \\ C_{s}H_{s}.C - O | & \\ C_{s}H_{s}.C - NH_{s} + H_{s}O. \end{array}$$

This reaction of an aldohyds with a dicarbonylcompound and ammonia, in which benzilimide is produced, is not known to occur in the ease of free aldchydes.

If the reaction occurs according to Equation I. of the general reactions, benzilam is formed:

These formulæ for honzilimide and honzilam account for the readiness with which the former is converted into the latter hy the action of dehydrating agents.

If the reaction occurs according to Equation II., lophine is produced:

Finally, if it occurs according to Equation IV. the product is imabenail:

But free benzoic aldehyde, with benzil and ammonia, yields only lophine.

The foregoing equations express the formation of the various compounde obtained from benzil with alcoholic amuionia. The reactions in which the complex compounds discovered by Lanrent are formed thus really occur in two stages, of which the first consists in the formation of benzoic aldehyde, the second in a benzilaldshyde-ammonia condensation (Japp, B. 16, 2686; Japp a. Wynne, C. J. 49, 477).

BENZILIC ACID C₁₄H₁₃O₂ = (C₄H₁₃O₄ = Diphenylglycollic acid. [160°] (Jena). BENZILIC ACID C_{1,H1,O2} — contacts (J.).

H₁,C(OH).COOH. Diphenylglycollic acid.

O'] (Jena).

Formation.—1. By warming benzil with ethyl alcohol is saturated with gaseous HCl and

alceholic potash: C.H., CO.CO.C.H., *KOH=(C.H.), CO(CH.), COOK (Liebig, A. 25, 25; Zinin A. 31, 329).—2. By boiling diphenyl-bromo-acetic acid, (C.H.), CBr. COOH (obtained by passing the vapour of bromine over heated diphenyl-acetic acid), with baryta water (Symons a. Zincke, A. 171, 1721). 171, 131).

Preparation .- 1. Benzil is added to five times its weight of metted potash to which a little water has previously heen added. The whols solidifies, cwing to the formation of potassium benzilate (E. Fischer, B. 14, 326 footnote). The mass is dissolved in water, and the benzilic acid is precipitated by hydrochloric acid and recrystal. lised from hoiling water .- 2. It can also be prepared from henzoin. 15 g. benzein, 20 g. KOH and from 250 to 300 c.o. water are heated in a current of air until everything has dissolved. The solution is extracted with other to remove a small quantity of diphenyl-carbinol which is formed by the decomposition of the benzilio acid, and solid caustic potash is added. This causes the separation of nacreous lamina of potassium benzilate, which are removed by filtration, washed with a solution of caustic potash. and finally decomposed with sulphurio acid (Klinger, B. 19, 1868).

Properties. - Small white monoclinic needles with a satiny lustre. Heated above its meltingpoint it turns red. Conc. H.SO, colours it deep red; the colour disappears on the addition of water. V. sol. alcohol, ether, and boiling water;

sl. sol. cold water. Bitter taste.

Reactions.—1. Heated for several hours to 180° it yields a deep-red liquid, and, on cooling, solidifies to an amorphous mass, from which, by treatment with alcohol, dibenzilic acid G₂H₂O₃ [196°], benzophenono, and other products can be isolated (Jena).—2. Chromic mixture oxidises it to benzophenono: (C₂H₁),C(OH).COOH + O =

(C,H,),CO + CO, + H,O. Benzophenono is also produced when silver

benzilate is heated, either alone or with water (C₄H₃).CH.COOH + I₁ + I₁.O.

Zine and hydrochloric acid, and sodium amai-

Zinc and hydrochloric scid, and sodium amai-gam, are without action $(J_1, -k_2)$. Barium benzi-late, distilled with $\frac{1}{10}$ th of its weight of soda-lime, yields diphonyl-carbinol: $(C_2H_3)_2C(0H).COOH = (C_2H_3)_2CH.OH + CO_3$ $(J_1, -k_2)_2C(0H).COOH = (C_2H_3)_2CH.OH + CO_3$ chloride, $C_4H_4O_2Cl_1$ a heavy, colourless liquid (270^9) , which in contact with moist air is rapidly decomposed into housilis and hydrochloric acids decomposed into benzilie and hydrochloric acids (Cahours, A. 70, 46).

Salts .- Beazilio acid is monebasic. C₁,H₁,O₂,K: v. sol. crystallins salt (Zinin).— (C₁,H₁,O₂),Ba6aq: v. sol. crystallins crusts with a fatty lustre (J.); separates from alcohol in anhydrous needles (Symons a. Zincke).— (C.H.A.) Department of the control o (C,H,O,).Ph: pulverulent precipitate, ohtainse hy adding lead acetate to an aqueous solution of henzilio acid; fuses on heating to a red liquid (Zinin). -C, H,O,Ag: easily decomposable pre-

on diluted with water. Not volatile without

lecomposition (I.).

Ethyl-bensilic gold, C_{1e}H_{1e}O₂, isomeric with the foregoing, is a resinous substance obtained by heating benzoln in alcoholic solution with sodium ethoxide at 150°. Scarcely soluble in potash and ammonia, save in presence of alcohol (Jens a. Limpricht, A. 155, 96).

Dibensilic acid, C.H.,O. [196°], obtained by heating benzilic acid to 180° (v. supra).

heating benzilie acid to 180° (v. supra), crystallises from alcohol in minute needles. It is an anhydride, and, by heating with water at 180°, is converted into benzilio acid (Jena, B. 2. 385). F. R. J.

BENZILIMIDE v. BENZIL, AMMONIA-BERI-VATIVES OF

BENZIMIDE v. BENZOIC ALBERTDE.

DI-BENZIMIDE OXIDE v. BENZONITRILE.

BENZIMIDO-ACETATE v. BENZONITRILE, Combination 6.

BENZIMIDO-BENZOATE v. BENZONITRILE, Reaction 5.

BENZIMIDO BUTYL - ETHER MITRILE, Combination 5.

BENZIMIDO . ETHYL . ETHER NITRILE, Combination 5.

BENZIMIDO - NAPHTHYLAMIDE NAPUTHYL-BENZAMIDINE.

BENZO-ANILINE v. AMIDO-DENZOPHENONE.

BENZG-TRI-CHLORIDE C.H.Cl., i.e. C.11, CCl. Mol. w. 195-5. (214°), S.G. 14 1-38.

Formation .- 1. From benzeyl chlorido and PCl. (Wöhler a. Liebig, A. 3, 265; Schischkoff a. Roeing, C. R. 46, 367; Limpricht, A. 134, 55; 135, 80; Bl. 1866, ii. 468).-2. From benzyli-"deno chloride by chlorination (Cahours, C. It. 56, 703) .- 3. By chlorination of toluene (Naquet, O. R. 55, 407; 56, 482).

Preparation.—By passing chlorine (3 mols.) into cold toluene (1 mol.) exposed to direct sunshine (Schramu, B. 18, 608) or into boiling

Properties .- Pungent liquid, insol. water, which slowly converts it into benzoic acid (the change is rapid at 140°). Alcohol at 130° forms benzoic ether.

Reactions. - 1. Sedium line no action. -2. Ag.O forms benzoicanhydride. - 3. Aqueous ammonia at 140° gives benzoic acid, benzamide, and benzonitrile, reacting thus: O.11,CCl, +4NH,= 3NH₄Cl+C₆H₄CN.—4. Aniline forme di-phenylbenzamidine, C₆H₄C(NC₆H₄).NHC₆H₄.—5. Reacter with aromatic bases thus: C.H.CCl. + 2C.H.NXY = C,H,CCl(C,H,NXY), + 211Cl=

C.H. OCC, H. NXY C.H. NXYCl+2HCl; whence

forms the carbinol C₄H₃C(OH)(C₄H₄NXY). The reaction requires presence of a metallic chloride or other condensing agent; it takes place most easily with tertiary bases, least readily with primary bases. The products are dyes, the primary bases giving violet, the secondary and tertiary green, coloure...6. It acts similarly on phenols. C.II, CCI, +2C, H,OII = 2HCI + O, H, CCI (C, H,OH),. The products are converted by treatment with water into carbinols of H COMPIC H O(H), the blighter relief C.H. C(OH)(C.H. OH), the alkaline salts of which are colouring matters (Doebner, A. 217, 226).—

T. Converted by heating with dry oralic acid, acid, into Ph.CO.Cl, then Into (Ph.CO)(O) 2. From hippuric sold. Urino of horses or C.H.C(OH)(C.H.OH), the alkaline salts of which

(Anschüte, A. 296, 90).—8. Copper produces, on neating, C.H. CCI, CCI, C.H. (Onufrowies, B. 17.

BENZO-CUMIDE O. PHENYL AMIDO-CUMYL KHONE.

BENZO-FURILIO ACID v. BENZFURILIO ACID. 180-BENZOGLYCOL C. H.O. i.e. C.H. (OH)2(?), [471°]. A crystalline substance formed by tha electrolyeis of a mixture of benzene, alouhol, and dilute H.SO. (Renard, C. R. 91, 175). Sol. water, alcohol, and other. Reduces Fehling's colution and ammoniacal AgNO,

Diacetyl derivative C. II. (OAc), [1219]. (300°). Insol. water, sol. alcohol and other.

DI.BENZO-HYDROQUINONE v. DI-PHENTE DI-OXY PHENYLENE DI-KETOKE,

BENZOIC ACID O.H. O. i.s. C.H. CO.H. Mol. w. 122. [12149] (Schiff). (249'2 cor.) (Kopp. A. 94, 303). S.G. 21 1'20 (Mendeldeff); 1.837 (Rüdorff, B. 12, 250); 1.292 (Schröder, B. 12, 562). S. 156 at 0° (Ost, J. pr. [2] 17, 232); 172 at 0°; 207 at 10°; 425 at 81°; 178 at 75° (Bourgoin, J. Ph. [4] 30, 488). S. (ether) 66 S. (alcollo) 47 at 15° (Bourgoin, Bl. [2]
29, 245). H.F. 94,533 (Stohmann, J. pr. [2]
36, 2). S.V. 126 (Ramsay). S.V.S. 112-09
(S.). R.S. 54-24 (in a 6 p.c. benzeno solution, Kanomikoff).

Occurrence.- In various resins, e.g. gnm benzoin, dragon's blood, storax, and balsams of Peru and Tolu (Blaiso de Vigenère, Traits du feu et du sel, 1608; Liebig a. Wöhler, A. 3, 249). In castoreum (Wöhler, A. 67, 360), in tha spindle-tree (Euonymus europæus). In putrid nrino (Liebig, A. 50, 168). In cranberries (Loew, J. pr. [2] 19, 312). In the higher boiling phenolic portion of coal-tar oils (Schulze, B. 18, 615).

Formstion.-1. By exidation of benzoic aldehyde, benzyl alcohol, teluene, ciunamic acid. &c.-2. In small quantity, by passing a current of dry CO, through a nearly boiling inixtura of aluminium oblorido and benzene (Friedel a. Crafts, C. R. 86, 1368) .- 8. In small quantity, by the action of H₂SO₄ and MnO₂ on benzone, especially when formic acid is added (Carius, A. 148, 51, 59). - 4. By distilling calcic phthalate with lime (Depouilly, Bl. [2] 3, 163, 469). -5. By the action of H_2SO_4 and MnO_3 on casein or gelatin (Gnckelberger, A. 61, 80).-6. By fusing potassium benzene autphonato with sodium formate (V. Meyer, B. 3, 112) .-- 7. From benzonitrilo by eaponification. - 8. By passing CO, into sodium in bromo-benzeno (Kckulé, A. 137, 129).

Preparation .- 1. From gum henzoin by sublimation or by extracting with line-water or acctic acid (Mohr, A. 29, 178; Schoole, Opuso, 2, 23; Wöhler, A. 49, 245; Loow, J. pr. 108, 257; Guichard, Bl. [2] 19, 357). Some varieties of gum benzoin contain cinnamic acid, but this acid is absent from the benzoin of Siam or the l'alembang benzoin from Sumatra; the latter yields 10 p.o. benzoic acid (Saalfeld, Ar. Ph. [3] 16, 280). Benzoic said that has been sublimed from gum benzoin leaves a small quantity of oily reciduo when treated with aqueous Na, CO,; this oil consiets of gualacol,

DESTRUCTION AND

oxen is left for some days to patrely, when the hippuric acid is split up into glycocoll and benzile acid; milk of lime is added and the liquid concentrated; excess of lime is ppd. by CO₂, and the filtrate ppd. by Fe.Cl.; the ferric benzoate is decomposed by ICl. Benzoic acid prepared in this way crystallises in platos and smells of urine, but by sublimation it may be freed from the smell and then crystallises in needles (Dymond, Ph. [8] 14, 463).—3. From benzotrichloride by decomposing it with water under pressure, with lime or baryta-water, or with McCl. and glacial HOAc (2 mols.) at 100° (Jacobsen, B. 13, 2013).—4. From benzyl chloride by boiling with dilute 11NO, (Lungs a. Petri, B. 10, 1275; cf. v. Rad, D. P. J. 231, 538).

533). Properties.—Needles or pearly plates. When pure it does not melt under water, but slight impurities greatly affect its physical properties; the so-called salylic neid was impure benzoio acid (Kolbe a. Lantemann, A. 145, 187; Kekule, A. 117, 159; Griess, A. 117, 31; Reichenbach a. Beilstein, A. 132, 309; Kolbe, J. pr. [2] 12, 151). Volatile with steam (1g. passing over with about 2,000 c.c. water). It dissolves in conc. 11,80, and is reppd. by water. It is not attacked by boiling dilute 14XO, or CrO, (which convert cinnamic neid into benzoic aldebyde); its neutral salts give a buff-coloured pp. with Fe,Cl.

Reactions .- 1. Passage of the vapour through s red-hot tube gives CO, and benzene .- 2. Distillation with lime produces benzene. -3. Fusion with NaOII produces benzene (75 p.c. of the theoretical amount) and a little diphenyl (Barth a. Scuhofer, B. 12, 1256). - 4. Fusion with KOII produces chiefly p-oxy-henzoic acid, but also oand m- oxy-benzoio acids, oxy-iso-phthalic acid, diphenyl o-, m-, and p- carboxylic ccias, and a brown amorphous substance (Barth a. Schreder, M. 3, 799). -5. MnO, and H.SO, form CO., furmic acid, and small quantities of phthalic and tereputhalic acids (Carius, A. 118, 50; Oudemans, Z. [2] 5, 84).—6. Hydrogen peroxide and H.SO, produce salicylic acid (Hanriot, C. R. 102, 1250). - 7. Vapours of benzoio acid passed over heated zinc-dust form benzoic aideliyde (Baeyor, A. 110, 295) .-- 8. Sodium amalgam reduces it to benzyl alcohol, and henzoleic acid C, H₁₀O₂, and an oil C₁, H₁₄O₂ (Kelbe, A. 118, 122; Hermann, A. 132, 75).—9. PCl, forms benzoyl chloride. - 10. Distillation with KSCN or Pb(SCN), gives benzonitrile. -11. Benzene and P.O. at 190° give I-mzophenono (Kollarits a. Merz, B. 5, 417).—12. Dimethylaniline and P.O. givo C. 11, CO. C. 11, NMe. (O. rischer, B. 10, 958).

13. Chlorine produces chloro-benzoio neids.—

14. Bromine forms bromo-benzoio noids.—15. Iodine in presence of 1110, forms iodobenzoic acid. -16. Cone. 11NO, forms m-nitro-benzoic acid .- 17. Fuming 11.SO, forms sulphobonzoic acid.—18. In the animal organism it is converted into hippurie acid and excreted in the urine (Wöhler). -19. Cr.F. forms di-fuoro-benzoic acid (Jackson a. 1lartshorn, B. 18, 1993).

Salts. — Benzeic acid decomposes carbonates, but an alcoholio solution of potassium benzente is decomposed by CO. Calcium benmate gives on distillation benzophenone, and smaller quantities of benzene, anthraquinone,

and letra-phanyl-methane (Rekull as Franchi-mont, B. 5, 909). Calcium benzonte distilled with calcium formate gives benzole aldehyde, Potassium benzonte distilled alone or with sodium formate gives terephthalic and isophthalic acids (Richter, B. 6, 876; Conrad, B. 6, 1395). Cuprio benzonte gives on distillation benzenc, benzole PhOBz, and phenol (List a. Limpricht, A. 90, 190). Cyanogen bromide acts upon potassium benzonte thus; Ph.CO,K+CNBr=Ph.CN+CO,K+KBr (Calcours, A. Ch. [3] 52, 201). Potassium benzonte when electrolysed gives K and benzole anhydride; in presence of excess of KOH acetylene is also formed (Bourgoin, Z. [2] 4, 566).

formed (Bourgoin, Z. [2] 4, 506).

Al₂A'₁(OII), aq: crystals (Sestini, Gioognanl, a. Zavatti, Bl. [2] 13, 488).—NII, A': deliquescent; on distillation it givos benzonitrile.—NH, HA'_x—BaA'_x2aq.—CdA'_x2aq.—CaA'_x3aq: S. 3·5.—CoA'_x3aq.—CoA'_x2aq.—CuA'_x2aq: needles.—CrA'_xxaq.—CrA'_xxaq.—CrA'_x(OII), 2aq (Schiff, A. 12, 169).—FoA'_x(OII), 5aq: bull-coloured pp.—IaA'_x3aq.—PbA'_xaq: plates.—PbA'_x2PbO.—MgA'_x3aq: S. 4·5 at 25°.—MnA'_x4aq: lurge flat prisms, S. 6·55 at 15° (Senbert, B. 20, 791).—HgA'_xaq.—HgA'_x.—NiA'_x3aq.—KA'3aq.—XaA': S. (alcoliol) 5 at 20°.—NaA'aq.—SnA'_xaq.—ZnA'_x—Msthyl ether C₆H₃CO_xMa. Mol. w. 136.

Methyl ether C₈H,CO,Me. Mol. w. 186, (199°) (Kopp); (195°5) at 768 min. (Stohmann, J. pr. [2] 36, 4). S.G. ¹²I·10 (Kopp); ²g·19862 (Brühl), S.V. 149·8 (Ransay), S.H. 363 + 90075L, H.F. 84,024 (S.). µs 1·5289. R. © 61·30 (S. Porned by distilling wood spirit (1 pt.), benzoic acid (2 pts.), and H.SO, (2 pts.) (Dumas a. I'eligot, A. Ch. [2] 58,50; Malaguti, A. Ch. [2] 70, 387; Carius, A. 110, 210).

[2] 70, 387; Carius, A. 110, 210). Ethyl ether C. 11, CO_Et. Mol. w. 150. V.D. 5:53 (enlc. 5:2) (Troost, C. R. 89, 851). (211·2° cor.) (Linuemann, A. 160, 208); (211·4°). (Stohmann, J. pr. [2] 36, 4). S.G. $\frac{3}{4}$ 10473 (Brühl); $\frac{12}{2}$ 10:50 (L.). S.H. 374 + 00075t (R. Schiff, A. 234, 300). H.F. 91,693 (St.). μ_{θ} 1:517 (B.). It $_{\infty}$ 68·82 (B.). Formed by saturating a solution of benzoio acid (3 pts.) in alcohol (2 pts.) with HCl and distilling the liquid. Converted by Br at 270° into benzoio acid and citylone brounde (Naumann, A. 133, 199). Ferms crystalline compounds with titanio chloride: BzOEtTCl_{θ} – BzOEt2TCl_{θ}, and with aluminium chloride: BzOEtAlCl_{θ} (Gustavson, B. 13, 157; Scheele, Onuscula, 2, 141; Dunnas a. Boullay, A. Ch. [2] 87, 20; Wöhler a. Lielig, A. 3, 274; Peville, A. Ch. [3] 3, 188).

Propyl ether Call CO.Pr. (230°). 8.6. 14 1-032 (L.); 1: 1-025 (S.). H.F. 98,990 (Stohmanp, J. pr. (2, 36, 4). S.H. 383 + 00075t (Schiff, A. 234, 300).

Isopropyl ether C.11, CO.Pr. (218°).
S.G. 9 1-023 (Silva. Bl. 12, 225). According to
Linnemann (A. 161, 51) the ether splits up on
distillation into propylene and benzoic acid.
n. Butylether C.H., CO.C.H., (247.3 cor.).
S.G. 22 1-00.

S.G. ⁹² 1·00. Isobutyl ether. (234°) at 755 mm. S.G. ¹² 1·002. H.F. 105,628 (St.).

Isoamyl ether C.11., CO,C.H.,. Mcl. w. 192. (261°) (Kopp, A. 94, 311); (253°) (Stohmann, J. pr. [2] 36, 4). V.D. 6-71 (calc. 6-65,

Troogl. C. E. 39, 851). B.G. #1-004 (K.); 12 998

R.). H.C. 1.570,048 (St.).
Formed by heating ethyl benzoate with isoamyl alcohol at 280° for 60 hours (Friedel a. Crafts, Bl. [2] 2, 100).

Octyl ether C.H. CO.C.H ... (306°) (Zineke, A. 152, 7).

Decyl ether C.H. CO.C., 11, (over 280°) (Borodin, J. 1864, 338).

Cetylether Call CO Call .. [30°] (Becker, A. 102, 221).

Allyl ether C.H. CO C.H. (242°) (Zinin. 4. 96, 362); (230) (Berthelot a. de Luca, A. 100, 860); (280') (Cahoura a. Hofmann, A. 102, 2971

Ethylene ether (C.H.CO.) C.H. (360°).

Propylene other (C.11, CO.), C.II. [72].

Isoamylene other (C,11,.CO,),C,11,.[133]

Mayer, Bl. (2, 2, 451).

Other ethers of benzoic acid are described as benzoyl derivative, of the hydroxylic compounds from which they may be derived.

References. - V. also Aldenyno., Amino., Bio. мо-, Вкомо-амідо-, Вкомо-кітко-, Вкомо-кітко-OXY-, BROMO-OXY-, BUTYL-, CHLORG-, CHLORG-10DO-, Chloro-oxy-, Cyano-, Fluoro-, Iodo-, Iodo-oxy-, NITRO-, NITRO-BUTYL-, NITRO-OXY-, NITRO-PROPYL-, OXY-, SULPHO-, DENZOIC ACID.

Orthobenzeie acid C.11,.C(OH),.

alcohol.

ý

Ethyl ether C.H.C(OEO, (220 1 225). From benzotrichloride and NuOEt at 100 (Limpricht, A. 135, 87).

Tri-acetyl derivative C.11.C(OAc). From C.II. CCl. and AgOAc. Readily splits up into Ac.O and C.H.CO.O.Ac.

Salphinide of Lenzeic acid v. lands or sul-

PROBENZC C ACID.

BENZOIC ALDEHYDE C.II,O i.e. C.II, CO.II. | Deneallehyde. Oil of bitter almonds. Mol. w. 106. (179-). S.G. 3° 1-0455 (Brühl); ½1-0504 (Mendelčeff, J. 1860, 7). S. 33 (Flückiger, J. 1875, 482). μ_B 1-5624. R_A 51-65 (B.). H.F. 23,254 (Stohmson, J. pr. [2] 36, 3).

Formation. -1. From almonds (q. v.). -2. By oxidation of benzyl alcohol (Cannizzaro, A. 88, 130), einnamic acid (Damas a. Peligot, A. 14, 50), and proteids (Guckelberger, A. 61, 60, 72, 86).—2. By boiling benzyl ch.oride with water and nitrate of lead (Lauth a. Grimaux, A. 143, 80), nitrate of copper, or sodium nitrate. ... 8. By heating benzylidene chloride with water or alkalis (Cahours, C. R. 56, 222) .- 4. By mixing benzylidenc chloride with conc. 11,80,. diluting, and distilling (Oppenheim, Z. [2] 5, 411). 5. By passing vapour of benzoic or plathalic acid over heated zino-dust (Bagyer, A. 110, 295).

6. By reducing benzoic acid with SoCl. (Obsert, J. 1862, 263), or sodium amalgam in slightly acid solution (Kolbe, A. 118, 122) .- 7. By distilling calcium benzoate with calcium formate (Piria, A. 100, 104) .- 8. From benzylidene ehloride and silver oxalate (Golowkinsky, A. 111, 252) or potassium carbonate (Meunier, 3s. 12) 2, 38, 159).

9. From tolucne by successive treatment with Cro.Cl., and water (Etard, C. R. 90, 534).—

10. From benzylidene chloride, acetic acid, and Ph.CHCl. + CH.CO.H = Ph.CHO + CH.COCl + HCL ZnCl,:

Preparation.-I. Bennyl ablaride (1 pt.) Experience.—I. Senny: aniorany to pro-is cohobated at 100° with water (19 pts.) and lead nirate (12 pts.), a corrent of OO, being passed through the apparatus. The product is distilled and the light oil fractionated. It is shaken with a saturated solution of NaHSO, and thoresultingorystalline compound is washed with alcohol, erystallised from water, and then decomposed by boiling aqueons Na OO, (Lauth a. Grimaux, A. 143, 80 ; Bertagnini, A. 85, 183). --2. Crude benzylidene chloride is heated at 110°-130° with an equivalent quantity of dry ozalic acid, the product is distilled in vacuo: 1'hCHCl_c+H₁C₁O₁ ~ 1'h.CHO+2HCl+CO₂+CO (Anschütz, A. 226, 18). 3. A mixture of benzyl chloride (2 mols.) with benzylidene chloride (1 mol.) obtained by chlorinating toluene till the S.G. is 1.175 is boiled with water and MnO. (2 mols.) (Schmidt). -4. By heating benzylideno chloride with aquenus KOH under pressure, or by boiling it with milk of lime. 5. Bitter almonds are freed from almond oil by pressure. The press-cake (12 pts.) is made into a paste with boiling water (110 pts.); after 15 minutes the paste is allowed to cool. The smulsin is destroyed by boiling, and therefore a second quantity of the press cake (1 pt.) is mixed with cold water (6 pts.) and added to the first. After 12 hours' maceration the whole is distilled with steam. The yield is 2 p.c. of the press cake (Pettenkofer, A. 122, 77; cf. Liebig a. Wöhler, A. 22, 1). In this operation amypdalin is split up by the unorganised ferment emulsin, the products being benzoic aldehyde, prossic seid, and placose

C, H2, NO11 + 211 O ~ C, H2O + CNH + 2O, H4, O4 aldeliyde so prepared contains Benzoio prassic acid, which appears to be combined in the form of the cynnhydrin Call, CH(CH).CN; for a nexture of benzoic aldehyde and prussio acid yields methylamine on reduction, while crude oil of bitter almonds yields amide phenylethane C.H. CH, CH, NH, again, a mixtura of benzoic aldebyde and prussic neid, on treatment with chlorine, yields Call, Cl.CO.Cl, while oil of bitter almends yields, by similar treatment, Call, Cl.(Oil). CO.N. Cl.Call, (Fileti, O. 9, 446). Prussic acid may be removed by shaking with FeSO, and lime or potash, or by digesting with HgO and water. The uldehyde is then purified by means of NaIISO, as described under 1.

Properties .- Colourless oil. It is not polschools. It exidises rapidly in the air, but the addition of a little prissic acid hinders the oxlds. tion (Dusart, Bl. 8, 459). It does not reduce Felding's solution.

Reactions .- 1. Oxidised to benzoic acid by air or other oxidising agents. Conc. HNO however, forms v.. (and a little o) nitro-benzal-dehyde. - 2. Aqueous or alcoholic potash gives benzyl alcohol and potassium benzoate. -8. Led over red-hot pumice it is split up into CO and benzene (Barreswil a. Boudault, A. 52, 860) .-4. PCl, forms benzylidene chloride (Cahours, A 70, 39). COCl, acts similarly (Kempf, J. pr. [#] 1, 412). - 5. Chlorine forms benzoyl ohlorid and a compound of that body with benzoic aldahyde, C. II, CliCl(OBz) (Laurent a. Gerhardt, J. 1850, 489). Bromine acts similarly, forming C.H. CiBr(OBz) [70°] (Liebig a. Wöhler, A. S. 266; Claisen, B. 14, 2475) .- 6. Succingl chie

ride produces succinic acid and benzylidene chloride (Rembold, A. 138, 189) .- 7. Sodium amalgam reduces it, in presence of water, to benzyl alcohol, hydrobenzoin, and isohydrobensoln .- 8. Polassium cyanide produces benzojn. 9. Aqueous HI (S.G. 2.0) at 280° reduces it to teluene (Berthelet, J. 1867, 316).—10. H.S forms thiobenzaldehyde.—11. Aqueous NU, torms hydrobenzannide (C.H.CH)₃N₂.—12. Ammonium sulphide forms thiobenzaldine C21 II. NS. -13. NII. and sulphide of carbon form NH. CSS. N(CH.C. H.) .. -14. With acetic anhydride and sodium acetate, on heating, it forms sedium cinnamate (v. Penkin's Synthesis, p. 108). The reaction prebably takes place in two stagee: Call,CHO + CH, CO, Na =

C.II..CH(OH).CH, CO, Na =

H2O+O4H3.CH;CH.CO2Na.-15. With Ac2O and sodium succinate it gives the lactone of CO₂H.CH₂.CH(CO₂H).CH(OH).C₂H₃.—16. With sodium isobutyrate and isobutyric anhydride it torms C.H. CH(OH).CMe, CO, II, v. Oxy-PHENVL-VALERIO ACID (Fittig, A. 216, 119) .- 17. With Ac₂O and sodic butyrate at 100° it gives only phenyl-angelic acid, whereas at 180° the chief product is cinnantic acid (Slownm, A. 227, 53). — 18. The reaction PhCHO+Cll XY= HO+Ph.CH:CXY takes place under influence of dry IICl or aqueous er ulcoholic KOII en condition that X or Y is of the form CO.Z, e.g. benzoie aldehyde acting on acctone, mesityl oxide, acetophenone, pyruvic acid, malonio ether, and acete-acetic ether. Perkin's reaction is of a cimilar nature. Occasionally intermediate compounds of the ferm Ph.Cll(OH).CllXY are tormed (Claisen, A. 218, 121). - 19. Sodium malonate and Ac,O react in the cold, giving off CO, and forming cinnamic acid, as follows: Ph.CHO + CH₂(CO₂H)₂ = l'hCH:C(CO₂H)₂ + H₂O = PhCH:CH₂(CO₂H + CO₂ + H₂O (Stuart • J, J, 43, 404) .- 20. Sodium isosuccinate and Ac O act similarly, forming phenyl-iso-crotonic acid:
Ph.CHO+CllMe(CO₂H)₄ =

Ph.CH:CMe.CO₂H + CO₂ + H.O.

91. Acetyl chloride and sinc-dust form diacetylhydrobenzein; while benzoyl chleride and zinc dast form di-benzoyl-hydrobenzoïn (Paal, B. 15, 1818; 16, 636; 17, 909).—22. Reacts with nitro-paraffins thus: Ph.CHO + H.C(NO.).CH, = H₂O + PhCH:C(NO₂)CH₂ (Priebs, A. 225, 319). -23. A solution of aniline in cone. HCl pps. yellow crystals of a molecular compound. They are only stable in presence of conc. HUl (Elbers, A. 227, 357). If SnCl, be also present a compound (NPhH,IICl), (C,H,O), SnCl, is formed (E.).—24. Anilino fofus benzylidene aniline, O.H. CII: N.C.II.; e-toluiding ucts similarly. When heated in presence of HCl or ZuCl, aniline forms di-umido tri phenyl-methane. Dimethylanthins in presence of ZuCl, gives C.H.OH(C.H.NMe.). (Fischer, B. 10, 1623); dimethyl m. (but not o- or p.) tolnidine behaves similarly (Fischer, B. 13, 807).—25. Ethylene-diamine forms (C.H., CH.N.C.H., [54°] (Masen, B. 20, 267) .- 26. (8) Naphthylamine forms benmylidene-(B) naphthylmine and then phenylnaphthacridine dihydride PhCH<C...II (Claisen, A. 237, 261).-27. Resorcin in pre-

cis, and orcis act similarly (Michael a. Ryder, B. 19, 1388; Am. 9, 130).—29. (β)-Naphthol left to stand for several days in the cold with an acctio acid colution of benzaldehyde treated with a few drops of HCl forms di (8) naphthyl henzaldeliydate (di-naphthyl-ortho-benzoic aldehyde) C.H.CH(OC10H.). [205°]. It is a crystalline pp., sl. sol. all ordinary selvents; insol. aqueous alkalis. By warming with acetic acid and a few drops of HCl it is converted into the isomeric di-oxy-di-naphthylphenyl methane Call CII(CoH. OII), which at the same time loses II_O, giving the compound C_bH_b .CH $< C_{10}H_b > O$ (Claisen, B. 19, 3317).— 29. Accione in presence of aqueeus NaOH produces C_aH CH:CH:CO.CH₂ and the compound C_aH₂.CH:CH.CO.CH:CH.C_aH_a. In general, compounds containing the group CH. CO react with benzoic aldehyde, exchanging the H. fer CHPh (Claisen, B. 14, 349, 2468; v. Benzyldens-acktone).—30. Prussic acid ferms mandelonitrile or the cyanhydrin of benzeic aldehyde, C.H. CH(OH) CN (v. MANDELLE ACID). This is converted by alcohol and HCl into mandelic imido-ether, C. II. CH(OH). C(NH). OEt (Völckel, A. 52, 361; Tiemann, B. 14, 1967). Benzoic aldehyde (4 vols.) mixed with nearly anhydrous prussic acid (1 vol.) and shaken with alcoholic KOH forms benzimide C, H10N,O,, [167], a flocculent substance, insol. water, alkalis, and acids (Laurent, A. Ch. [2] 59, 397; 66, 193; Zinin, A. 34, 188; B. 2, 552; Gregery, A. 54, 372). 31. Hydrogen iodide forms a pungent compound CaH at O [28°] insol. water (Genther a. Cartmell, A. 112, 20).-32. SO, forms a disulphonic acid C. II, (SO.II), CHO (Engelhardt, J. 1864, 350).-33. By treatment with a methyl-alcoholic solution of sodium methylate a white solid compound CaHaC(OMa)(OC,H)(ONa) formed. The same body is formed by the action of sodium methylate on benzyl benzonte or of sodium benzylate on methyl-benzeate. By treatment with acctic acid it is split up inte a mixture of benzyl benzoate, methyl benzoate, benzyl alcohol, and methyl alcohol.-34. If benzaldchyde is heated with a small quantity of sodium beneylate for several days at 100°, it is slowly polymerised to benzyl benzoate. Probably the compound C.H. C(OC, H.) ONa is first formed, and then decomposes into benzyl benzoate and sodium benzylnte, which latter again reacts upon a further quantity of benzaldeliyile, preducing more of the intermediate compound, and so on (Claisen, B. 20, 646) .-35. By boiling with ammonium formate it yields tri-, di-, and mone- benzyl-amine and their formyl derivatives, together with other products (Leuchart, B. 19, 2128).—36. Ammonium sulphocyanide at 110° forms benzylidene-thiobiuret C.H.CH NH.CS NH [237°] (Brodsky, M. 8, 27) .- 37. Benzene azo benzene and ZnCl, forms 'benzylidene benzidine' C. II. N2O2(?) [239] (Barzilowsky, J. R. 1885, 366).—38. Acts upon an alcoholic selution of sodium acetoapon an arco.tone solution of solution decision decision decision of the solution of the solution alkalis (Michael, J. pr. [2] 35, 450),—39. Hydrazine benzoic acid, NH₂NH₂Cl₂CO₂H₁forms sence of Hel forms a resin C₂H₂O₄ (Baeyer, benzyli lene-hydrazine-benzoic acid O₁₄H₁₂N₄O₄
B. 5, 25). Phenol, pyrocatechin, Phloroglu[1725 uncor.] (Roder, A. 236, 171).

C.H.CH(OH).SO, Na jaq. Small crystals, v. e. sol. water, insol. cold alcohol. Decomposed by boiling water, boiling dilute acids, or cold alkalis or alkaline carbonates (Bertagnini, A. 85, 188). — C.H.CH(OH).SO,K: laming. — C.H.CH(OH).SO,NH, aq: formed by action of 80, on an alcoholic solution of hydrobenzamide from the Na salt and BaCl. -2. With SO₂ and aniline: (C₄II₂CII)(C₄II₃N₃SO₂ (Schiff, A. 140, 130). - p. Toluidine forms, similarly, (C,H,O), (C,II,N). SO, ... Amido acids shaken with aqueous solutions of SO, and henzoic aldchyde form crystalline compounds, e.q. (from glycocoll), C.H. CH(OH).SO, NH, CH, CO.H, and C.H. CH(OH).SO, NH, C.H. CO.H. (from amido benzoic acid) (Schiff, A. 210, 123).—3. With inorganic salts. C.H. 013CaCl. (*) (Ekmanu, A. 112, 175). - C. H. OBF. (Landolph, J. 1878, 621).

Oxim v. BENZALDOXIM.

Phenyl-hydrazide C.H., CH, N.HC, H., [153°]. Formed by adding a solution of phenylhydrazine hydrochloride and sodium acetate to an aqueous solution of benzaldehyde; the white pp. is distinctly visible with a solution of 1 pt. of benzaldehyde in 50,000 pts. of water (Fischer, B. 17, 571). Can be crystallised from alcohol. Insol. water. Acetyl derivative Collin N. Ac [120], long needles (Schroeder, B. 17, 2096). Benzoyl derivative C.H.,CH.N.NBzPh : [122]. Formed from benzaldehyde and u benzyl-Idenyl-hydrazine. Very thin silky needles. V. sol. alcohol (Michaelis a. Schmidt, D. 20, 1717).

Methyl-phenyl hydrazide PhMeN.N.CH.Ph. [102°-1015°].-1. Formed in small quantity from the methyl phenyl-hydrarido of phenyl-glyoxylio acid (q. v.) at 120 : Benzoic ablehyde and methyl miline are also formed .- 2. From benzoic aldehyde and methylphenyl-hydrazine in alcoholic solution. White needles (Wallach, A. 227, 352).

Derivatives of Benzoic ortho-aldehyde are described as BENZYLDENE derivatives

BENZOIC ALDEHYDE, AMMONIA-DERI-VATIVES OF. The most important of these are hydrobenzamide, amarine, and lophine.

1. HYDROBENZAMIDE C21H (aN. i.e. (Call, CII) AN.

Tribenzylidene diamine. [1102].
Formation.—By the action of ammonia upon benzoic aldehyde (Laurent, A. Ch. [2] 23; 66, 18), upon benzylidene acetate C.H. CH(O.C.11,0)2 (Wicke, A. 102, 368), or upon benzylidene-dichloride (Engelhardt, A. 110, 78).

Preparation .- Benzoic aldehyde, which must be free from hydrocyanic acid, is left for some days in contact with strong aqueous ammonia. The crystalline mass which separates is washed, first with water and then with other, and finally recrystallised from alcohol. Heat accelerates the action, but diminishes the yield. Equation: $BC_aH_{a}CHO + 2NH_{a} = (C_aH_{a}CH)_aN_a + 3H_aO_a$

Properlies. - Crystallises from hot alcohol in colourless rhombic octahedra, which are generally wedge-shaped. Insol. water: sol. alcohol Has a sweetish taste and is not and other. poisonous.

Reactions .- 1. When heated for several hours to 120°-130' it is converted into the iso-

Combinations. — 1. With bisulphites. | merio amarine (Bertagnini, A. 88, 187). -By H.CH(OH).SO.Na jaq. Small crystals, v. c. | destructive distillation it yields lophine O₂₁H₁₁N₂ (Laurent).—2. Boiling with aqueous potash copverts it into amarine (Fownes, T. 1845, 263), whilst alcoholic potash breaks it up into benzoie aldehyde aud ammonia. This last decomposition is also effected by prolonged boiling with alcohol alone. Pasion with potash produces very complex decomposition, yielding among other products lophine. - 3. Dilute acids hydrolyse it readily on boiling, slowly in the cold, into benzoic aldehyde and ammonia. The case with which this hydrolysis occurs is best accounted for on the assumption that hydrobenzamide is tribenzylidene diamine -- 4. Dry hydrobenzamide absorbs guscous hydrochloric acid. During the process a non-nitrogenous substance volatilises, and the residue, when treated with water, yields benzoic aldehyde and ammanium chloride. If instead of treating the residue with water it is heated, benzonitrile and benzyl chloride distil over, whilst a complex mixture of basic substances remains (Ekmann, A. 112, 151; Kuhn, A. 122, 308). -5. It unites with 2 mols.of hydrocyanic acid to form hydrobenzamidedihydrocyanide, a yellow crystallino mass melting at 55°, which, when treated with hydrochloric acid, is decomposed into henzoic aldehydo and phenylamidoacetonitrile: C₂H₁₈N₂2HCN + H₂O - C₂H₆O + 2C₆H₅CH(NH₂).CN, the latter compound being subsequently hydrolysed to the corresponding acid (Plochl, B. 13, 2119). Under other conditions the benzoio ablehydo and phenylamidoacetonitrile thus formed may unite with elimination of water to form beneagl-arotide, $C_1H_12N_2$, thus: $C_2H_1O+C_2H_2CH(NH_1)$, $C_1H_{12}N_2+H_2O$ (Plach), B. 14, 1142). When an other all solution of hydrohenzamide is in xed with 1 mal, of hydrocyanic acid, and gaseous hydrochlorio acid is passed into the liquid, a hydrochloride of the monahydrocyanide of hydrolenzumide, C21H18N2,HCN,HCl, separates. On bailing this precipitate with concentrated hydrochloric acid, it is decomposed into benzoic allehyde, ammonia, and the hydrochloride of an anhydride of the formula C, 11, N,O: c, 14, N,10:N+2H,O=C, 16, N,O+C,H,C, 10:O+N, The free anhydride mells at 161° and sublines without decomposition; the neid C₁₈H₁₈N₂O₂ melts at 120° (Plöchl, B. 14, 1139).—6. Dissolved in absolute alcoholand treated in the cold with 3 p.c. sodian analyam it yields benzylidene-dibeneylimide Call, CH(NH.CH, C.H.)2. Astronger ionalgam, aided by heat, converts it into henzylamine and toluene (O Mischer, B. 19, 748). Hydroberczamide is also acted upon by chlorine, sulphurous anhydride, sulphuretted hydrogen, and ethyl i lide, but the reactions

> 2. AMARINE: Callinna Isomerio with hydrobenzamide. Probable constitution: C,H,CNH

are not of importance.

CH.C. II. (E. Fischer, A. 211, 217; C.H.C.NH Japp a. Robinson, C. J. 41, 323); or Call.CH.NH C.C.Il, (Claus, B. 15, 2338). CH_CH_N [100].

Formation .- 1. By the action of ammonla upon an alcoholic solution of benzoic aldehyde

(Laurent, O. R. 19, 858) .- 2. By holling hydrobenzamide with aqueous potash (Fownes, T. 1845, 268).—8. By heating hydrobenzamide to 120°-130° (Bertagnini, A. 88, 127).-4. Together with lopline by distilling the double compound of benzoic aldehydo and aminonium hydrogen sulphite with excess of slaked lime (Gössmann, A. 93, 329).—5. In small quantity when ammonia acts upon a mixture of benzoin and

benzoio aldehyde:—C₁,H₁₂O₂ + C,H₄O + 2NII₃ = C₃,H₄N₄ + 3II₄O (Radziszewski, B. 15, 1495),

Freparation.—Hydrobonzamido is heated from 2 to 4 hours at 120°-130°. The vitrous mass thus obtained is dissolved in boiling alcohol, and an excess of hydrochloric acid is added. The amarino hydrochloride, which separates in white crystals, is purified by recrystallising Iron boiling alcohol and is then decomposed with

Properties. - Deposited from alcohol in lustrous prisms. Melts at 100° (Fownes), but by boiling for some time with water, is converted into a granular substance melting as high as 126°. This modification is reconverted into prismatic amarine, melting at 100°, by recrystallisation from alcohol (Claus, B. 18, 1678). Insol. water, v. sol. alcohol and other; the alcoholic solution having an alkalino reaction. Has a bitter taste, which is noticeable only after a time. Very poisonous. Combines with acids to form sparingly soluble salts which have an intensely bitter taste. — C₂₁H₁₈N₂,HCl. an intensity officer (aste. — $C_{21}H_{11}N_{21}HCI)$. Needles, sparingly soluble in boiling water. — $(C_{21}H_{11}N_{21}HCI)_{21}PtCI_{12}$. Yellow crystalline saft (Gössmann). — $C_{21}H_{11}N_{21}HN$ (Borodine, 4. 110, 79). — $C_{21}H_{11}N_{21}HNO_{4}$ (Fownes). $(C_{21}H_{11}N_{12})_{11}HNO_{4}$ (Fownes). $(C_{21}H_{11}N_{12})_{11}H_{2}CI_{2}O_{2}$. Yellow precipitate, almost insol, water.

Reactions. 1. By destructive distillation it yields ammonia, toluene, and lophine, C, H16 E, (Fownes; v. also Itadziszewski, B. 10, 70). - 2. By limited oxidation, as when the chromate is boiled with glacial noetic acid, it is converted into lophino: $C_{21}11_{18}N_{2} + O = C_{21}11_{18}N_{2} + 11_{*}O$ (Fischer a. Troschke, B. 13, 707). Excess of chromic neid oxidises it to benzoic acid. - 3. When a hot alcoholio solution of amurine is acidified with acetio acid and a hot concentrated solution of an alkaline nitrite is added, nitroso-amarine, Cally (NO) No separates, and, by recrystallishtion from alcohol, may be obtained in rhombie tables, which, when heated to 150°, decompose with formation of lophine (Horodine, B. 8, 934) .- 4. With mitric acid in the cold, mononitro amarine, CnH1. (NO.) No. is formed; hot nitric acid converts it into dinitro amarine, C2, H10 (NO.) N2. The latter compound yields, with reducing agents, diamido amarine, C21H16(NH2), N2. Salts ol these nitro- and amido- bases have been prepared (Claus a. Witt, B. 18, 1670).-5. With acetyl chloride an ethercal solution of amarine yields a product which, when trented with alcohol, is separated into amarine hydrochloride and diacetyl amarine, C₂₁II₁₆(C.II₃O)₂N₂ [268°] (Bahrmann, J. pr. [2] 27, 295).—6. Amarine reacts with the halogen compounds of the alkyls to lorm substitution and addition compounds. Thus when amarino, methyl iodido, and ether are allowed to stand together in the cold, methytamarine hydriodide, Call, McN, III, is formed.

Ammonia is without action upon this sait, but alcoholic potash liberates methylamarine [184°]. When amarine is heated with methyl icdide, a dimelhylamarine hydricdide, C₂₁H₁₆Me₂N₂,HI, is formed. Like the monomethyl compound it is not decomposed by ammonia; by treatment with alcoholio potash, dimethylamarine [146°] is obtained, and this base unites with hydriodic acid to lorm a hydriodide isomerio with the loregoing and differing from it in being decomposed by ammonia even in the cold (Claus a. Elbs, B. 13, 1418). Dimethylamarine and the other dialkyl-amarinos do not combine with the

lulogen compounds of the alkyls.

Further derivatives of amarine. — Ethylamarine, C., H., Eth. [163°] (Claus a. Scherbel, B. 18, 3080); dielhylamarine, C., H., Et., N. [110°-115°] (Borodine, A. 110, 82).—Benzyl-amarine, C₂₁H₁₇(C,H₂)N₂ [123°-124°]; dibenzyl-amarine, C₂₁H₁₆(C,H₂)₂N₂ [139°-140°] (Claus s. Eibs, B. 13, 1418; Claus, B. 15, 2330; Claus a. Kohlstock, B. 18, 1819. - Ethylbenzylamarine, Kohlstock, B. 18, 1819).—Ethylbenzylamarine, C₁₁II_{1n}Et(C,II₁)N₂ [135⁻] (C. a. K.).—Hydromethylbenzylamarine, C₂₁II_{1n}Me(C,II₂)N₂O[208^o]; hydrotrimethylamarine, C₂₁II_{1n}Me₂N₂O [158^o] (Claus, B. 15, 2326).—Other derivatives: C₂₁II_{1n}N₂(C,I,I), (C,I,I), (C,I,I) a. Elbs, B. 16, 1272); diamarine silver nitrate, (C₂,H₁₈N₂)₂,AgNO₃ aq [218°] (C. a. K.). When amarine silver is mixed with one mo-

lecular proportion of an alkyl-halogen compound (the latter diluted with benzenc) and allowed to stand in the cold, a double compound of amarinesilver with the alkyl-halogen is obtained. At the samo time a small quantity of monalkyl-amarine is formed from the double compound by climina. tion of silver iodido. In this way the following compounds have been prepared (Claus a. Schercompounds have been prepared (claus a. Scher-bel, B. 18, 3077); amarine-silver methylo-iodido, C_nH₁,AgN_mCl₄1 [173°]; amarine-silver ethylo-iodide, C_nH₁,AgN_mC.11, [115°]; amarine-silver isoprepylo-bromide, C_nH₁,AgN_mC₂HBr, [140°]; and finally amarine-silver benzylo-chloride, C_nH₁,AgN_mC,H,Cl, [250°]. When amarine-silver is treated with benzyl ethloride it yields benzoyl-amarine, C,H,,(C,H,O)N,, [180], which forms salts with neits and also unites with benzoyl chloride to form benzout-amarine benzoul chloride, C., H., (C,H,O)N., C,H,OCl, [312°], and with benzyl chloride to form benzoul-amarine benzylo-chloride, C., H., (C. H., ()) N., C., II, Cl [351°], which latter is isomerie with benzyl-amarins benzoyl chtoride, C₂₁H₁₂(C II,)N₂₀C,H₂OCl, [340°_350°], obtained by the action of benzoyl chloride upon benzyl nuarine. The bases corresponding with these two chlorides are also isomeric. Benzoyl-amarine also unites with methyl iodide and with ethyl iodide to form additive compounds melting at 318° and 354° respectively (C. a. S.).

C.H. C.NH 3. LOPHINE C₁₁H₁₄N₂ = C.C₄H₄, C.C₅H₄, C.H₄, C. N. Triphenylolyoxaline (Japp a. Robinson, C. J. 41, 323). [275⁵].

Formation.—1. By the destructive distillation of the control of the c

tion of bydrobenzsmide (Laurent, A. Ch. 19, 369), of amarine (Fownes, T. 1845, 263), or of tribenzylamine (Brunner, A. 151, 135). By the limited oxidation of amarine with chromite soid (B. Fresher a. Troschke, B. 18, 708), 2. By passing ammonia into a warm alcoholic solu-tion of a mixture of benzil and benzoic aldehyde: C.H., CO + C.H., CHO + 2NH, -

C,H,C,NH C,H,C,NH C,H,+311,0

(Radziszewski, B. 15, 1493; cf. Japp, B. 15, 2410). -4. By heating together p-oxybenzoic aldehyde, benzil, and ammonia, p-oxy-lophine, C2, H14 (OH) N2, is formed (v. Benzil, ammonia-DERIVATIVES OF, p. 466); and this, by distillation with zinc dust, is converted into lophino (Japp a. Robinson, C. J. 41, 323).—5. By warming oyanphenine, Callan, with acctic acid and zine dust, or by distilling it with potash and iron flings, it is converted into lophino and ammonia: $C_{11}H_{14}N_1 + 2H_2 = C_{21}H_{16}N_2 + NH_1$ (Radziszcwski).

Preparation. - Hydrobenzamido is hented in a retort until the more volatile products of its decomposition -hydrogen, ammonia, and toluene - have been given off. The residue, which can only be distilled at a high temperature, is treated with ether, and then dissolved in glacial acetic acid. From this solution water precipitates lophine, which is finally purified by recrystallising it from boiling alcohol (Radziszewski, B.

10, 70).

Properties .- Crystallises in very slender, colourless silky needles [275°] (Radziszewski). Distils without decomposition at a high temperature. The vapour-density agrees with the formula $C_n\Pi_{to}N_n$ (Fischer a Troschke). Insol. water, sl. sol. alcohol and other, Its solution in alcoholic potash phosphoresces when air is admitted, owing to a process of exidation, in which the lophine is slowly converted into henzoic acid and ammonia (Radziszewski). Feebly basic: the salts are partially decomposed by water, in which they are for the most part insuluble; but they may be crystallised from alcohol.

Salts .- (Laurent, A. Ch. 19, 369; Atkinson a. Gössmann, A. 97, 283; Brunner, A. 151, 135). CnH, NyHClaq: oldained by adding hydrochloric acid to an alcoholic solution of lophine. According to Laurent and Brunner this salt is anhydrous. - C21 II 18 N22 211 C1: formed by the action of gaseous hydrochloric acid upon lophine. (C_nH_{1s}N₂, HCl)_nP(Cl_s: rhombic plates. Contains 5H₂O (Brunner). — C_nH_{1s}N₂, HI. — C_nH_{1s}N₂, HNO₂. — With silver nitrate it gives rise to: CallisNa AgNOs; 2CallisNa AgNOs; and 2C21H16N2,3AgNO, (A. a. G.),

Reactions. -1. Oxidation with chromic acid in acetio acid solution converts lophino into

a mixture of benzamide and dibenzamide:

C₁H₁N₁+1(O + O₁=

C₂H₁CONH₂+(O₁H₂CO₂)NH

(Fischer a Troschet:—2. By the action of bromine upon lophine hydrobromide, an perbromide of the formula, CnH14N2Br4, IIBr(?), is obtained (F. a. T.). 8. Nitric acid yields, according to temperature and concentration, either diritiolopkine, $C_nH_{14}(NO_2)_2N_2$, or trinitrolophine $C_nH_{14}(NO_2)_2N_2$ (Laurent; Ekmann, A. 112, 161) .- 4. Heated with concentrated sulphuric acid to 1600-1700 lophine is converted into lophine-disulplumic substance is formed when benzoic aldehyde and C₁,H₁,N₂(SO,H)₂ (F. a. T.).—5. With is mixed with ammonium thio carbs mute is that indide at 100° it yields lophine hydriodide 2C,H₂O + CS₂ + 2NH₂ = C₁,H₁,N₂S₂ + 2H₂O (Malentine in the contraction of t

and distilliophesium todide, CaH., EtN., Ett and this latter compound, by treatment with moist silver oxide yields the corresponding base (Külın, A. 122, 826). (A. 122, 313) an isomeride of lophine,

C., H₁₆N., laq, is obtained, together with ordinary lophine, by saturating hydrobenzamide with gaseous HCl, and heating the mixture to 230°. Needles [170°], v. sol. boiling alcohol. Forms salts.

AZOBENZOILIDE, C., 71, N. (?) (Laurent, A. Ch. [8] 1, 306) and DIAGRED VILLIAMO, CHILLISMO (Robson, C. J. 4, 225), are also compounds which are stated to have been obtained by the action of ammonia upon benzoic aldehyde. They have been very little studied. BENZOIC ALDRITTLE, HYDEOCYANIC ACID, AND

AMMONIA.—A number of compounds are obtained when ammonia acts upon benzoie aldehydo containing hydrocyanic acid. Some of these are, however, more readily prepared from the hydrocyanides of hydrobenzamide (q. r.) and have therefore been described under that head. -1. Henroy'azotole, Chille N. Formed along with other products when a mixture of benzoic aldehyde, hydrocyanio acid, and ammonia is allowed to stand for some works (Laurent, A. Ch. [2] 66, 180; vide supra, 'Hydrobenzamide'). Crystalline puwder, consisting of minute rhombic prisms, sl. sol. boiling alcohol. By destructive distillation it yields amarone, C₁₀H₁₁N, together with lopline and other products. Amarone forms colourloss

needles [233], sl. sol. boiling alcohol (Lauront,

Revue Scient. 18, 207). 2. Azobenzoyl, CzilliaNa

is formed according to the equation:
3C.H.O.+.HON.+.NII. C...HI.N.+.3HI.O.
(Beilstein a. Reinecke, A. 136, 175). White
crystalline powder, insol. water, v. sol. cther. When warmed with alcohol and hydrochloria acid it is decomposed into HCN, marine, and a haso of the formula CitHieN, which latter crystallises in lamina, melting at 122° (Müllor a. Limpricht, A. 111, 140) .- 8. Benzhydramide. Cz.II , NzO, is formed, along with the foregoing, from the same generating substances according the equation; 3C, H,O+HCN+NH, = Cz,Ha,N,O+211,O, and differs from it by containing the elements of a molecule of water more. Migroscopic crystals. Sl. sol. alcohol,

Laurent a. Gerhardt, A. 76, 302). Banzoio aldenyde, ammonia, and suc-eduarted hydrodin. -1. Penzoic aldehyde and ammonima sulphide if mixed and allowed & stand for some weeks, yield this-benzaldin

v. sol. ether (Laurent, A. Ch. [2] 66, 180;

 $3C_1H_4O + 2H_2S + ...H_3 = C_{21}H_{19}NS_2 + 3H_2O$ (Laurent, A. Ch. [3] 1, 291; also [3] 36, 342 Deposited from other in monoclinio crystals [125°]. When hoiled with alcohol it gradually evolves sulphuretted hydrogen.-2. By the action of ammonium sulphide upon crude oil of bitter almonds Laurent obtained a compound C42113, N , S3 (?).

Benzoic aldehyde slowly reacts in the cold with CS, and NII, to form a compound, C₀, II₁₄N,S₂ (Quadrat, A. 71, 13). The same substance is formed when benzoic aldehyde der. A. 168, 238). Prismatic crystals, melting with decomposition at 100°. Cannot be dissolved in alcohol or ether without decomposition. The compound may be regarded as dibenzylidene - ammonium dithio - carbamats. NH, CS.S.N(CH.C.H.)". F. R. J. BENZOIC ALDEHYDE CARBOXYLIC ACID

W. ALGENYDO-BENZOIC ACID. BENZOIC ALDEHYDE GREEN v. Tetra-METHYL-DIAMIDO-TRIPHENYL-CARBINOL.

BENZOIC ANHYDRIDE C₁₄H₁₀O₂ i.e.
11 COLO Benzoul oxide. Mol. w. 226. (C.H.,CO)₂O. Benzoyl oxide. Mol. w. 226. [42°]. (360° i.V.). S.G. (liquid) ²² I·227. H.F. 104,815 (Stohmann, J. pr. [2] 36, 3).

Formation .- I. From BzCl and NaOBz, BaO, H₁C₂O₃ or K₂C₃O₄; or from NaOBz by the action of PCl₃, POCl₃ or S_cCl₂ (Gerhardt, A. Ch. [3] 37, 299; Wunder, J. pr. 61, 498; Heintz, P. 92, 458; Gal, A. 128, 127; Anschütz, B. 10, 1882). 2. From benzotrichloride (I pt.) and H₂SO₄ (3 pts. ol 95.4 p.c.) (Jenssen, B. 12, 1495).—8. By the action of the dry nitrates (6 mols.) of Pb, Ag, Ilg, Cu, or Zu upon benzoyl chloride (I mol.) (q. v.) (Lachowicz, B. 18, 2990).

Properties .- Trimetric prisms, insol. water, m. sol. alcohol and ether. Slowly converted into bonzoic acid by boiling water, more rapidly by alkalis. Hot NH, Aq forms benzamido and ammonium benzoate. Combines with brontine (I mol.). Gaseous HCl forms BzCl and HOBz.

Mixed Ashydrides. - Mixed auhydrides of the form BzOR are formed by the action of benzoyl chloride on the alkaline salts of various acids, or by the action of various alkoyl chlorides on sodium benzoate. They are generally decomposed by heat into a mixture of two anhydrides; and by water, more rapidly by alkalis, into two acids.

Benzo-acetic anhydride R.O.B. v. ACRTO-BENZOIC OXIDE, p. 17.

Benzo-isovaleric anhydride Bz.O.C,11,O. Oil (Chiozza, A. 81, 108).

Benzo-heptoic anhydride Bz.O.C.H.O. Oil. S.G. 41-013 (Chiozza a. Malerba, A. 91, 102).

Benzo-pelargonic anhydride BzO.C, H., O. Gil (Chiozza, A. Ch. [3] 89, 209). Benzo-myristic anhydride

BzO.C, II., O. [380].

Benzo-stearic anhydride BzO.C, II, O. [70°] (C. a. M.).

Benzo angelic anhydride BzO.C.II.O. Oil.

Benzo-cinnamic auhydride

Bz.O.O, II.O. O.I. 43, G. 23 1-184. Benzo-cuminic anleydride Bz.O.C., H., O. Oil. S.G. 23 1-115.

BENZOIC BROMIDE & BENZOYL BROMIDE. BENZOIC CHLORIDE v. BENZOYL CHECKIDE.

BENZOIC CYANIDE P. BENZOYL CYANIDE. BENZOICIN v. Tri-benzoyl-GLYCENIN.

BENZOIC OXIDE C. BENZOIC ANHYDRIOE. BENZOIC PEROXIDE C. BENZOYL PEROXIDE. **BENZOIN** $C_{14}H_{19}O_3 = C_8H_3.CH(OH).CO.C_8H_3$

Phenyl-benzoyl-carbinol. [137°]. First prepared by Stange (R. P. 16, 93), and almost simultaneously by Robiquet (A. Ch. [2] 21, 254), by acting with a solution of potassium hydroxide or barium hydroxide upon crude oil of bitter almonds containing HCN.

Formation .- 1. By partial oxidatiou

hydro - benzoin, C,H, CH(OH).CH(OH).C,H, (Zinin, A. 123, 128).—2. By treating benzii C,H, CO.CO.C,H, with zine and alcoholic hydrochleric acid (Z., A. 119, 177), with acetic acid and iron filings, or with K2S.

Preparation .- 200 g. of pure benzoic aldehyde are heated for a short time with a solution of 20 g. of KCN in 800 g. of 50 p.c. alcohol, and the liquid allowed to cool. Benzoin separates and is removed by filtration. The filtrate, on heating with more KCN, yields a fresh quantity of benzoin (Zincke, A. 198, 151). Two mois. of benzoie aldehydo unite to form I mol. of benzoin: 2C, H, CHO = C, H, CH(OH). CO.C, H, The action of the KCN is not understood.

Properties .- Colourless, lustrous, six-sided prisms. Sl. sol. boiling water. V. sol. hot al-cohol; sl. sol. in cold. Conc. H₂SO₄ disselves it with a violet colour.

Reactions .- I. Partially decomposed by distillation. Repeated distillation breaks it up for the most part into 2 mols, of benzoic aldehyde; a smaller portion yields benzil and deoxybenzoin: ${}^{2}C_{11}H_{12}O_{2} = C_{11}H_{16}O_{2} + C_{11}H_{12}O + H_{2}O$ (Zinin, B. 6, 1207).—2. Treated in alcoholio solution with zinc and hydrochloric acid it is converted into deoxybenzoin (Zinin, A. 126, 218); nt the same time deoxybenzoin-pinacone, CaH21(OII), and hydrobenzoin are formed (Goldunberg, A. 174, 332). Sodium amalgam reduces it to hydrobenzoin. Heating with cone. hydriodic acid for some hours to 130° converts it into di-benzyl C₁₄H₁₁ (Goldenberg). By distillation with zinc-dust it yields stilbene, Culle, together with an oily hydrocarbon isomeric with stilbene (Jena n. Limpricht, A. 155, 90). -3. Nitric acid oxidises it to benzil (Zinin); but chromic acid, or potassium permanganate, converts it into benzaldehyde and benzoic ucid (Zincke, B. 4, 839). It reduces Fehling's solution in the cold, a property common to all compounds containing the group CO.CH(OII) (E. Fischer, A. 211, 215) .--4. Pusion with caustic potash converts it into benzoic acid with evolution of hydrogen (Liebig a. Wöhler, A. 3, 276). When distilled with sodatime, the benzoate which is first formed is decomposed by the excess of sodu-lime, and benzene is obtained (Jena a. Limpricht). The action of atcoholic potash is complex: when benzoin and alcoholic potash are heated with access of air, benzoio acid, a small quantity of benzilie acid (C₁₁H₂₀O₃), benzon ether (C₂H₂₂O₃ [157°]), and a compound C₂₄H₁₂O₄ (to which Limpricht a. Schwanertgave the name othyl-dibenzoin, assigning to it the formula C to H to O1) are formed—this last, however, is produced from benzil generated by the air-exidation of the benzoin (v. Benzil). If the benzoin is heated with alcoholic potash in a sealed tube at 100° the products are benzoic acid, hydrobenzoin, and ethylbenzilio acid (C_nII₁O₄). With very concentrated alcoholic potash at 160° benzoin yields henzoic acid, stilbene, a compound C_nII_nO_n and a small quantity of ethyl-benzilic acid. By heating benzoin with a solution of sodium cthylate in alcohol, ethyl-benzoin, C14H11O2.C2H, [95°], is formed, together with the various products already mentioned. Prisms, with a vitreous Instro, v. sol. alcohol (Jena a. Limpricht, A. 155, 89; Limpricht a. Schwanert, B. 4, 836; Japp a. Owens, C. J. 47, 90).—5. Chlorine acts like nitric acid, converting bensoln into bensil (Laurent, A. Ch. [2] 59, 401).—6. When heated with Imming hydroschlorio acid at 130° for 6 or 8 hours it yields lepidene, C₂₀H₂₀O (g. v.), benzil, and a thick yellow oil (Zinin, J. pr. 101, 160).—7. When boiled with dilute sulphuric acid it parts with tha elonients of water, yielding oxylepidene:— 2C₁H₁₂O₂ = C₂H₁₂O₃ + 2H₁O (Limpricht a. Schwanert, B. 4, 335). Concentrated sulphuric acid converts it into benzil (Zinin). 8. Heated with alcoholic ammonia for some hours at 100° in scaled tubes it yields benzoinam, C₂H₁₂N₁O (silky needles, v. sl. sol. alcohol, melling with decomposition), benzoinalam, C₃H₂NO₂ / [199°] (granular crystals, sl. sol. alcohol), and tetra-C₃H₁, C—N—C₄H₁

phenyl-azinc, | | | [246°], to-

gother with some lophine, Callin Na. Tetraphenyl-azine is best prepared by heating benzoin with ammonium acctate until the salt is volatilised, dissolving the product in the strongest alcoholic hydrochloric acid, and ppg. with alcohol. V. sl. sol, alcohol, v. sol, alcoholic hydrochloric acid, v. sol. boiling benzene, sal. with blood-red colour in cold code, 11,80, Sublimes without decomposition. Heating with sodalime converts it into tetraphenylene - azine C₂,H₁₀N₂ (Laurent, A. Ch. [2] 66, 181; Erdmann, A. 135, 181; Japp a. Wilson, C. J. 1886, 825; Jupp a. Burton, C. J. 1886, 813; 1887, 98). — 9. Bouzon reads with the primary anines of the benzenoid series, when heated with them to 200°, climinating 1 mol. of water and generating feelily basic compounds which by boiling with acids are decomposed into with actors are decomposed in their generating substances. Anilhanziin C_aH_x,CH(OH).C(N.C_aH_x), [OP²], from unifine and benzoin, forms yellowish needles, v. sol. most organic menstrna. Yields with nitrons acid nitroso anilbenzon C_all_aCH(OH).C(N.C_aH_aNO).C_aH_b (140°), with acetic anhydride a monacetyl-derivative (153°), and with bromine a monobromo - derivative [168°]. Sodium amalgam reduces anilbenzoin in alcoholic salution to hydrobenzoin anilide, C₆H₅.CH(OH).CH(NI1.C₆H₅)C₆H₅ [119°], which forms with sulphuric acid a salt not decomposed by boiling with the dilute acid. - p. Tolilbenzoin C_aH, CH(OH).C(N.C_aH, CH₄),C_bH, [I442] resembles in ils properties and its behaviour towards reagents the aniline compound. It. yiolds with nitric acid a mono nitro derivativo [125°] and a di-nitro derivativo [195°]. 8. Naphthilbenzoin [130] also resembles the aniline compound (Voigt, J. pr. 34, 1).—10. Benzoin reacts, with hydroxylamine and with phenyl-hydrazine .-- Benzoin-oxim, C,H,,CH(OH).C(N.OII).C,H, [\$12-1522]. An alcoholio solution or benzoin is mixed with an aqueous solution of hydroxylamine and allowed to stand for a week. Microscopic prisms, soluble in benzeno (Wittenberg a. V. Meyer, B. 16, 504). -- Benzoin phenylhydrazide, C.H. CH(O11).C(N.H.C.H.).C.H. [1552]. Benzoin and phenylhydrazino, together with a little alcohol, are heated at 100°. Needles, sol. benzens (Pickel, A. 232, 223).—11. By heating with aci-chlorides the hydroxylic hydrogen of benzoin may be replaced by acid radicles to form ethereal malts. Benzoin acetate, C.H. (C.H.O)O. [75°],

From benzoln and acetyl chloride. Monoclinie prisms or tables, v. sol. ether and alcohol (Zinin, A. 104, 120; Jena a. Limprloht, A. 155, 92).—Benzoln benzoate, C.,H.,I.(C.H.,O)O, (125°). By warming benzoin with benzoyl chloride. Slender needles. Sol. hot alcohol. Yields a mono-nitro-compound [137°] (Zinin).—Benzoln succinate, (C.,H.,I.O.),C.,II.O., [129°]. Hy heating benzoin with succinyl chloride to 100°. Leaflots from alcohol. Sol. also in ether and CS, (Lakanin, B.5,331).—12. When benzoin is heated with hydrocyanic acid and alcohol to 200° the process of its formalion is reversed and it is broken up into benzoic aldehyde. A part of the benzoic aldehyde undergoes a further change, yielding amongst other products ethylic henzoate (Michael a. Paluer, Am. 7, 192). 13. Phenyl cyanate forms Ph.CH(O.CONPh11).CO.Ph [163°] (Gumpert, J. pr. 12: 32, 280).

BENZOIN, GUM. A resin which flows from

BENZOIN, GUM. A resin which flows from the bark of Styrax benzoin, a tree growing in Sumatra, Borneo, Java, and Siam. Gum benzon contains, besides various resins, benzoid acid and, frequently, cinnamic acid. Siamese and Palemhang henzoins are free from cinnamio acid. Potash fusion produces henzoic, p-oxybenzoic, and protocatechine acids, and pyrocatechin. Distillation with zine-dust gives toluene and a little o-xylene, naphthaleno, and methyl-multithalene.

References.—Unverdorhen, P. 8, 397; Van der Vliet, A. 34, 177; Kopp, G. B. 19, 1262; Kolbe a. Lautemann, J. 115, 143; 149, 136; Deville, A. Gh. (3) 3, 192; Ashoff, J. 1861, 400; Wiesner, J. 1872, 1050; Theeparten, J. 1874, 922; Chamician, B. 11, 274; Saulfeld, A. Ch. [3] 16, 280.

BENZOIN DI-p-CARBOXYLIC ACID

C₁₀H₁₂O₁₆, C₆H₄(CO₂H), CH(OH), CO, C₆H₄(CO₂H), Fermed by exidation of dip-addelyda-benzom with KMnO₁. Short felted needles, Sublimable, Infusible, Ag₂A". Dimethyl ether McA": [126] (Oppenheimer, B. 19, 1816). BENZOLEIC ACID C₂H₁₀O₂ Hydrobenzoic

BENZOLEIC ACID C.11_{in}O₂ Hydrobenzoic acid. Formed, together with benzyl alcohol, by the action of sodium amalgam on an aqueous solution of benzoïc acid kept acid by HCl (Hernann, A. 132, 75; Otto, A. 134, 303). Oil, smelling of vuleric acid, heavier than water, v. sol. alcohol and ether. — EtA': oil.

BENZQLINE. A mixture oI paraffins (hoxane, heptane, octane) boiling between 70° and 100° obtained by distilling petroleum or paraffin oil. The mixture is also called petroleum spirit or ligioin.

BENZOLON. Pdentical with lophine (v. BENZIL).

BENZODIMETHYLANILINE v. DIMETHYL-AMIDG-BENZO-DI-METHYL-DI-FURFURANE, m. BENZO-DI-METHYL-DI-FURFURANE, C₁₂II₁₆O₂ i.e.

$$HC \stackrel{(1)}{\underset{(2)}{\leftarrow}} C_{\bullet}H_{2} \stackrel{(5)}{\underset{(6 \text{ or } 4)}{\leftarrow}} CH.$$

[e. 27°]. (270° at 720 mm.). Obtained by evaporating (β) benzo-di-methyl·di-IurInrane-di-carboxylic ether C_eH₂(

CMe

CO₂Et)₂ with alcoholio KOH and lime, and dry distillation. Prisms. By warming with conc. H₂SO₄ a pure

bine colouration is produced (Hantzsch, B. 19, 2988; 20, 1887).

p-Benzo-di-methyl-di-furfurane

 $0_{\rm eH_2}(<_{\rm O}^{\rm CM_0})_{\rm cH})_{\rm r}$ [108°]. Obtained by heating the potassium salt of the di-carboxylic acid with lime. Large pearly tables with bluish Auorescence. V. sol. alcohol, ether, etc. (Nuth, B. 20, 1337).

Benze-tri-msthyl-tri-furfurane C15H12O3 i.c. C.(<CMc>CII), [115°-120°]. Needles. Very soluble in ordinary solvents. Formed by

evaporating the tri-carboxylic ether C(CMe C.CO,Et), with alcoholic KOII and

lime, and dry distillation (Lang, B. 19, 2936) o-BENZO. D1 . METHYL - DI -FURFURANE DI-CARBOXYLIC ACID

CMc C.CO.11)2. Obtained by saponification of the ethyl-ether which is formed by dissolving in cone. II SO, the product of the action of chloracetoacetic ether (2 mols.) upon di-sodium pyrocatechin (1 mol.). Amorphous solid.—BaA" 2aq.

Di-ethyl ether Ft.A": [155°]; short prisms

(from nleohol), or long white needles (from ether) (Nuth, B. 20, 1337).

(a)-m-Bsnzo-di-msthyl-di-furfarane di-carboxylle acid C, 11,00, i.e.

HO,C.C CMO ^{Мо}>С, H₂<СМе >С.СО, П. (4 or 6)

above 310°]. Formed by suponification of the di-ethyl-ether. This ether is obtained, together with a much larger quantity of the (3)-isomerido by the action of chloro acetacetic ether upon dicodinm resorein Calla(ONa)2 in presence of alcohol, extracting with benzene, dissolving the undissolved portion in conc. H.SO, pouring into water, and extracting with other; when the othereal solution is mixed with hot alcohol and allowed to cool the (a)-ether crystallises out, whilst the (3) other remains in solution. The two others are also formed (the a in very small quantity) by heating oxy-methyl-commarilic ether

C.H. (OH) CMc C.CO.Et with chloro-aceto-

acotic other and alcoholic NaOEt, and dissolving the product in H_sSO. The acid is a white microcrystalline solid. Scarcely soluble in water, more readily in alcohol. The salts of the heavy metals are all sparingly soluble.

Warm Il, SO, produces a pure blue colouration.

Diethyl ether A"El, [1869]; needles; al. sol. alcohol (Hantzsch, B. 19, 2930). (\$)-m. Benzo di . msthyl-di-furfurane-di-car-

above 810°]. Formed by saponification of its di-ethyl ether. Microcrystalline solid. Scarcely soluble in water, more easily in alcohol. Gives sparingly soluble pps. with the salts of the heavy metals. The soid and its other give a pare blue colouration with warm H.SO.

Disthyl sther R'Et .: [1419]; small white needles; more soluble than the (8)-isomeride. Formed, together with a small quantity of the (B)-ether, by the action of chloro-acetacetic ether upon dry di-sodium resoroin, extraction with benzene, dissolving the residue in conc. H.SO., pouring into water, and extracting with ether. Also by heating oxy-methyl-coumarilic ethor

 $C_vH_3(OH) < CMC O_vEt$ with chloro-acetacetic

ether and alcoholic NaOEt, and dissolving the product in H.SO. By evaporation with alco-holic KOH and lime and dry distillation it (β) - benzo -di - methyl - di - furfurane yields C.H. (< CMo > CH). (Hautzsch, B. 19, 2930).

p. Bsuzo-di-msthyl-di-furfurane di-carboxylio Obtained by saponification of its, ethyl-ether which is formed by dissolving in conc. 11.SO, the product of the action of chloro-nectacetic ether (2 mols.) upon di-sodium hydroquinono (1 mol.).

Amorphous solid (containing aq).
Salts.—Ag.A": white pp.—BaA" 2aq: sl.

sol. yellowish white powder.

Ethyl ether Et.A": [150°]; glistening greenish plates; v. sl. sol. all solvents (Nuth, B. 20, 1334)

Bsuze-tri-methyl-tri-furfurane-tri-carboxylic acid C18H12O0 i.e. C. (< CMc > C.CO2H),

Formed by saponification of its ethyl other which is obtained by treating dry powdered tri-sodiumphloroglucol (1 mol.) with chloro-acctactio ether (3 mols.), dissolving the product in H₂SO₄ and pouring into water. Gelatinous pp. (containing aq). Scarcely soluble in alcohol and ether. Its salts are mostly insoluble and gelatinous. A", 7aq: microcrystalline. Ethylether Et, A": [c. 298°]; small white Bn,A"

glistening needles; sl. sol. all solvents, most rendily in chloroform (Lang, B. 19, 2935).

BENZONAPHTHONE r. NAPHTHOQUINONE.

BENZONAPHTHYL-THIAMIDE v. Thiobenroyl (a) NAPHTHYLAMINE

BENZONITRILE C,H,N i.e. C,H, CN. Mol. w. 103. [-17] (Hofmann). (190.5°). S.G. 2 1.023; H 1.008 (Kopp, A. 98, 373). S. 1 at 100°. S.V. 123.7 (Rnmsay).

Formation .- 1. By the dry distillation of amnionium benzoate (Fehling, A. 49, 91).—2. From benzamide by heating it alone or with CaO (Anschütz a. Schultz, A. 196, 48), BaO (Wöhler, A. 192, 362), P.O. (Hofmann a. Buck-(Woller, A. 192, 502), 4.0. (Holmann a. Duorston, A. 100, 155), PCl. (Hencke, A. 106, 276), or Pr.S. (Henry, A. 2, 307).—3. By heating hippuric acid alone (Limpricht a. Uslar, A. 88, 133), or with ZnCl. (Gössmann, A. 100, 74).—4. By the action of BzCl or Bz_O on benzamide.— 5. By the action of BzCl on oxamide, on potassium sulphocyanide (Limpricht, A. 99, 117), or ou potassium oyanate (Schiff, A. 101, 93) .-- 6. By heating Bz.O with potassium cyanate or sulphocyanide.-7. By the action of HgO on thiobensamide. - 8. By heating benzoic acid with sulphocyanide of lead (Krüss, B. 17, 1767), or of potassium (Lotts, B. 5, 673).—9. From potassium benzoata and avenue.

4. OA. (8) 52, 200).—10. By distilling a mixture of smillne and oxalic acid (Hotmann, C. R. 64, 888).—11. Formed by distilling formanilide over sine-dust; the yield is usarly 20 p.c. of the formanilide (Gasiorowski s. Merz, B. 17, 73; B. 18, 1001).—12. Formed together with sodium formate, by the action of dilute NaOH upon anilino di chloro-acetate (Cech a. Schwsbol, C. C. 1877, 134) .- 13. Formed by running an aqueous solution of diazobenzene chloride into a hot solution of Cu₂(CN)₂ (Sandmeyer, B. 17, 2653).— 14. By heating tri-phenyl phosphate with potassium oyanide or ferrocyanide; the yield is 25 p.o. of the theoretical (Scrugham, 4, 92, 318; Heim, B. 16, 1771).--15. By henting potassium benzeno sulphonate with KCN (Merz, Z. 1868, 83).-16. From K.FeCy, and chloro or bromobenzene at 400° (Merz a. Weith, B. 8, 918; 10, 749) .- 17. From iodo benzono and AgCy (Merz a. Soholnberger, B. 8, 1630). -18. Together with tsrephthalonitrile, by passing a mixture of benzeno and cyanogen through a red-hot tube (M. a. S.). - 19. By phssing dimethylaniline through a red-hot tube (Nietzki, B. 10, 474). 20. By the action of CyCl on benzene in presenes of aluminium obloride (Friedel a. Crafts, Bl. [2] 29, 2). -21. From bromo-benzene, Cy (Cl., and Na (Klason, J. pr. [2] 35, 83) .- 22. By boiling phenyl thiocarbimide with finely divided Cu .-23. From phenyl carbamine by intramolecular changs at 240° (Weith, B. 6, 213).—24. By the action of acetic anhydrids on bsnzaldoxim (Lach, B. 17, 1571).

Properties.—Colourless oil, smelling of almonds; sinks in cold, but swims in hot water;

miscible with alcohol and other.

Reactions .-- 1. Cold aqueous potash has no action, but on boiling it forms NII, and KOBz; dilute acids not similarly .- 2. Heating with polassium gives KCy, cyaphsnino, (C_a11₃),Cy, and other bodies (bingley, Chem. Gaz. 1851, 829; Hofmann, B. 1, 198). When the boiling alcoholio solution is treated with sodium, the greater part is saponified whilst a smaller portion undergoes reduction to bonzylamino and to benzeno (Bamberger a. Lodter, B. 20, 1709).-3. Il,S or ammonium sulphide forms thiobenzumide.-4. Zn and HCl in alcoholio solution form mono-, di-, and tri-benzylmmino (Mendius, A. 121, 129; Spica, G. 10, 515). -5. Furning H.SO, forms, on heating, benzene sulphonio acid; at 20° it forms cymphonine. Benzonitrile (10g.) cooled with ice and trented with tuming H.SO, (7 g.) added slowly, forms di-benzamide NHBz, [148°], and 'benzimide-benz-amide 'NHBz,C(NH),C,H, [100°], called by Pinner a. Klein 'benzimide-bonzoate' nnd dibenzimido-oxide, respectively. Dilute 11Cl converts the former into the latter (F. Gumpert, J. pr. [2] 30, 87; Pinne., ibid., 125). - 6, Beried with sinc ethyl it gives off ethane (1 vol.) and an olefins (I vol.) and forms a product, whence, by treatment first with alcohol, and then with aqueons HCl, eyaphenine (q. v.) and the hydro-chloride of a base which crystallises in six-sided plates, C₁H₁, N₂Cl, is formed. This body (387°) is readily soluble in alcohol, but sparingly o in water. Potash liberates the bass as a sologriess oil (Frankland a. Evans, C. J. 37, 566) .- 7. By exhaustive chlorination with SbCl. yields penta-chloro-cyano-benzene C.Cl.(CN) (Mers a. Waith, B. 16, 2835).—S. With mathetic and H.SO, it gives the bensoyl derivative of methylena diamine (q. v.).—9. With diphenylomine hydrochloride at 180° it forms al-phenylomine hydrochloride. Think prisms (from bedzene), containing benzsno of crystallisation; yollow tholtes (occasionally from benzono); long thin prisms (from alcohol). The tablets are monoclinio, above 5875:1:5014, L=51° 23' (Bodewig). Solubls in benzene and ether, slightly in alcohol. Its alcoholic solution is neutral. Acetyl chloride does not net on it, Salts: BHICI. Narrow red prisms. [Above 220°.]—(B'IICI).PtCl.

Combinations. - I. With metallic ohloridss.— (C,H₃N)₂AuCl₁.— (C,H₁N)₂PtCl₄.— (C,H₃N)₂SnCl₄.—(C,H₃N)₃TiCl₄ (tlenke, A. 100, 284).—2. With halogons. C₄H₄CBr:NBr.— (C,11,CN) Br₂: needles (Engler, A. 133, 137).— 3. With hydrogen obloride: C,11,N 2HCl (Pinner a. Klein, B. 10, 1891; cf. Gerhardt, Traité, 4, 762). - 4. With hydrogon bromido. C.H.N2IIBr. [70] (Engler, A. 149, 307). — 5. With alcohols. —Bonzimido. ethyl ether, C.H. C(OEt):NII. The hydroeliloride, B'HCl, is formed by passing dry HCl gas into a mixture of ethyl alcohol and bonzo. nitrile, diluted with other. Large glistening prisms; decomposes on heating to about 120 into ethyl ohlorido and benzamide (Pinner, B. 16, 1654). Benzimido isobutyl ether C.11,-C(OO,H,):NII. The hydrochloride B'2HOI is formed by passing l1Cl into a cooled mixture of benzonitrile and isobutyl alcohol (Pinner a. Klein, B. 10, 1890); it gradually loses HOl bocoming B'HCl, {135°},—B',11,1'tCl_a—B'H,3O_c—6. With acids.—Benzimido acctate C_aH_a,C(OAc):NH. [116°]. From benzimido isobutyl ether and Ac,O (Pinner a. Klein, B. II. 9). - 7. With mercaptans. - Benzimido. ethyl thio-ether C.H. C(SEt):NH. From benzonitrile, mercaptan, and HOl, or from thiobenzamido and Etl (Berntlisen, A. 197, 848). Oil; decomposes readily into merenptan and benzonitrile, -- B'HCl. [188°]. -- B'₂H₂PtCl₂.--B'HI. [142°]. Benzimido-isoamyl thio, ether O₂H₂C(SC₂H₃):NH. Tho hydrochlorido, B'HCl, is formed by passing HCl into a mixtura of benzoni'rilo and isoamyl mercaptan (Pinner a Klein, B. 11, 1825). Tho free base is an oil. Benzimido - benzyl - thio - ether C.H. C(SC,H.):NH. Prepared like the ethyl ether. - B'HCl [181°].

Derivatives of Benzonitrile are described as BROMO-, NITHO- &C. BENZONITRILE.

BENZO-PHENOL v. OXY-BENZOPHENONE.

BENZOPHENONE C.; II., O i.c. C. H., CO.O. II., Di. phenyl. ketone. Mol. w. 182. [48°]. (806° i.V.),

Formation.—By the dry distillation of calcoim benzoate (Peligot, A. 12, 41; Chancel, A. 72, 279).—2. From BzCl and ligPh₂ (Otto, B. B. 197).—3. From BzCl and benzeno in prosence of Al₂Cl₂.—4. From benzeic acid, benzens, and P₂O, at 190° (Kollarits a. Merz, B. 6, 446, 538).
5. From benzeno, COCl₂, and Al₂Cl₄ (Friedel, Crafts, a. Ador, C. R. 85, 673).—6. By oxidation di-phenyl-methane (Zincke, A. 156, 377).

Preparation.—From BzCl, O₂H₂, and Al₂Cl₄

the yield is 70 p.c. of the calculated (Elbs, J. pr. [2] 85, 465).

Properties .- Prisms; insol. water, v. sol. alcohol and ether.

Reactions .- 1. Reduced by HI to di-phenylmethane (Graebe, B. 7, 1624).—2. Reduced by sinc-dust to di-phonyl-methane, tetra-phonylethylene, and tetra-phonyl-ethano (Staedel, A. 191, 307) .- 3. Potash fusion gives benzoic acid and benzene .- 4. Reduced to di-phonyl-carbinol by sodium-amalgam or by heating with alcoholic potash. - 5. Animonia has no action .- 6. Zn and alcoholic H.SO, reduce it to benzpinacone and (a) and (b) benzpinacolin (Zincke a. Thorner, B. 11, 1396) .-- 7. AcCl in presence of zinc-dust acts on an othereal solution forming crystalline (α)- and (β)- benzpinacolin.—8. PCl, forms di-chloro-di-phonyl-methane.—9. Passage through a red hot tube slightly decomposes it; the product contains benzene, di-phenyl, and p-di-phenyl benzene, while gaseous carbonic oxide, hydrogen, and acetylone escape (Barbier g. Ronx, C. R. 102, 1559),-10. Whon heuted with ammonium formute at 200° 220° it yields the formyl derivative of di-phenyl-carbinylamine (Calla), Cl1.NH.CHO (Leuchart a. Bach, B. 19, 2129). -- 11. P.S. at 100° forms C. H. S. [153], crystallising in histrous flat monoclinic needles. At 200° it turns deep blue (Japp a. Raschen, C. J. 49, 481). - 12. P.S. at 140° forms C₂₆H₂₆P₂S₅ [227°], crystallising in minute plates, insol. alcohol, sl. sol. bot benzene. On melting it turns deep blue. It is exidised by CrO, in

HOAc to benzophenone (J. a. R.).

Oxim (C.H.), C:NOH. Di-phenyl-kctorim.
[140]. Prepared by boiling an alcoholic solution of benzophenone (30g.) with hydroxylamine hydrochloride (28g.) and a little HCl for a day (Beckmann, B. 19, 988; Janny, B. 15, 2782). Silky needles, v. sol. ether and nectone, m. sol. benzene and ligroin, v. sl. sol. cold water. Sol. acids and ulkalis. Resolved by acids into its constituents. By PCI, or POCI, it is converted into w-ohloro-benzylideno-aniline, C,H .N:CCl.C,H, produced by intramolecular change from (C.H.), C:NCl (B.). By warming with cone. 11,80, to 100° it is converted by similar isomerio ohango into benzamlide (Beckmann, B. 20, 1507). Salts. - Cullio N(ONa): crystalline powder .-C14H10N(OII), IICl: white powder.

Methyt.oxim Cullin N(OMe) : (92°]; yellow crystals.

Ethyl-oxim C15H10N(OEt): (276°-279°);

Benzyt.oxim CisliaN(OC,H1): [56°]; white crystals.

Acetyl oxim C₁₃H₁₀N(OAc): [55°]; white crystals (Spiegler, B.•17, 810; M. 5, 203).

Phenyt hydrazide Ph₂C:N₂HPh. [137°]. Got by boiling benzophenone with phenyl hydrazine and alcohol; or by heating the oxim with phenyl hydrazine, N₂ and NII, being evolved. Needles (from alcohol). Insoluble in water, not very soluble in alcohol. Heated for some time with diinte (20 p.e.) HCl, it is resolved into benzephenone and phenyl hydrazine (Pickel, A. 232, 228; Fischer, B. 17, 576; Just, B. 19, 1206).

Isomeride of Benzophenone. (26°). (305°). Sometimes formed in oxidising di-phenyl-methane or in distilling caloium acetate with

calcium benzoate (Zincke, A. 159, 867). Readily changes into ordinary benzophenone

Derivatives of benzophenone are described as AMIDO-, BROMO-, CRLORO-, CYANO-, NITEO-, OXY-, &c., BENZOPHENONE, and as Di-PHENYLENE-KETONE

BENZOPHENONE-CARBOXYLIC ACID v. BENZOYL-BENZOIC ACID.

BENZOPHENONE-DI-p-CARBOXYLIC ACID [4:1] C_sH₄(CO₂H).CO.C_sH₁(CO₂H) [1:4]. Formed by boiling di-p-cyano-benzophenone with alcoholic KOH (Brömme, B. 20, 522). Also by the oxidation of di-tolyl-methane, or di-methylbenzophenono, with chromic mixture (Weiler, B. 7, 1185; Ador a. Crafts, B. 10, 2173). Microscopic needles. Sublimes at a high temperature without melting. Sl. sol. alcohol, benzene, and ether. S. (hot water) = 002. The NH salt gives pps. with salts of Fe, Co, Cu, Ba, and Ca, but not with salts of Pb, Cr, Zn, Mg, and Ni.

Silver salt A"Ag, Ag,O: insol. water. Di-methyl etha: A"Me,: [138°]; large ncedles (B.). Ag.A' (A. a. C.).

Benzophenone dicarboxylio acid

C.H. CO.Ĉ. II. (CO.II)... Benzoyl-isophthalic acid: [280°]. From benzyl-isoxylene and chromic mixture (Zincke, B. 9, 1762). St. sol. hot water and CHCl, v. sol. alcohol. Converted by Zu and HCl into the luctous of C. H. C(OH)H. C. H. (CO.H)g-Salts: CaA"aq.—BaA"aq.—Ag.A'

Methylether McA". [118]. Ethylether Et A". [95]. Benzophenone dicarboxylic acid

C₆11,.CO.C₆11₃(CO.H), [1.2.5]. Benzoyl-terephthalic acid. [290°] (W.);

[285°] (E.). Formation .- 1. By oxidising benzyl-cymene

with chromic mixture (Weber, J. 1878, 402) .-2. From benzoyl-p-xylene and 11NO₃ (S. G. 1·15) at 170° (Elbs. J. pr. [2] 35, 479).—3. From phenyl p-cymyl ketone and dilute 11NO₃. Properties.-V. sl. sol. water, v. sol. alcohol.

Reduced by Zn and HCl to

C_sII_sCEI(OII),C_sII_s(CO_sH)_s.—CaA" aq.— BaA" 5aq.—A g.A". Methyl ether Me_sA". [101°]: nocdles. Ethyl ether Et_sA". [101°]: prisms.

Bonzophonone tetra-carboxylic acid

C.H. CO.C.11(CO.H). From benzoyl iso durens and KMnO. (Essner a. Gossin, Bl. [2] 42, 170).

BENZOPHENONE CHLORIDE is exo. Dr-Chloro-di-Phenyl-Methane (q. v.).

BENZOPHENONE OXIDE v. Di-PHENYLENS. KETONE OXIDE.

BENZOPHENONE SULPHONE C, H.SO, i.e. $SO_2 < \frac{C_s 11}{C_s 11} > CO$. [187°]. From benzophenone and fuming H.SO. (Beckmann, B. 6, 1112; 8, 992). V. sol. other, sol. alkalis; converted by water at 190° into an isomeride (?) [175°].

BENZOPHENONE DI-SULPHONIC ACID C₁₃H₁(SO₁H)₂O. From benzophenone and fuming H₂SO₁ by warming (Staedel, A. 194, 314). Converted by potash fusion into phenol and poxy benzoic acid. - BaA"

Chloride CO(C.H.SO,Cl), [122°] (Beck. mann, B. 8, 992).

BENZOPHENYL- v, BENZOYL-PRENTI-

RESTOPMOSPHINIC ACID is CARDOTY-RENEOPINACOLIN is BENTPINACOLIN (q. v.).

RENZOPINACONE is BENZPINACONE (q. v.). BENZOQUINOL is HYDROQUINONE (q. v.). BENZOQUINONE is QUINONE (q. e.). BENZORESORCIN v. DIOXYBENZOPHENONE. DI-BENZO-RESORCIN v. DIOXYPHENYLENE-

DI-PHENYL-DI-RETONE BENZOSTILBINE is lophine (v. p. 474).

BENZO-SUCCINIC ACID v. BENZOYL-SUCCINIC

BENZOTHIAMIDE P. THIO-BENZAMIDE. BENZO-TOLUIDINE C. PHENYL AMIDO-TOLYL

BENZTROPEINE e. BENZOYL PROPEINE. BENZOXAMIDINE v. BENZAMIDONIM. BENZ-OXIMIDO-AMIDE & BUNZAMIDOXIM. BENZ-OXIMIDO-ETHYL-ETHER C.H., NO. i.e. C.H., C(OEt); NOII. Formed by

the action of hydroxylamine hydrochloride on benz-imido-ether (c. Benzonium.r), (Pinner, B. 17, 181). distillution. Colourless Suid. Decomposes on

BENZOXY- v. Benzoyt-Oxy-

BENZOXY-PROPIO-CARBOXYLIC ACID v CARBOXY-BENZOYS-PROPIOSIC ACID.

BENZOYL. The radicle Call CO. Benzoyl derivatives obtained by displacement of H in amidogen, imidogen, or hydroxyl, are described under the compounds from which they are thus dsrived.

Di-benzoyl is called Brazu. (a. r.) BENZOYL - ACET - CARBOXYLIC ACID v. ACETOPHENONE 161-CARROXYLIC ACTO.

BENZOYL-ACETIC ACID CHIO, i.e. C.H. CO.CH .CO.H. Acetophe..one w-carborylic acid. [104"].

Formation. I. From the other by leaving it 24 hours with cold dilute KOH (3 per cent.), oooling to 0° and then adding dilute II SO. The acid is then ppd. as white flakes (Baeyer a. Perkin, B. 15, 2705; 16, 2128; W. 11. Perkin, jun., C. J. 45, 176). 2. From phenyl propiolic acid and cone. II, SO, the solution being poured upon ice .- 3. From the ether by allowing it to stand for 14 days with 20 vols. of conc. 11 SO, and then pouring upon ice (Perkin, C. J. 47, 210).

Properties. Minuto needles, which polarece light (from benzens at 70° containing a little light petroleum). At 104° it melts, and pives off CO.. Sl. sol. light petroleum, v. sol. alcohol, ether, hot benzene, and hot water. Fe Cla colours its alcoholic or aqueous solutions reddian-

Reaction .- I. Heated alone or with dilute H.SO, it gives acctophenone and CO,-2. The ammonium salt gives with AgNO, a pp. of AgA';

with Fe₂Cl, a blackish-violet pp; with FeSO, no pp.; with CuSO₄, a preenish-fellow pp.

Methyl ether Mc. An oil preparel by the action of cone. H₂SO₄ on methyl plenyl-propiolate (W. H. Perkin, jun., a. Calman, C. J. 49, 154). Fe₂Cl, gives a violet colour in alcoholate (W. H. Culling and the colour in alcoholate (W. H. holio solution. Sodium ethylato gives a white amorphous sait, C.H., CO.CHNa.Co.Me. This salt is v. sol. water and hot alcohol.

Ethyl ether. A'Et. (265°-270°) at 760 mm.; (230°-235°) at 200 mm.

Formation.-I. From phenyl-propiolic ether (100 g.) and H.SO. (3000 g.) at 0°. After three H.SO. gives ucctophenone, benzoic acid, and CO. VOL L

hours the product is poured upon powdered los, and the new body extracted with ether,—2. By heating diazo-acetic other with benzoic aide hyde (Buehnsr a. Curtius, B. 18, 2371),-3. By healing EtONa (140 g.) with benzoic ether (300 g.) at 100°, mixing the product with acetia ether (350 g.) and heating for In hours at 100° (Claisen a. Lowman, B. 20, 651). 4. By the action of cold cone. 11,50, upon (e)-bromocinnamic ether (Michsel a. Browne, B. 19, 1892).

Properties. Colourless oil. Partly decomposed when distilled. Sl. so', water, sol, alcohol

and ether.

liteactions .-- I. The alcoholio solution gives with Fe.Cl. a violet colour .- 2. Boiled with water, or dilute H.SO,, it gives acctophenone, ulcohol, and CO_x -3. Boiling for 8 minutes produces deliver o-benzoyl-acetic acid (q. v.). Boiling for 30 minutes forms two isomerides (Callada). Oue of these (n = 3?) crystallises in plates, [275°], no sol. hot alcohol, v. sl. sol. bonzone; sol. alcoholic NaOH lott ppd. by CO, second (n 4?) is an acid, not being ppd. by CO, from its solution in alcoholio NaOH; cone. H.SO, forms a yellow solution, turning violet when warmed (Perkin, jun., C. J. 47, 262) .- 4. NaNO, and H.SO, added to the sodium derive tive produce an oxim of benzoyl-glyoxylio ether, Ph.Co.C(NOII).Co.Et [121], whence alkalis

produce a substance C. H.O. [125°].

Metallic derivatives. BzCHNaCO.Et. Got by adding NaOEt to alcoholic solution of the other. Silky needles, turns brown in sir; insol. ether. — $(C_{11}II_{11}O_3)_a$ Ba. — $O_{11}II_{11}AgO_4$. (CoII oo), Cu: pale green; solublo in aqueous NaOII. On holling Cu,O is ppd.

Nitrile v. Benzoyl-Acetonitelle. Benzylidene-benzoyl-acetic ether

C.H. CH:CBz.CO,Et. [99°]. From benzoylacetic ether and benzoic aldehyde, cither by passing HCl at 0° into the mixture, or by heating in a sealed tube (l'erkin, jun., C. J. 47, 210). Monoclinio prisms: a:b:c = 1.2730:1:.7460; $\beta = 86^{\circ}$ 36'. Sol. hot methyl alcohol. Cono. H.SO, forms a yellow solution, which becomes colourless on heating.

Benzylideno-di-benzoyl-di-acetic acid

(CO.H.CHBz),CHPh. [130°]. The others of this acid are formed by dropping diazo-nectio ethers (2 mols.) into benzoic aldehyde (8 mols.) at 170 (Buchner a. Curtius, B. 18, 2374). They dissolve in cone. H₂SO₄ forming rose coloured solutions, which turn brown on warming. The acid and its ethers give off a smell of hyacinths when burnt.

Methyl ether [113°]; prisins.
Ethyl ether [103°]; tubles; NaOEt led to its ethereal solution gives added (CO Et.CNaBz) CHPo.

Di-benzeyl-acstic acid CIIBz, CO.H. [1090]. Formed by the action of BzCl on sodinm benzoyl-acetic ether, and saponification of the product with KOH (Bueyer a. Perkin, jun., B. 16, 2133; C. J. 47, 240). Slender felted needles, sl. sol. cold alcohol and water, v. sol. ether; sol. aqueous alkalis. FezCl, gives a red colouration. Cone. H.SO, gives no colour on warming.

Reactions. 1. Boiling water splits it up

into di phenyl methylens di ketone. (Calla CO) Cli, and CO. 2. Boiling dilute

Balt .- Aga: Ethylether EtA'. Oil; not solid at -10°. Methyl-benzoyl-acetic acid is (a)-BENZOYL-

PROPIONIC ACID (q. v.).

Ethyl - benzeyl - acetic acid BzCHEt.CO.H. [1110_1150]. From NaOEt and EtI on alcoholie BzCH, CO, Et, and saponifying the oily product by allowing it to stand for some days with alcoholic KOH (Bneyer a. Perkin, jun., B. 16, 2130; C. J. 45, 180; 47, 240). Small needles, melts about 115° with slight decomposition. Easily soluble in alcohol, ether, and benzene. Boiled with dilute alcoholic KOII, it gives plicaylpropyl-ketone; benzoic and butyrio acids are also formed, especially if the potash be strong.

Ethyl ether EtA'. (232°) at 225 mm. Propyl-boazoyl-acetic ether BzCHPr.CO.Et.

(239) at 225 mm. Prepared like the preceding. Alkalis from phenyl-butyl-kotone. PCl, forms \$-chloro-a-propyl-cinnamic ether.

1sopropyl-beazoyl-acstic ether

BzCHPr.CO.Et. (237°) at 225 mm. From bonzoyl-acetic other, Na, and Prl.

Iso-butyl-beuzeyl-aestic other

Bz.Cl1(Cl1, Cl1Me,).CO, Et. (247°) at 225 mm. Prepared like the preceding (Perkin a. Calman,

C. J. 49, 165).

Di-ethyl-honzovi-acstic acid BzCEt.CO.II. [128°-130°]. From Bx.Cl1Et.CO.Et by NaOEt and Etl. The diethyl-benzayl-acetic other is saponified by standing for weeks with dilute alcoholic KOH (Baeyer a. Perkin, jun., B. 16, 2131; C. J. 45, 183). Heated alone or with dilute 11,80, it gives off CO. Boiling dilute alcoholie KOH forms benzoic acid, diethylacetic soid, and di-othyl-acetophonone (v. amyl-phenylketone).

Trl-beazoyl-acetic ather. CBz, CO,Et. From ethyl di-benzoyl-acetate, NaOEt, and BzCl (Perkin, jun., C. J. 47, 240). Thick yellow oil; sol. alcoholic KOH but reppd, by water. Boiling

dilute 11.80, forms acctophenone.

BENZOYL ACETIC-ALDEHYDE C,H,CO.CH,CHO. Prepared by dissolving sodium (1 atom) in 20 or 30 times its weight of absolute alcohol, cooling to 02, and adding nectophenone (1 mel.) and formic ether (1 mel.). On long standing the sodium compound separates as a granular pp.; this is dissolved in water and the aldehyde ppd. by nectic neid. Colourless un-stable oil. Cupric acetate gives a pp. of bright green needles, which soon clininge to dark green prisms. It reacts with amines very readily.

Anilide CaH, CO.CH. CH: NCaH,: [141°];

yellow prisms or plates; sol. hot alcohol. p.Toluide C, II, CO.CH, CH:NC, II, : [160°-

163°; small yellow crystals.

(B)-Naphthylamide C.H., CO.Cll. CH: NC, 11,: [182°]; small bronzy crystals; sl. sol. almost all solvents (Claisen a. Fisoher, B. 20, 2191).

BENZOYL - ACETIMIDO - ETHYLIC ETHER

C.H. CO.CII. C(NII).OEt. [89°-5 corr.]. Bsnzoyl-acetonitrile C.H. CO.CH. CN treated with slcoholic hydrochloric acid gives rise to C.H. CO.CIL, C(NHHCI)OEt and this loses HCl when treated with ammonia giving the imido-ether (Haller, Bl. [2] 48, 24; C. R. 10t, 1418).

Properties .- Prisms or tables. V. sol, ether. Reactions. - KNO, and II, SO, give the nitrosoderivative C.H. CO.CH. C(N.NO).OEt [117], has no action upon it.

which gives Liebermann's reaction. The hydrochloride of the base dissolved in aqueous alcohol deposite MH,Cl and benzoyl-acetic ether is left in solution.

[140° corr.]. Hydrochloride B'.HCl. Entangled aecdles. Insol. aq aad ether. Strongly irritates the mucous mombranes.

BENZOYL-ACETO-ACETIC ETHER v. p. 21. BENZOYL - ACETO - CARBOXYLIC ACID .

ACKTOPHYNONE CARNOXYLIC ACID.

BENZOYL-ACETONE C₁₀H₁₀O₂ i.e.

CO.CH., CO.CH₃. Phenyl methyl met C.H.CO.CH.CO.CH. thylene di-ketone. Acetyl-acetophenone. Acetylbenzoyl-methane. [61°], (261°).

Formation .- By the action of dry NaOEt upon a mixture of accteae and benzoic ether

(Claisen, B. 20, 655).

Preparation. - 1. Beazoyl-accto-acctic ether, formed by the action of benzoyl chloride upon sodio-aceto-acetio ether, is boiled with water for a few hours; the yield is 25 p.c.-2. Prepared by adding acetophenone (1 mol.) to a cooled mixture of acetic ether (about 2 mols.) and alcohol-free sodium ethylate; yield: 80-90 p.c. of the acetophenone (Beyer a. Claisen, B. 20,

Copper compound (C10H100,)2Cu: formed as a pale green pp. by adding cupric acetate to the alcoholio solution. It is m. sol. alcohol and bouzeno, from which it crystallises in bright

green needles.

Amide Calla-CO.CH_C(NH).CH3: [143°]; clear glistening trimetrio crystals, a:b:c = '9927:1: '8820.

Anilide C. H. CO.CH, C(NPh).CH,: [110°]; plates. By warming with II, SO, (10 pts.) it is converted into (Py. 1:3)-phenyl-methyl-quino-line (Beyer, B. 20, 1770).

Properties. -- Crystals; distils undecomposed; volatile with steam. Sol. hot water, alcohol, and other. Dissolves in alkalis with a yellow colonr. Sl. sol. strong acids. Fo₂Cl.

gives a dark-red colouration.

Reactions. - By warming with alkalis or by long boiling with acids it yields acetophenone. It has slightly acid characters, the H of the central CII, group heing replaceable as in aceto-acetic ether, since it lies between two CO groups. By heating with strong aqueous NII, at 120° it is converted into the imide

C,H,C(N11).C11,.CO.CH, or Cally CO.CH. C(NII).CII. It condenses with (I mol.) of phenyl hydrazine with elimination of 21LO, forming methyl-di-phenyl-pyrazol (Fischer a. Bulow, B. 18, 2131).

Salts. .. C,oll,O,Na: small yellowish plates. -C1011,OAR: white pp.; v. sl. sol. water

(Fischer a. Kuzel, B. 16, 2239).

Oxim C10H,102Ni.c.C6H,C(NOH)CH2CO.CH or C.H., CO.Ch , C(NOH).C11, [66°]. Formed by hearing benzoyl-acetone with hydroxylamine hydrocliforide in alcoholic solution (Ccresole, B. 17, 812). White glistening scales. Volatile with steam. V. sol. sectone, benzene, and CS. insol, water.

Di-beazoyl-aceteae (C.H.,CO),:CH.CO.CH, 12°]. Formed by the action of beazoyl [1020]. chloride upon sodio benzoyl acetone (Fischer a. Bülow, B. 18, 2133). Small needles. alcohol and ether, v. sl. sol. water. Sodiam

BEHEOYL-ACETONIMIDE CHEHINO &.e.

C.H. O(NH).CH, CO.CH, or C.H. CO.CH, O(NH).CH, [143°]. Obtained by heating benzoyl acctone with strong aqueous NH_a at 120° (Fischer a. Bülow, B. 18, 2134). Distils undecomposed. Small plates, or largo quadratic crystals. V. sol. dilute acids, by quadratic crystals. heating with which it is converted back into benzoyl-acetone and N11.

BENZOYL-ACETONITRILE C.H,NO

C.H. CO.CH. CN.

Cyano-acetophenone. [81° cor.], Formed by the action of boiling water on benzoyl-cyanacetic other C.H., CO.CH(CN).CO.Et (Haller, Bl. 12) 48. 23). White needles, sol. boiling water, alcohol, ether, and alkalis. Boiling cone. KOII C.H., CO.CH., CN + 2KHO + H.O thus: $= NH_1 + C_2H_3 \cdot CO_2K + CH_2 \cdot CO_2K$. In alcoholio solution gaseons HCl gives a body C_HH₁₁O NCl (probably C_H, CO.CH, C(NHHCl)OFt 110corr.]; whence aminonia in the cold gives the imidoether C.H. CO.CH. C(NH).OEt). If the action of the alcoholic HCl is prolonged, the products me the same as with KOII.

Silver salt Call, CO.CHAg.CN. White pp. Insol. aq and alcohol; sel. ammonia.

BENZOYLACETOPHENONE v. Districtive

METHYLENE DI-KETONE.

BENZOYL . ACETYL ETHANE v. ACETO -PHENONE-ACETONE, p. 36. Di-benzoyl-di-acetyl-ethane

C.H.CO.CH.CO.CH, C20H11O i.c. Di phenyl-C.H. CO.CH.CO.CH,

di-methyl-acetylene-tetra-ketone [175°]. Formed by the action of an otherest solution of iodino upon 2 mols, oI sodio-benzoyl acctone (Fischer a. Bulow, B. 18, 2133). White needles. Sol. hot alcohol, sl. sol. ether, insol. water and dilute alkalis. Decomposed by boiling with alkalis.

BENZOYL-ACRYLIC ACID C.H. CO.CH:CH.CO.II. White plates, [64°], from water; after fusion its melting-point is altered to [97°]. Long needles, [99°] Irom toluene. Sl. sol. cold water and ligroin, v. sol. other solvents. Prepared by the action of Al,Cl, on a mixture of benzene and maleie anhydride. By alkalis it is decomposed into acctophenone and glyoxylic acid. On heating water.

by itself or with Ac O, it gives a red condensation product (Peclmann, B. 15, 885). Bromine addition product [135°]. Colour-

less crystals.

BENZOYL-ALLOPHANIC ACID v. p. 127. BENZOYL-ALLYL-ACETIC ACID v. ALLYL-BENZOYL-ACETIC ACID, p. 135. BENZOYL-AMIDO- v. AMIDO-

BENZOYL-AMIDO-ACETIC ACID v. Hip-

PURIC ACID.

BENZOYL . AMMELINE D. H. N.O. C,H,BzN,O. From sodium cyanamide and BzCl (Gerlich, J. pr. [2] 13, 272). Brown resin, insol. water and other, sol. alcohol and aqueous alkalis. Resolved by distilling in a current of hydrogen into benzonitrilo, carbonio oxide, and eyanamide.

BENZOYL-ANILIDE v. ANILINE.

BENZOYL - ANILINE v. AMIDO - BENZOPHE-

BENZOYL-ANISIDINE v. Benzoyl-methyl-AMIDO-PHENOL.

BENSOYL-AZOTIDE v. p. 475. BENZOYL-BENZENZ v. BENZOPHERONS, Di-benzoyl-benzene v. PHTHALOPHENONE.

BENZOYL-BENZIDINE v. Di-Anido - Di-

BENZOYL-BENZOIC-ACETIC ANHYDRIDE C.H., CO.C.H., CO.O.CO.CH., [112°]. Prepared by henting o-henzoyl-henzoio neid with acotio unhydride to 100° (Freiher a. Peelmann, B. 11, 1865). Largo crystals. Insol. alkulis. At 200 it decomposes into acctic and benzoylbenzoic anhydrides.

O BENZOYL-BENZOIC ACID C, Il, O. i.s. Call, CO.Call, CO.H [1:2]. Bensophenone carboxylic acid. Mol. w. 226. [87] (Z.); [949] (Hemilian, B. 11, 838).

Formation .- 1. By exidation of o benzyltolneno (Zineke a Plaskuda, B. 6, 907), phonylo-tolyl-ketone (Behr n. Van Dorp, B. 7, 17), or di-benzyl-benzene (Zincke, B. 9, 32) with chromic mixture.

Preparation, -150 grms, of Al₂Cl₄ are slowly added during 3 hours to a solution of 100 grms. of phthalic anhydride in 1000 grms, of henzone (pure), the benzene is then poured off and can be used at once for a fresh operation, whilst the solid residue is washed with dilute HCl and with water, dissolved in Na₂CO₂ and the acid precipitated from the solution by HCl, and finally recrystallised from xylene (3 pts.); the yield is 60 p.e. of the phthalio milydride used (Friedel n. Crafts, C. R. 86, 1368; 92, 833; Freiherr a. Pechmann, B. 13, 1612).

Properties. -- Triclinie needles (containing

aq). When dry it melts at 1281

Reactions. -1. P₂O₃ at 190° forms anthra-quinone, -2. Hot funing sulphuric acid forms anthraquinono sulphonic acid (Liebermann, B. 7, 80a). - 3. Sodium amalyam first reduces it to C₂H₂Cl1(Ol1).C₂H₄CO₂H and then to C₃H₂Cl1₂CO₂H₃CO₃H. A. Resorcin, pyrogatlol &c., on heating, form phthaleins. 5. With phonyl-hydrazine it gives a condensation-pro-C -Ph

N₂Ph [182°] (Roser, B. 18, 805). čo

This forms small needles, sl. sol. alcohol, insol.

Salts. - CaA', - BaA', - ZnA', 2aq. -

CuA, aq.

Methal ether MoA'. [52]; prisms.

Anhydride (Bz.C.II, CO),O. [120] (Pechmann, B. 14, 1866).

m-Benzoyl-benzolc acid I'm CO.C. Il. CO.H.

[1:3]. [1616]. Formation.-I. From planyl m tolyl methane (10 g.), K₂Cr₂O₂ (60 g.), H₂SO₄ (90 g.) and water (270 g.) by boile 3 for 3 days (Botering; Senff, A, 220, 237). It is purified by reduction to Ph.CH(OH).C.H.,CO,Na by sodium amalgam, crystallising this salt from water and oxidising again with H SO, and K Cr2O,-2. Aquaatitative yield is obtained by treating phenyl-m-tolyl-methane at 130° with bromine-vapour sufficient to form Ph.CH, C.H, CH, Br and treating the product with chromic mixture for 12 hours,-8. From Bz.() (I mol.) and BzCl (2 mols.) in presence of ZuCl. (Docbner, A. 210, 277; B. 14, 648) .- 4. Formed as a by-product in the pre-

Properties .- Long silky needles (from water or glacial acetic acid), or small plates (from alcohol). Sl. sol. cold water, v. sol. benzene of tolueno, v. e. sol. nlcohol or other. May be sublimed as pintes. Dissolves in conc. H.SO, giving no colour. l'otash fusion gives benzoic acid. Reduced by sodium-analgam to exo-oxy-benzylbenzoic acid (q r.).

Salts. BaA' 3aq: white crystalline powder, BaA', 4aq: small plates.—CaA', 2aq: white crystalline powder. - AgA': white leaflets.

Methyl ether Mi A' [62]

p-Benzoyl-berzoic acid C.H. CO.C.H. CO.H [1:4]. [191]. Formed by oxidation of phenylp-tolyl-methane, phenyl-p-tolyl-ketone, p-phenylbenzophenone or di-benzyl benzene (Zincke, A. 161, 98; B. 6, 907; 9, 32; Goldschmiedt, M. 2, 438). Monochnic plates (from water). V. sl. sol. cold water; sl. sol. hot water (difference from the o acid), v. sol, alcohol and ether, sl. sol, benzene. Sublines in plates.

Salts. CaA', 2aq: needles.-BaA', 2aq.

AgA': v. el. sol, water,

Methyl ether McA', [107°]. Satiny plates. Ethyl other EtA'. [52°]. Monochine.

DI BENZOYL BENZOIC ACID Calling, i.e. (C, H, GO) C, H, CO, H. Two acids of this composition are formed, together with an acid $C_{12}\Pi_{10}O_{21}$ by exidising the hydrocarbon $C_{13}\Pi_{24}$ obtained as a by-product in the preparation of benzyl-tolnene (Weber n. Zineke, B. 7, 1153).

(a) Acid. (824). Resinous, and forms resinous salts. Potash fusion forms benzuie neid

and a small quantity of an acid C₁ H₁₀O₂.

(B) Acid. (212°). Needles, insol. water, v. sol. ulcohol and other. Its salts are sl. sol. water. Ethyl ether EtA', (1079).

DI BENZOYL. BENZYLIDENE. DT. ACFTIC ACID v. Benzylideno di Benzontina acrite acris.

BENZOYL BENZOTRICHLORIDE P. PHENAL TRI-CHLORO-TOLVE KETONE.

BENZOYL-RENZYL ANILINE U. BENZYL.

BENZOYL-BENZYL CHLORIDE P. PRENYL CHLORO-TOLYL BE DINE.

BENZOY) BENZYLIDENE CHLORIDE r.

PHENTE DESCRIOR STOTAL RETORY. BENZOYL-DROMANILINEY, BROMO-ANILINE,

BENZOYL BROMIDE C.II, CO.Br. 219%, V.). [a little below 0°]. S.G. 10 1.67. Colourless

fluid, which fumes in the air. Preparation - Benzoic acid (500 pts.) is warmed with phosphorus tribromide (740 pts.)

and the product is separated from the phosphorous neid by distillation in racuo; the yield is 400 pts. (Claisen, B. 11, 2473). The so-called benzeyl brounde of Liebig and Wöhler (A. 3, 260) and of Patern's was probably benzylidene bromide benzoate, C.H.CHEr(OBa), a compound of benzoic aldehyde with benzoyl bromide.

BENZOYL . BROMO - NITRANILIDE RHOMO-NITRO-ANILINE

BENZOYL - BROMO - PHENOL v. BROMO.

DI-BENZOYL-ISOBUTYRIC ACID

paration of isophthalophenone by the action of ing di. 5 bensoyi di methyl malenie seid Al.Cl. on a mixture of benzene and iso-phthalyl (Bs.CH.), C(CO.H). Silky needles. V. soi. sico chloride (Ador, B. 18, 821). ligroin. - A'Na: glistening needles (Kues a. Paal. B. 19, 3147)

BENZOYL-CARBAMIC ACID. C.II,NO.

Ethyl ether CaH3.CO.NH.CO.Et. [110°] Formed by boiling benzoyl-thiocarbamic ether in alcoholic solution with PbO (Lösener, J. pr. [2] 10, 254). Needles (from dilute alcohol), sl. sol, water. Decomposed by aqueous KOH into KOBz, alcohol, NII, and K.CO. Alcoholio KOH gives a pp. of C.H. CO.NK.CO.E1, v. e. sol. water.

BENZOYL-CARBINOL Ph.CO.CH,OH [86°]. Ph.C(OH), CH.OH [74]. Exo oxy - phenylmethyl ketone. Oxy-acctophenone. Aceto-phe-

none alcohol. Phenicyl alcohol.

Formation. - 1. By the action of alkalis on the acctate or chloride, C.H., CO.CH., Cl.-2. From phenyl-glycol (2 g.) and HNO, (6 o.o. of S.G. 136). The mixture is warmed and, as soon as reaction sets in, it is cooled. The product is diluted, neutralised with Na CO, and the crystalline carbinol filtered off. Ether extracts a further quantity from the filtrate (Hunaeus n. Zineke, Ji. 10, 1187).

Properties. Prisms or pintes (from benzol-Large erystals (from alcohol or other). V. sol. ether, alcohol, or CHCl, Largo plates containing II,O (from water or dilute aloohol). In drying these, decomposition readily occurs, benzoic aldehyde being formed. The carlinol combines with NaIISO, Readily reduces ammoniaeal silver nitrate or Fehling's solution, the chief product of the oxidation being numbelic acid C₈H₂CH(OH),CO₂H₄ (Breuer a. Zincke, B. 13, 635).

lie utions.-1. Heated alone it gives off benzoic aldehyde and n pungent body.— 2. Heated with aqueous NaOll or baryta it becomes yellow and forms benzoic aldehyde.-3. Water at 140° acts similarly, -- 4, With IICN it yields the nitrile of atroglyceric acid CH_(OH).CPh(OH).CO2H (Plocht a. Blumlein, B. 16, 1290).

Acetate Ph.CO.CII,OAe. From chloro-acetophenone and AgOAc (Graebe, B. 4, 34); or from the earbinol and Ac.O (Zincke). Trimetric tables (from benzoline). V. sel. alcohol, ether, or chloroform. M. sol. benzoline.

Benzoate Ph.CO.CH,OBz. [1170] (Zineke). From chloro aceto-phenons and AgOBz or from benzovl - carbinol and Bz O. Small tables (from dilute alcohol). V. sol. ether, benzene. or CHCL

BENZOYL-CHLORANILIDE v. CILORO-ANI-

BENZOYL CHLORIDE C.II, CO.CI. (Liebeh, A. 178, 43). (198°). (194°) (Brühl, A. 235, 11). S.G. 1: I-21; 29 I-2122 (B.). V.D. 4.99 (calc. 4.90). $\mu_D = 1.5537$. S.V. 126.3 (Ramsay).

Formation.-1. From chlorine and benzois aldehyde (Liebig a. Wöhler, A. 3, 262), -2. From benzoic seid and PCl, (Cahours, A. Ch. [8] 28, 334 .. - 3. From benzoates and POCI, (Gerhardt, O.H., CO.CH.), CH.CO.H. Di-phenacul-acetic the action of Clupon mandelic acid or on benseid. [183°]. Formed by loss of CO, by heat zoic ethers (Malaguti, A. Ch. [2] 70, 374).-

B. 19, 8232).—10. In considerable quantity by ployed (Lenckart, B. 18, 2341). White plates heating benzaldehyde with glycocoll (Curtius a. | (from hot alcohol); v. al. sol. water. When Lederer, B. 19, 2462).—11. Together with di- and | heated for a long time with MeI or EII at 150°. tri- benzyl-amino, as a by product, in the proparation of di-benzyl-hydroxylamine from hydroxylamine hydrochleride, benzyl chloride and NaOH (Walder, B. 19, 3293).

Properties .- Liquid, unscible with water, alcohol, and ether. Separated from water by KOH. Strongly alkaline, absorbs CO, forming a crystalline carbonate, and fumes with IICl. With cyanogen it forms a compound (C, II, N), (CN), [140°] which crystallises from alcohol, and forms a hydrochloride (C.H.N), (CN), 2HCl (Strakosch, ; B. 5, 693).

Salts.-B'HCl: large leaflets or flat tables. -B'HBr.-B', H.PtCl,: orange tables or ye low plates, st. sol. water. -B', H.SO,.

Acctyl derivative C. H. Cll. NHAc [61°]; (300°); crystalline solid, sol. water (Amsel a. . Hofmann, B. 19, 1285; Strakosch, B. 5, 697; Budolph, B. 12, 1297).

Di-benzylamine CarloN i.c. NH(CH, Call). S.G. 14 1 033.

Formation .- 1. By the action of NII, on C.H.CH.Cl or by reduction of benzonitrile (v. supra). 2. By the action of bromine-water on tribenzylamine (Limpricht, A. 144, 313). - 3. By boiling henzoic aldehyde with ammonium formate (Leuckurt a. Bach, B. 19, 2128). 4. Occurs together with mone- and tri-benzylamine as a by-product in the preparation of di-benzyl-hydroxylamine from hydroxylamine hydrochloride, benzyl chloride, and NaOH. 5. Formed by the action of PCl, upon di-benzylhydroxylamine and treatment with water, the

hydroxynamic and beams; reaction probably being: (C,H,),N,O,PCl,+11Cl and (C,H,),N,O,PCl,+31I,O = (C,H,),N,N,PCl,+21IC(Wulder,B, 19, 3287).

Properties, Liquid, insol. water, v. sol. alcohol and ether. Does not absorb CO, from the air. On distillation it decomposes into s-di-phenyl ethane, s-di-phenyl ethylene, lophine, and various bases (Brunner, A. 151, 133).

Salts.—B'HNO₄: [186°], very sparingly soluble thin glistening needles.—B'HCl [256°]. -B'HBr [276°]. -B'HI [224°]. -B',H,PtCl, golden yellow needles.

Nitrosamine (C.H.Cll.), N.NO: [61°]; white crystals: v. sol. alcohol and other, insol. water (W., cf. Rohde, A. 151, 366).

Picryl derivative (C, H, CH,) N.OC, H, (NO),* (171°); orange plates.

Formyl derivative (C. II, CII,) N.CIIO: [52°]; (above 360°) (Lenckurt a. Buch, B. 19, 2128).

Disulphonic acia C₁₄II₁₃N(SO₂II)₂ (Linpricht, A. III, 317).—BaA".

Trl-benzylamine C₂₁H₂₁N i.f. (C₄II₂,CJ₁)₄N.

Formation,-1. From benzyl chloride and NH, (v. supra).—2. By heating di-benzylamine with benzyl chloride at 100° (Walder, B. 19, 3287) .- 3. Tegether with mono- and di- benzylamins as a by-product in the preparation of di-benzyl-hydroxylamine from hydroxy:amine hydrochloride, benzyl coloride, and NaOH (W.). 4. By heating benzaldehyde with rather more than an equal weight of ammonium formate; the yield is 40 p.o. of the benzaldehyde em-

benzyl iodide and tetra-methyl- (or ethyl-) ammonium iodido are formed (Marquardt, B. 19, 1027). Fuming sulphuric acid forms

C_HII_{I,C}(SO₂H)₃N (Limpricht, A. 144, 311).

Salts.—B'IICl: [228°]; tbick prisms or iridescent plates, v. sol. hot alcohol, insol. water. -B'₄H.Cl₄! transc vellow needles.—B'IINO₃: [125°]; insol. water. B'IIBr: [208°]. -B'HBr, -B'HI: [178] -B'HAl(SO,), 12aq: [110]; sol. water.

Methylo-iodide B'MeI: [1819]; needles or plates; sol, hot alcohol, sl. sol, cold water. Methylo-hydrate B Me(OH): crystalline solid; alkaline reaction; v. sol. water. On heating it evolves McOH forming tri-benzylamine.

Methylo-chloride-platinum-salt (B'MeCl), PtCl4: [197°]; orange pp.; insol. cold water and alcohol.

Ethylo-iodide BEH: [190°]; colomless rhombic crystals; sol, alcohol and hot water. Isopropyla iodide B'FrI: (170°1; needles; sl. sol. hot water.

BENZYLAMINE. W. DI-SULPHONIC ACID C_bH_wCH(SO₅H),NH(SO₅H). The di sodium salt A"Na23aq is formed by shaking henzuldoxim with a 30 p.c. sodium bisulphite solution. It crystallises in small white needles, v. e. sol. water, insol. cold alcohol. By warming with dilute acids it is split up into benzaldehyde, sodium sulphute, and ammonium sulphite: C_oH₃.CH(SO₃Na).NH(SO₃Na) + 2H₂O

 $= C_nH_s.CHO + Na_sSO_s + (NH)_sHSO_s.$ Alkalis decompose it in the cold, and water on boiling (Pechmann, B. 20, 2539).

BENZYL-ANILINE Calls, NH(C,11), [33° uncor.]. (above 360°). Obtained by reducing thiobenzeyl-miline (Bernthsen a. Trompeter, B. 11, 1760). Fermed also by boiling diazobeuzenebenzyl anilide (50 g.) with 11Cl (200 c.c.); the yield is 20 g. (Friswell a. Green, B. 19, 2036). Yellowish crysta.

Salts.—B'HCl [203° nucor.]; white plates; decomposed by water.—B', H, Cl, PtCl, [168° uncor.]; slender yellow needles; tolerably easily soluble in water. - B'₂C₁H₂O₄, --B'CdCl₂.

Benzoyl derivative [104] (Floischer, A. 138, 229).

Di-benzyl-aniline Call, N(CII, Call,)2. [67°]. (above 300°). Prepared by heating a mixture of ancline (51 pts.), benzyl chloride (150 pts.) and NaOH (30 pts.) on the water-bath for three or four weeks. After cycling the solidified cake is pressed, distilled with steam to remove excess of benzyl chloride, washed with hot water, and crystallised from alcohol. Colourless needles. V. sol. other, benzene, hot alcohol and hot acctic acid, sl. sol. cold alcohol and cold acetic acid, nearly inso!, water. Weak base.

Salts. - B'HClaq: glistening prisms. B'ALCI PtCl.: thin orange yellow scales. Picrate B'C.H.(NO.), OH: [132], long yellow needles (Matzudaira, B. 20, 1611).

TRI-BENZYL-ARSINE v. p. 322. BENZYL-BARBITURIC ACID v. BARBITURIO

BENZYL-BENZENE v. Di-phenyl-methans, Di-benzyl-benzene CzoH, i.e. (C.H.CH.),CH, [86°]. Formed, together with the o-isomeride and di-phenyl methane, by the action of sine on a mixture of benzyl chloride and benzene, or by the action of H,SO, on a mixture of benzene and methylal, CH₂(OMc), (Zinoke, B. 6, 119, 221; 9, 31). Transparent lamine, sl. sol, ether, v. sol. benzene and hot alcohol. CrOs forms (a)-dibenzoyl-benzene and p-benzoyl-benzoie acid.

o. Di. benzyl. benzene $(C_uH_u.CH_z)_2C_uH_4$. [78°]. Sifky needles (from alcohol); v. sol. ether and alcohol. CrO, lornis o-di-benzeyl-benzene and o benzoyl benzoic steid.

BENZYL BENZOATE C, II, 20, i.e.

C, II, CH, C.C.C, II, (324° i.V.). [21°]. S.G.
(fluid, at 19°) 1·12°24. From benzyl alcohol and BzCl (Kraut, A. 152, 130). Formed also by several days' heating of benzaldehyde at 100° with a small quantity of sodium benzylate;

probably the compound C.H., C(OC,H.), ONa is first formed and then decomposes into benzyl benzoate and sodiom benzylite, which latter again reacts upon a further quantity of benzaldehyde, producing more of the intermediate product, and so on. Large colourless crystals

(Claisen, B. 20, 616).

O BENZYL BENZOIC ACID C.H.O. i.e. C.H. CH, C.H. CO.H. Mol. w. 212. [114]. From o benzoyl-benzoic acid and sodium amalgam (Rotering, J. 1875, 598; B.9, 633). Slender needles; may be sublined; al. sol. cold water, v. sol. alcohol and other. CaA', 2 aq. -CaA', 3', aq. BaA',5) aq. AgA', McA', in-Benzyl-benzelc acid

Ph CH, C, H, CO H. [1089].

Formation. 1. From excoxy-benzyl-ben zoie acid, Ph.CH(OH).C.H.,CO.H. und cone. HI at 170 : -2. From exc-bronco in tolnic acid, CH_Br.C.H. CO.H. beazene and Al_Cl. (Scuff, .1. 220, 247). Yield 50 p.c. of theoretical from toluic acid. 3, A small quantity from benzoic other, benzyl eldoride, and ZaCl, by boiling.

Properties. Short slender needles (from hot water), small plates (from hot dilute alcohol); sl. sal. cold water, to, sol, hot water, v. n. sol, alcohol, ether or chloroform. Conc. H.SO, forms a colourless solution. K.Cr.O. and 11.SO, give m - benzoyl - benzoie acid. — CaA'2 nq. — BuA', Inq. AgA'.
p Bounyl-benneic acld

Ph.CH, C.H, CO,H [1:4]. [155°].

Formation. I. By oxidising p benzyl-tolnene with dilate 11,80, (Zincke, A. 161, 106),--2. By reducing exo-oxy-p-beazyl benzoic neid with HI. 8. From p-benzoyl henzoic acid and sodiumamalgam or Ill and & (Graebe, B. 8, 1054).

Properties. - Minute needles (from water); may be sublimed; sl. sol. cold water, v. sol. alcohol and other. Chromic mixture oxidises it to p-henzoyl-benzoic acid - CaA'.HA' .--

BuA', 2nd. AgA'

BENZYL BROMIDE C. 11, CII, Br. (199°). **8**.G. 🖓 1 4380.

Fermation. - I. From benzyl alcohol and HBr (Kekulé, A. 137, 190) .- 2. From Br and boiling toluene (Beilstein, A. 143, 369; Jackson a. Field, Av., 2, 11).-3. From benzyl chloride and Ashr, (Brix, A. 225, 163).

Preparation.-By the action of bromine (1 mol.) upou cold toluene (1 mol.) in direct sunshine; the yield Is quantitative (Schramm, B. 18, 608).

couple acts vigorously upon it, producing two isomeric benzylenes. In presence of ether, the zino-copper couple produces dibenzyl, ZnBr. and C,H,ZnBr, whence water produces toluene: 2C,H,ZnBr+2H,O=2C,H,+ZnBr₂+Zn(OH), In presence of alcohol, the couple produces toluene and EtOZnBr. In presence of water, the couple produces dibenzyl and a little toluene (Gladstone a. Tribe, C. J. 47, 448) BENZYL BUTYRATE C,H,O, i.e.

Properties -- Pungent liquid. The sine-copper

C.H. CH. O.CO.Pr. (240°). S.G. 16 1.016 (Conrad a. Hodgkinson, A. 193, 320)

Benzyl isobutyrate Me_CH CO_CH_Ph. (228° i. V.). S.G. #3 1 016. Prepared by boiling au alcoholie solution of benzyl chloride and potassic isobutyrate for five days with inverted condenser. The product is mixed with water and the oil distilled.

Properties .- Oil, with pleasant odour.

Reactions .- When benzyl isobutyrate (90 g.) is heated with sodimu, (8g.) a violent action occurs, the products being hydrogen, sodic isobutyrate, benzylie benzyl-isobutyrate (q.v.), sodio benzeate, toluene, and an oil (C, Il 180), (340°-350°). The principal reaction is:

4Me,CH.CO,,CH,Ph + Na, = 2Me,C(CH,Ph).CO,C,H, + 2Me,CH.CO,Na + H, (W. R. Hodgkinson, C. J. 33, 496), a-BENZYL-ISOBUTYRIC ACID Call 140, i.e.

Me_C(CH_Ph)CO_H. Benzyl ether (C,H,)A'. (280°-285°). S.G.

1 0285. Prepared by the action of sodium on benzyl-isobutyrate (q. v.).

Reactions. - 1. Hented with sodium a violent action occurs, toluene, sodie beuzoate, sodie benzyl isobutyrate and an oil, CulluO (350' 355°) being formed .- 2. It is attacked by alkalis with great difficulty, the saponification gives isoflutyric not benzyl-isobutyric acid (W. B. 11odgkinson, C. J. 33, 503; A. 201, 171). BENZYL CARBAMATE NIL.CO.O.C.II,

[86]. From benzyl alcohol and area nitrate at 140° (Campisi a. Amato, B. 4, 412) or solid eyanogen chloride (Caunizzaro, B. 3, 518). Large plates (from water); sl. sol. hot water, v. sol. ulcohol. Decomposes above 200° into benzyl

alcohol and cyanuric acid.
BENZYL-CARBAMIC ACID C,H,NH.CO.OH. Benzyl-ammonium salt C,H,N11.CO,NH,C,H,. [99°]. From benzylamine and CO2. Formed also by heating a amido-phonyl-acetic acid at 260°; the yield being nearly the theoretical (Tiemann a. Friedländer, B. 14, 1969). Plates, sol. water and alcohol, insol. ether; volatile with steam. Decomposed by

acids or alkalis into CO, and benzylamine,
BENZYL DECARBOXY GLUTACONIC ACID

v. Discarboxy-diotaconic acid.
DIBENZYL CARBOXYLIC ACID v. Di-PHENYL-FTHANK-CABBOXYLIC ACID

DIBENZYL DI-CARBOXYLIC ACID v. Dr. PHENYL-SOCCINIO ACID and DI-PHENYL-ETHANE DI-OARBOTTLIC ACID.

BENZYL CARBINOL v. PHENYL-ETHYL AL-

BENZYL CHLORIDE C,H,Cl i.e. C,H,CH,CL Mol. w. 126.5. a. Chloro - toluens. (178°) at 754 mm. S.G. 47 0.9453. S.V. 138.45 (Schiff, B. 19, 563; A. 220, 98); 133-18 (Ramsay).

(Cannissare, 4. 88, 129; 96, 346; Deville, A. Ok. [8] 3, 178).—2 By distilling toluene in a surress of chlorine (Lanth a. Grimaux, Bl. 1867, i. 105).

Preparation.-By passing ohlorino (1 mol.) into cold toluene (I mol.) oxposed to direct sunshine; the yield is nearly theoretical (Schramm, B. 18, 608).

Properties .- Oil, sol. alcohol and ether.

Reactions .- I. Boiling alcoholic KOH forms O.H.OEt. - 2. Alcoholie KOAc forms C.H.OAc. -3. Alcoholic KCN forms C.H.CN. -4. Alcoholio NH₂ forms, on heating, mono., di., and tribenzylamine.—5. Hot dilute HNO, (or a nitrate) forms benzoic aldehyde. - 6. Boiling Pb(Oll), forms benzyl alcohol. - 7. KOPh forms phenyl benzyl oxide. -8. Water at 180 gives a product which, on distillation, yields henzyl-tolucue and anthracenc. Before distillation the product is perhap: C.11, CH...C.11, CH Cl (Van Dorp, B. 5, 1070; Zineke, B. 7, 276).—9. Long boiling with water (30 vols.) produces benzyl alcohol. 10. Sodium-amalgam produces a little s-di-phenyl-ethylene.—11. Aromatic hydrocarbons in presence of powdered zinc give off HCl and form condensation products (Zincke, B. 6, 137) .- 12. Chloroformic ether and sodium form di-phenyl ethano exo-carboxylio ether, PhC11, CHPh.CO.Et (Wartz, C. R. 70, 350).

13. Heated with Al.Cl, it gives off HCl, forming tolueno and authraceno (Perkin, jun. a. Hodgkinson, C. J. 37, 726).-14. In carbon disalphide solution yields, when chromyl chloride is added gradually, a brown precipitate of composition PhCH,Cl, CrO,Cl., slowly converted by moist air into benzoic atdeliyde;

3Ph,CHCl,O,Cr(OH)Cl, + 3H .0 =9HCl+Cr₂O₃ CrO₃ + 3PhClIO.

The compound heated to 170° loses HCl. forming a compound PhCHClCrO Cl, which also yields benzoic aldehyde.

3PhCHCl.O.CrO.Cl + 3H₂O

=6HCl + Cr₂O₉₃ + CrO₃ + 3PhCl 10 (Etard, A. Ch. : [5] 22, 235),—15. HI reduces it to tolucue,—16. Zinc dust gives toluene, phenyl-tolyl-methane, and anthracene (Frost, Bl. [2] 46, 249).

BENZYL . CHLORO . MALONIC ACID v. CHLORO-BENZYL-MALONIC ACID.

BENZYL-CINCHONINE v. CINCHONINE.

a-BENZYL-CINNAMIC ACID C, H ,O2 i.e. [157°]. C,H,C11:C(C11,Ph).CO,H. Formed by the action of alkalis on the compound Ph.SO, C(C11,Ph), CO,Et (Michael a. Palmer, Am. 7, 70). Large white needles, insol. water, sol. alcohol.

p.BENZYL-CRESOL C.H., CII, C.11, Me.OII. (240°) at 40 mm. From benzyl chloride, cresol, and zinc (Mazzara, G. 8, 303; 11, 438; 12, 264).

Reactions. - 1. Chloro - acetic acid and Reactions.—1. Unioro - acetto deta and ROHAq form C.H., Cill., C.H., M.H.O. C.H., CO., LI [1119].—2. (a) - chloropropionic acid forms C.H., C.H., C.H., Me.O. C.H., C.H., C.H., C.H., and Na forms C.H., C.H., C.H., (0H), CO., Na. Acetyl derivative C₁₄H_{1,2}AcO. (245°) at

BENZYL CRESYL OXIDE C.H. CH, O.C. H. CH, v. Tolyl ether of BENZYL ALCOHOL .

C.H., CH., NH.CN. [83°]. Formed by passin CyCl Into bensylamine in ether (Strakosch, H 5, 694). Plates (from other); insol. water, v. e sol. alcohol and ether. On keeping it changes to isomerio tri-benzyl-melamine. Boiling HC forms benzyl-urca.

Di-benzyl-cyanamide (C. II CH.) N.ON. [549] From CyCl and dibenzylamine in alcohol (Limpricht, A. 114, 317). Plates; insol. water

BENZYL CYANATE O.H. CH. N.CO. (1750. 2003). Formed, together with benzyl evanurate, by the action of silver cyanate on benzyl ohloride or bromido (Letts, C. J. 25, 416; Ladenburg a. Strave, B. 10, 46). Pungent liquid. Changes spontaneously into the cyanmute. Alcoholio NII, converts it into benzyl-urea.

BENZYL CYANIDE v. PHENYL-ACETONITRILE. BENZYL CYANURATE (C.H.CH.), N.C.O., [157], (above 320°). The chief product of the action of silver cyanate on benzyl chloride (r. supra); formed by isomeric change from benzyl cyanate. Silky needles (from alcohol), insol, water. Potash fusion gives K CO, and benzyl-

BENZYL CYMENE Call a i.c.

C.H. CH. G.H. McPr. [2975] (Mazzarn, G. 8, 508; (308 | (Weber, J. 1878, 402). S.G. P 97. From benayl chloride, cymene, and zinc. On oxidation if gives benzoyl-terephthalicacid.

Benzyl-cymeae disulphonic acid

C. II., (SO. II) (M.). BENZYL DURENE v. BENZYL-TETRA-METHYL-

BENZYLENE (C.11a)2. Two hydrocarbons of this composition are formed by the action of the copper-zine couple upon benzyl brounde (or chlaride). (a)-henzyleno, [12]; μ_{ν} 5091, is a yellowish-red resin, sl. sol. alcohol, v. sol. ether and benzene. (β)-benzyleno is a brown resin, insol. alcohol or ether (Gladstone a Tribe, C. J. 47, 418).

BENZYLENE . BENZYLIDENE .

BENZYLENE-DIAMINE v. AMIDO BENZYL-

BENZYLENE-IMINE C,II,N i.e. CII2

[1:2]. Formed by reduction of o-nitro-henzyl-chloride with SuCl, in conc. HCl. Greyish yellow pewder. Sol. chloroform and acetic acid. Disselves in HCl to a red fluorescent solution. The salts are amorphous. The hydrochloride forms a reddish-yellow transparent solid (B HCl). The platinochloride (B', H.PtCl,) is an insoluble, amorphous, reddishbrown powder (Lellmann a. Stickel, B. 19, 1611).

BENZYL ETHER Call, O i.e. (Call CH2), O. Di benzyl other, Di-ben yl axide. (298° i. V.) Ε G. 18 1 036. μ 1 5525. Formed by heating benzyl alcohol with B.O. at 120° (Cannizzaro, At, 92, 115). Also by heating benzyl chloride with water at 190° (Limpricht, A. 139, 313). From benzyl chloride and sodium benzylate (Lone, C. J. 51, 700). Decomposed by heat into toloene and benzoic aldehyde.

BENZYL-D1-ETHYL-AMINE C, II, N i.s. C.H. CH NEt... (212° cor.). From benzylamine and Etl at 130° (Ladenburg a. Strave, B. 10, 47, 561, 1152, 1634); or from di-cthyl-amine and benzyl chloride at 100° (V. Meyer, B. 10, 810,

Ethyle-iodide C.H. CH. NEt,I. Large crystals, v. sol. water. On dry-distillation it gives triethylamine and henzyl iodide. C.H., CH., NEt. I. [87°] -- (C.H., CH., NEt. Cl.). PtCl., Di. benzyl-ethyl-amine C., H., N i.e.

(C.H., CH.), NEt. From di benzyl amine and Etl (Limpricht, A. 144, 315).—B'HCl.

Ethylo iodide (C.H., CH.) NEt,1. Sl. sol.

cold water.

BENZYL ETHYL BENZENE C, H, 6 i.e. C, H, CH, CH, Kt (1:4). Mol. w. 196. (295° i.V.). S.O. 12 '90. From henzyl chloride, ethylbenzene, and zinc (Walker, B. 5, 686) or from p. ethyl-benzophenoue, HI, and P (Söllscher, B. 15, 1682). Oxidation gives p-benzoyl-benzoic

RENZYL ETHYL OXIDE v. BENZYL ALCOHOL. BENZYL ETHYL KETONE C10 II 120 i.e. C. H. CH. CO.C. H., Mol. w. 148. (c. 226). 8.0. 25 1 00. From phenyl acetic chloride and with NaHSO, Oxidised by CrO, to benzoic and propionic acids.

BENZYL, p. ETHYLPHENYL - CARBINOL C.H., CH., CH., CH., CH., (350).

Liquid. Fermed by heating benzyl othylphenylketoos with alcoholic KOH at 160°. Boiled with dilute H.SO, it gives phenyl ethylphenyl ethylene (Söllscher, B. 45, 1681).

BENZYL ETHYLPHENYL p. KETONE O.H. CH. CO.C. II. C.H. [1:1]. Ethyl-desoxyben-zoin. [61]. Prepared by the action of Al.Cl. on a mixture of ethyl-benzene and phenyl-acetyl chloride (Sollscher, B. 15, 1680). Boils undecomposed. V.D. 8403 (ebs.). Small plates. Sol. ether, benzene and hot alcohol, sl. sol. cold alcohol. On oxidation it gives terephthalic acid. On reduction it gives phenyl-ethylphenyl-

BENZYL . ETHYLPHENYL . METHANE v.

PHENYL-ETHYLPHENYL-LTHANE. BENZYL DI ETHYL SULPHINE. chloride (C, H, CH, SEt, Cl) PtCl, From Etl and di-benzyl sulphide, the product being treated with AgCl and PtCl4 successively (Schöller, R. 7, 1271).

DI-BENZYL-ETHYL pseudo-THIOUREA

G, H, N, S i.e. C, H, S, C(NH,); NH. Formed by heating di-benzyl thiourea with ethyl iodide

Salts. B'HI: '93'); monochnic prisms; v. sol. alcohol, sl. sol. water. B'H SO, : large foursided rhombic soluble tables. -B',II.PtCl .: fine needles (Reimarus, B. 19, 2349).

BENZYL FLUORENE Callie i.c.

 O_aH_a , $CH_aC_aH_a$ $< \frac{CH_a}{C_aH_a} >$, [102°]. Formed by heating fluorene with benryl chloride and zinc dust (Goldschmiett, M. 2, 413). Plates (from alco-

hol).
BENZYL FORM ALDEHYDE v. PHENYL-

DI. BENZYL-GLYCOLLIC ACID C, H, O, i.e. (C.H.CH.).C(OH)CO.H. Oxatoluic acid. a.Oxydi-phenyl iso butyric acid. [157°].

Formation. I. From the nitrile and cono. HCl at 140°-160° (Spiegel, A. 219, 46; B. 13, 2219; 14, 1687).—2. By boiling vulpio acid (q. v.) with aqueous KOH: C₁₇II₁₄O₄+3H₄O

= CH,O + 2CO, + C,H, O, (Möller a. Strecker, 4. 118, 56).

Rhombie prisms (from alcohol): a:b:e= ·5113:1: ·3058, Finffy mass of needles (from benzene). Salt .- AgA'.

Reactions.-1. HNO, gives a visoid nitroacid (Möller a. Streeker). -2. Cono. aqueous KOH gives toluene and oxalio acid on boiling. -3. Treated with PCl, and H₂O successively a monoplosphate, C₁₄H₁O₂PO₁H₂, orystallising in prisms, [160°] is formed.

Acetyl derivative. [106°]. Plates in rosettes (from CHCl, mixed with petroleum).

Methyl other MeA': [71°]; neodlee. Anhydride C, II, O, [169°]. Got by heating the acetyl derivative. Prisms (from benzene). V. sel. alcohol and ether. Na CO, Aq converts it into sodium di-benzyl-glycoliate.

Nitrite(C.H.CH.), C(OH), CN. [113°]. From the benzyl kelone, KCN and HCl. Colourless flut rhombs (from alcohol). At I13° it splits up into HCN and di-benzyl ketone.

Amide. [1939]. From the nitrile and cone. IICl at 125°. Fluffy mass of long needles.

BENZYL - GLYOXALINE C₄H₄(C,H₄)N₂.

[71°]. (310°). Formed by the action of henzyl chloride on glyoxaline (Wallach, B. 16, 539). Colonriess crystals. Sl. sol. ether, insol. cold water.—B',H,Ci,PtCl,: yellow pp., insol. cold water.

DI-BENZYL-GUANIDINE C. H. N. i.e. (C.H.CH.NH), C;Nif. [100']. Formed by boiling benzylamine hydrochloride with benzylcyanamide in alcohol (Strakosch, B. 5, 695).-Lamina (from alcohol). Sol. water, alcohol, and other. - B'HCl (1759).

BENZYL HYDROXYLAMINE v. HYDROXYL-

BENZYLIDENE. The radiale C. H., CH, also called benzal or benzylene. The latter name is more appropriate to the isomeric radiels C.H.CH.

BENZYLIDENE DI ACETAMIDE

Cullin N.O. i.e. Call CH (NH.CO.CH.). Formed by boiling acetamide with benzoic aldehyde (Roth, A. 154, 72; Z. [2] 4, 650; 6, 680). Silky crystals, sl. sol. cold water and ether. Not affected by boiling KOHAq but decomposed by hot HClAq into benzoie aldehyde and NH2Ao.

BENZYLIDENE DI ACETATE C, II, O, i.e. CHPh(OAc)... Di acetyl-ben oic ortho-aldehyde. [45"]. From benzoic aldehydo and Ac₂O or from benzylidene chloride and AgOAc (Genther, A. 106, 251; Wicke, A. 102, 368; Hübner, Z. 1867, 277; Neuhof, A. 146 323; Limpricht, A. 139, 321; Perkin, Z. 1868, 172).

BENZYLIDENE ACETIC ACID v. CINNAMIO

Benzylidene-di-acetic acid v. PHENYL-OLU-TARIC ACID.

BENZYLIDENE . ACETO . ACETIC ACID v. p. 21.

BENZYLIDENE DI ACETO ACETIC ETHER

C,, 17, O, tr. C, 11, CH {CH.CH(CO,Et).CO.CH, }, (?). [153°]. Formed together with dehydro-benzylidene di acetoacetie ethor C10H22O3, hy the action of benzoic aldehyde (I mol.) upon acetoacotic ether (2 mole.) in presence of a primary amine. Long white needles. Sl. sol. cold alcohol and ether (Hantzsch, B. 18, 2588).

Behydre - bennylidens - di - acetoacetie - ether Ph C,H,O, possibly EtCO, -C-CH-C-CO,Et.

 $Me\ddot{C} = O = \ddot{C}Me$ [88°]. Formed as described abova. Glistening prisms. V. sol. cold or hot solvents (Hantzsch,

B. 18, 2583). BENZYLIDENE-DI-ACETONAMINE v. p. 27. BENZYLIDENE ACETONE CioHioO i.r.

Ph.CH:Cll.CO.Cll₃. Aceto-cinnamone. Methyl-styryl ketone. [42°] (152°) at 25 mm. (261° i.V.) at 760 mm.

Formation. - 1. From benzeic aldehyde, acetone and a little ZnCl, at 260° (Claisen a. Claparède, B. 14, 2461). · 2. By heating a mixhyde with Na and MeI at 130°, or by boiling cinnamic aldehyde with McOH and ZuCl. (Engler a. Leist, B. 6, 254).

Preparation. - From benzoic aldohyde (20 g.), acetone (40 g.), water (1800 c.c.) and aqueous (10 p.c.) NaOH (20 g.) in the cold. After four days the oil that has separated is extracted with athar, dried over CaCl, and rectified in vacuo

(Claisen a. Ponder, A. 223, 138).

Properties. - Plates, apparently rectangular. It has an odour of coumarin and rhuburh and attacks the skin. Easily solubla in ulcohol, ather, benzene, and chloroform, less in petrolaum-ether. In conc. II.SO, it forms an orange solution. Forms a crystallino compound with NaHSO3, and a di-bromide C10H110Br2 [125] crystallising in needles from alcohol.

Phenyt hydraside C, HaN2: [1567]; flat yellow needles; sol. hot alcohol, sl. sol. cold alcohol and ether, insol. water (Fischer, B. 17,

576; Knorr, B. 20, 1099).

Ph.C1I:Cl1.C(NOII).Cll3. Oxim (220°) at 100 mm. Forms a bromide, [115]; and an acatyl derivative [91°] (Zeliusky, B. 20, 922).

Di bsnzylldine acetone

PhCH:CH.CO.CH:CH.Ph. Cinnamone. styryl ketone. [1127]

Formation. - From benzoic aldehyde (20 pts.), acetona (6 pts.), and acetic acid (40 pts.), by

adding H.SO, (30 pts.) at 0° or passing in 11Cl. Preparation. - From benzoie aldehyde (10g.), acetone (3 g.), water (100 g.) and dilute (10 per cent.) NaOH (20 g.) left 4 days in the cold; or from benzylidene acetone (7 g.), benzoic aldeliyde (5 g.), water (200 g.), alcohol (150 g.) and diluto NaOH (20 g.) (Claisen a. Ponder, A. 223, 142; cf. Claisen a. Claparede, B. 14, 350, 2460; Schmidt, B. 14, 1459).

Properties .- Bright yellow monoclinic tablets (from acetona or CHCl₂) a:b:c = 4.986;1:1.378 $\beta = 78^\circ$ 43'. Readily soluble in chloroform or acetona, less in ether or cold alcohol. Aives

an orange solution in H.SO... Tetrabromide C₁₇H₁₄OBr₄: white needles [208°-211°] BENZYLIDENE. DIACETONE. ALCAMINE

H2C-CH(OH)-CH2 C.H.NO i.e.

 $(C_*H_*)HC - NH - C(CH_*)_*$ Oxy-phenyl di-methyl-tetra-hydro pyridine. Thick colonriess oil. Formed by reduction of an acid solution of benzylidene-di-acetonamine sidehyde and aniline hydrochloride dissolved

(v. p. 27) with sodium-amalgam. - B'HCl; amail (c. p. 27) when southern angents of the corystals, easily soluble in water and alcohol (Fischer, B. 16, 2936).

BENZYLIDENE-DI-ACETONINE v. p. 34.

BENZYLIDENE - ACETOPHENONE

C.H., CH:CH.CO.C. H., Phenyt styryl ketone, Phenyl cinnamenyl ketone. Benyli lene methyl phenyl ketone. Benzal-acetaphenone. (345° 348°). Formed by passing HCl gas into a mixture of acetophenone and benzaldehyde; by adding H.SO, to the two latter bodies di-Inted with neetic acid; by heating them with neetic anhydride to 170°; or by treating them with dilute NaOH (Claisen a. Claparède, B. 14, 2163; Claisen a. Ponder, A. 223, 148).

Preparation .- 12 pts. of acetophenone ara ture of calcium acctate with calcium cinnamate; mixed with 10.5 pts. of heuzablehyde and 3 pts. also in small quantity by heating cinnamic able of a 20 p.c. sodium methylate solution, and allowed to remain in the cold for a few days when the whole will have solidified to a crystalline mass; the yield is 90 p.c. of the theoretical (Claisen, B. 20, 657). Large trimetric tables. V. sol. chloroform, ether, benzene und CS, m. sal. alcohol, sl. sol. petrolemu ether. On axidation it gives henzoyl formic and benzoic acids. On boiling with dilute acids it is decomposed into acctophenous and benzaldelivde. By reduction with III and P it is converted into di-The 11Cl addition product henzyl-methane. C.H. CO.CHCl.CH .. Call., [120] forms colourless trimetric plutes, sparingly soluble in cold alcohol and ether. It is prepared by the addition of gaseons HCl to the ketone. The di-bromids gaseons HCl to the ketone. The di-bromids C.H., CO.CHBr.CHBr.C.H., [157] forms short colourless prisms sol. hot alcohol. Prepared by the addition of bromine to the ketone.

BENZYLIDENE . B . ACETYL . PROPIONIC ACID C .: H.c.O., i.e. CHPh: Cll., CO.CH, .. Cll., CO.H. Benzylidene-lavulic acid. Cinnamoyt-propionic acid. [120'-125']. Formed by heating levulic acid with menzoic aldehyde and sodium acetate. Small white crystals. The lead sult is insoluble. Dissolves in cold conc. H.SO, with a red colour-Boiling cone. KOH splits off benzaldehyde. Reduction in alkaline solution yields benzyl-valero-lactone C12H14O2 which forms large prisms of molting-point [85°] (Erdmann, B. 18,

3441). BENZYLIDENE-ISOAMYL-AMINE. An oil formed from benzoic aldehydo and isoamyl-

amine (Schiff, A. 140, 93).

BENAYLIDENT DI -180AMYL DI - OXIDE
C, H,CH(OO,H₁). (292° cor.). From benzylideno chloride and NaOC,H₁₁ (Wicke, A. 102,

BENZYLIDENE - ANILINE C. H. N C.H. CH:NPh. Aville of benzoic aldehyde, [49°] (Tiemann a. Piest, B. 15, 2028).

Formation. -1. By warming anilina with benzoic aldehyde (Laurent a. Gerhardt, Compt. chim. 1850, 117).-2. By heating di-phenyl-thio-urea with benzoic aldehyde (Schiff, A. 148,

Properties .- Laminæ; volatila with steam, insol. water, v. e. sol. alcohol and other. At 200° it changes to an isomeride which differs

from it in forming a salt L', H₁P'Cl_s.

Hydrocyanide C₁₁H₁₂N₂. [82°]. Formed
by passing HCN into fused benzylidene-aniline, or by the action of KCN on a mixture of benzois

in sloohol (Cech, B. 11, 246). It forms concentric needles, insol. alkalis and dilute acids.

BENZYLIDENE-DI-ANTIPYRINE v. DI-OXX-TETRA-METHYL-DI-QUINIXYI .- PHENYL-METHANE

DI BENZYLIDENE BENZIDINE C₂H₃N, t.e. O_cH₄(N:CHPh)₂. [232'] (C.); [239'] (B.). Obtained by heating hydrazo-benzene or benzene-azo-benzene with benzoic aldehyde and ZnCl, (Chève, Bt. [2] 45, 188; Barzilovsky, J.R. 1885, 366). Yellow scales (from benzene and chloroform). Resolved by HCl into benzeic atdeliyde and benzidine.
BENZYLIDENE. DI-BENZAMIDE

 $C_{ij}H_{ij}N_{ij}O_{ij}$ i.e. $C_{ij}H_{ij}CH(NH.CO,C_{ij}H_{ij})_{ij}$ Formed by heating benzoic aldehyde with benzamide (Roth, A. 151, 76). Long silky needles (from alcohol), insol. water. Resolved by hot HClAq into the parent substances.

BENZYLIDENE DI BENZOATE

C.H.,CH(OBz).. From benzylidene chloride and AgOBz (Engellardt, J. 1857, 471). Crystalline.

BENZYLIDENE BENZYL-AMINE

C_aH_aCH:N.CH_aC_aH_a (c. 300°). Formed by the action of PCl₁ upon di benzyl hydroxylamine, probably by intermediate formation of the chloride (C, H, CH,) N.CL. Od. V. sol. alcohol and ether, insol, water.

Salts. Buch: [251]; long plates tables; v. sol. nicohol, spiringly in cold water, more readily in hot. Bull lach : small golden yellow crystals (Wabler, It. 19, 1632).

BENZYLIDENE BROMIDE CH Br. C.H. CHBc. Benzal bromble, w Di bromo-toluene. (130° 140°) at 20 mm. From herzoic aldehyde and Pitr. Can only be distilled in vacuo. So tum at 180 forms toluene and dibonzyl (Michaelson a. Lippenann, Bl. (2) 4.

BENZYLIDENE BROMIDE BENZOATE

 $C_{\rm D}\Pi_{\rm H}B(O_{\rm g}) \text{ i.e. } C_{\rm g}\Pi_{\rm g}C\Pi \underset{\rm O}{\underbrace{\rm CO}_{\rm CO}}\underset{\rm CO}{\underbrace{\rm CO}_{\rm g}}\Pi_{\rm g}^{\rm o} = [70^{\circ}].$ Colourbes tables or prisms. Sol. alcohol, ether, and aceta acid. Prepared by mixing bearable hy lo and benzoyl bromide. On distillation it again decomposes into these bodies (Claisen, B. 14, 2475; cf. Liebig a. Wöbber, A. 3, 266).

BENCYLIDENE DI BUTYRAMIDE

C. H. S.O. i.e. C. H. CH(NH.CO.C.H.). Slender crystals formed by heating butyramide with benzoie ablebyde (Strecker, A. 154, 76).

BENAYLIDENE DI CARBAMIC ACID

CALCU-VILCOAD.

Ethyl other Et A". (171'). From carbamic other, bensoic aldehyde, and HC (Bischoff, B. 7. 634). Crystals; may be sublined. Uroppi ether Pr.A". [C13] (Bischoff, B. 7.

10 (2).

BENZYLIDENE CHLORAL AMMONIA

C.H.Cl.NO i.e. CCl., CHPOTONICH.C.H., [1307]. White leathets. Decomposed by dilute neids and by boiling water. Prepared by the action of benaoic aldehyde on chioral ammonia (Schiff, B. 11, 2166).

BENZYLIDENE CHLORIDE C.H.Cl. i.e. D.H., CHCl., Bental chloride, Chlorobenzol, Bentylene chloride, Mol. w. 161, (2047) at (204°) at 156 mm. S.G. 2 1.27. S.V. 154.25 (Schiff, B. 19, 563).

Formalion, -1. From benzoic aldehyde and Caliours, A. 70, 39; Suppl. 2, 253, 306).

1. By passing shieries 145 bolting tolume. (Belistein, 4. 115, 536; 146, 322; Lauth a. Grimaux, Bl. 2, 347).—B. From benzoic aldehyde and succinyl chloride (Rembeld, A. 188,

189) or OOCl. (Kempt, Z. 1871, 79).

Preparation.—1. By passing 2 mols. of chlorine into cold tolueno (1 mol.) exposed to direct sunshino (Schramm, B. 18, 608).—2. By heating tolucne (7 pts.) with PCl, (30 pts.) at 190°; the yield being nearly that calculated (Colson a. Gautier, Bl. [2] 45, 87).

Properties .- Oil, with faint odour. Ileactions. -1. Converted into benzoic aldehyde by water or aqueous K,CO, at 130°, or by warming with H,SO, at 50° and treating the product with water (Oppenheim, B. 2, 213).— 2. Alcoholio KHS forms benzyl disulphide and di-thio-benzoio acid.—3. Red-hot sodu lime forms benzene (Limpricht, Bl. 1866, ii. 467).— 4. Chlorine forms p-chloro-benzylidene chloride. 5. Nitric acid forms p-nitro-benzylideno chloride (Hübner a. Bente, B. 6, 803; cf. Beilstein, .1. 146, 333).-6. AgOAe forms C.H., Cli(OAc), 7. Silver oxalate forms benzoic aldehyde (Golowkinsky, A. 111, 252).-8. Na forms di-phenylethylene. - 9. McI and Na form onmene. 10. NH, forms hydrobenzamide. -11. ZnEt, diluted with benzene forms $C_{11}H_{16}$ di-cthylphenyl-methane and $C_{21}H_{16}$ (Dafert, M. 4, 618), 12. Copper at 100° gives CPhCl₂CPhCl₂ and CPhHCl.CPhHCl (Ounfrowicz, B. 17, 833)

BENZYLIDENE . D1 - CHLORO . CHROMIC ACID P. TOLUENE

DI BENZYLIDENE. ETHYLENE . DIAMINE C₁₀H₁₆N₂ i.e. C₂H₃(N;CH.C.H.)₂, [54°]. Formed by henting ethylene-diamine (1 mol.) with benzoic althehyde (2 mols.) to 120°. Large colour-less tables. V. sol. alcohol and benzene, insol. water. Decomposed into its constituents by acids (Mason, B. 20, 270).

BENZYLIDENE ETILYLENE DI SULPHIDE

S.CH, . (29°). Formed by passing \s.ċu.

If Cl gar into a mixture of equal neels, of benzaldehyde and ethylene sulphydrate. Crystals. Easily soluble in alcohol, ether, and benzene, insoluble in water. Very stable body. By the nction of bromine upon the chloroform solution

disthylene-tetra sulphide $C_1H_1 < \frac{S_2}{S_2} > C_2H_4$ formed (Fashender, B. 20, 460).

BENZYLIDENE DI ETHYL DI OXIDE

Call CityOEt), (222' cor.) Diethyl derivative of benzoic ortho-aldehyde. From benzylidene chloride and NaOEt (Wicke, A. 102, 363). BEN TYLIDEN & DI. HEPTYLENE TEIRA-

UREA C. H. N.O. i.e. C.H. CH(NILCO.NILC.H., NH.CO.NH.) From henzoic aldehyde and heptylene-diures (Schiff, A. 151, 195). Insoluble powder. BENZYLIDENE MALONIC ACID

Ph.CH:C(CO II), | 1962 (C.); [1932] (S.). Envantion.—1. From the other by aqueous baryta. -2. From benzoic aldeliyde, malonic acid and Ac O at 100° (Claisen a. Crismer, A. 218, 135). -3. From benzoic aldehyde, sodio malouate, and glucial acetic acid at the ordinary temperature: Ph.CHO + CH (CO,Na), =
Ph.CH.C(CO,Na), + H.O.

The product is diluted with water, shaken out

with other, addited, shaken again with other and the other distilled off (Stnart, C. J. 43, 405; p. also Claisen, A. 218, 129).

Properties.-Colourless glassy prisms. sol. cold water, v. e. sol. bot water, v. sol. alcohol, acetic ether or acetone, m. sol. ether or glacial scetic acid. Insol. benzene, chloroform or petroleum. Does not give in neutral solutions a pp. with BaCl2; but on warming such a mixture needles of the salt BaA" separate (characteristic reaction). Salt. -- Ag.A".

Reactions .- 1. At 200° it splits up into CO, and cinnamic acid .- 2. Boiling water Jecom. poses it into henzoic aldehyde and malonic acid, somo CO, and cinnamic acid being also formed. 8. Reduced by sodium amalgam to benzyl malonic acid Ph.Cll.Cll.(CO.II).—4. Browing acts on its solution in chloroform forming PhCHBr.CBr(CO.11), [96] whence water forms a-bromo-cinnamic acid (Stuart, C. J. 49, 360). 5. Cold alcoholic potash forms crystals of Ph.CH(OEt)CH(CO,K), whence a silver salt, PhCH(OEt)CH(CO,Ag), may be got. The free acid, if heated rapidly, melts at 120°-130°, sphtting up into EtOH and beuzylidene-malonic acid, which then solidifies again, and melts a second . time at 1903. By crystallisation from water the acid is partly split up into alcohol and benzylidene malonie acid. - 6. HBr forms & brome phenyl-isosuccinie acid Ph.CIIBr.CH(CO.H), which is decomposed by water into IIBr, cinnamic acid and CO, (Stuart, C. J. 49, 360).

Ethyl ether £t,A". [32]. (192°) at 17mm. S.G. 14 I'll. Formed by passing IICl into a mixture of malonic ether and benzoic aldehyde or by treating the mixture with Ac O at 160° (Claisen, B. 14, 348). Large transparent crystals (Stnart, C. J. 49, 360). Boils at 3085-3120 with decomposition.

BENZYLIDENE MESITYL OXIDE Calle, O i.e. (CH_a) C:CII.CO.CII:CII.C.H_a. (179°) at 14 mm. Oil. Prepared by passing HCl gas into a mixture of mesityl oxide and benzaldehyde (Claisen. Forms a tetrabromide [1182].

BENZYLIDENE METHYL KETOLE C., 11, N. [248°]. Colourless plates. Formed by reduction of dimethyl-resindelo CaH 20N2 with zine-dust and NH, It is oxidised in acctic acid solution by Fe,Cl, back to di-methyl-rosindole (Fischer a. Wagner, B. 20, 816).

BENZYLIDENE DI METHYL DI OXIDE C.H.CH(OMe), Di-methyl derivative of benzoic orthoaldchyde. (208° cor.). From henzylidene chloride and NaOMc (Wieke, A. 102, 363)

BENZYLIDENE D1 - METHYL p-PHENYL ENE-DIAMINE Call, CH: N.Call, NMog. Itenzylidene - amido - di - methyl - aniline. phenylene-diamide of benzoic allehyde (98°). Glistening plates or needles. So hot alcohol and benzene, sl. sol. cold alcohol. Weak base. Formed by mixing benzoic aldehyde and u-dimethyl-p-phenylene-diamine, either directly or in alcoholic solution. By HCl it is split up into its generators.—B"H,Cl,: white solid (Calm, B. 17, 2940).

BENZYLIDENE . (Py. 3) . METHYL . QUINOL . CH:CH

1,H, N: C—CH = CH.C,H,(?), [100°]. Formed NaOAe (1 pt.) (Gabriel a. Michael, B. 11, 1018). Vos. I.

by heating quinaldine (methyl-quinoline) with benzoic aldehyde or benzylidene chloride an ZnCl, (Jacobsen a. Reimer, B. 16, 2606). Glis tening colourless needles. Sublimable. Sol. ho alcohol, insol. water.—B'H,Cr,O, 23nq: very sparingly soluble reddish-yellow needles

BENZYLIDENE . (β) . NAPHTIIYLAMINE C. II. C. II. N. C. II. (103°). From benzoic aldehydd and 18) naphthylamine (Claisen, A. 237, 261).

BENZYLIDENE. (a) NAPHTHYLAMINE.p. SULPHONIC ACID C H (N.CH.C. H J. SO. 11[1:4]. The sodium salt (A'Na) is obtained by shaking a strong solution of so linu. (a) naphthylamine sulphonate with benzaldehyde. By long boiling with water it is split up into its constituents (Cahn a. Lange, B. 20, 2001).

BENZYLIDENE DI (8) NAPHTHYL OXIDE C.H.CH C. II.O. Anhydride of di-oxy-dinaphthyl-phonyl-methane. [190"]. Crystalline

solid. Insoluble in aqueous alkalis. Formation, -1. By heating a solution of (8) maphthol and benzoic aldelivde in acctic acid to 200', or with addition of H.SO, or HCl on the water-bath (Trzeinski, B. 17, 199).-2. By warming benzylidene di-naphthyl di-oxids Call_CH(O.C., H.), with nectio neid and a few drops of HCl. In these reactions di-oxy-dinaphthyl phonyl methane C.H. CH(C. II OH) must first be formed and at once split off H.O

(Claisen, B. 19, 3317). Benzylidene di-(8)-naphthyl-di-oxide

C₂₁H₂O₁ i.e. C₃H₁CH(OC₂H₁)₂. Di-naphthyl ortho-ben-aldehyde, [205]. From benzois aldehyde (5.3 pts.), (θ) -naphthol (7.2 pts.), glacial HOAc (30 pts.), and furning HCl (2 pts.) nt 0° (Claisen, A. 237, 269). Tables, sl. sol. CHCl, and CS_n, v. 81. 50.1 alcohol and other, insol. alkalis. Conc. II, SQ gives, on gentle warming, a deep-red solution. HOAc and some HCl slowly convert it at 100° into henzylidene. di-naphthyl oxide, a change which also occurs when it is heated at 210°.

BENZYLIDENE OXAMIDE C.H.N.O. Formed by warming oxamio other with benzoic aldehyde (Mcdieus, A. 157, 50). Lamine.

BENZYLIDENE . PHENYL . DIAMINE PHENYL-BENZAMIDINE.

DI-BENZYLIDENE.p-PHENYLENE AMINE C. all, N. i.c. C.H. (N.CIIPh), [140°]. From p-phenylencadinuine and benzoic aldelyde at 120° (Ladenburg, B. 11, 590). Plates (from alreadot). Resolved by hot HORA into the parent substances.

BENZYLIDENE.PHTHAB-ETHYL-1M1D1NE C1, H1, ON i.e. C. II. C (= CII.C, II.) NEt. Ethyl-

phthal.myi-benzyl. [c. 77°]. Formed by hoiling the ethyl-amide of decaybenzon-o-carboxylio acid C, II, (CO.NHEt) CO.CH, C, II, with acetic acid. Plates. V. sol. alcohol, benzene, benzeline, and CS, (Gabriel, B. 18, 2433).

BENZYLIDENE-PHIHALIDE C, II, O. i.e. $C_{\bullet}H_{\bullet} < C:(CHPh) > 0.$ [99°]. Benzylidene. phthatyl. Anhydride of deoxybenzoin carboxylic acid.

Formation. 1. By heating phthalic anhydride (5 pts.) with phenyl-acetic acid (5 pts.) and

2. By heating phthalyl-phenyl-acetic acid in cacto (Gabriel, B. 17, 2526).

Preparation .- A mixture of 100 g. phenylacetic scid, 110 g. phthalic anhydride and 21 g. dry sodium acctate is heated for 2 hours, and the product crystallised from alcohol; the yield is 75-78 p.e. (Gabriel, B. 18, 3170).

Properties.-Long prisms (from alcohel); insol. water, sl. sol. cold alcohol.

Reactions. -1. Hot aqueous KOH forms potassium deoxybenzoin earboxylate. - 2. By hoating with alcoholic NH, at 100' it is coninto droxybenzoin - earboxylamide C₄H₄CO.NH₂ which by solution in H.SO₄ or by boiling with glacial acetic acid loses II20 giving benzylidene-phthalimidine (phthalimidylbonzyl) C₄H₄ C (CII.C₄H₄) NII. -3. Similarly ethyl amine yields the ethyl-amide of deoxybenzoin earhoxylic acid, and this on boiling with acotic acid gives benzylidene-phthal-cthylimidino C, H, C(~ CII.C, II,) NEt (Gabriel, B. 18, 2133). - 4. By dissolving in benzeno and treatment with nitrous acid gas it yields the compound C, II, (C(NO₂).(CII(NO₂).C₄II₃)O CÓ (Gabriel, B. 18, 1251).

References .- V. Cyano. and Nitao-BENZYL-EDENC-PHTHALADE.

Benzylldene-phtbalide-di-bromide

 $\mathbf{C}_{a}\Pi_{a} \stackrel{\mathrm{CBr}(-\mathrm{CHBr}, C_{a}\Pi_{a})}{\sim} 0.$ [146°]. Formed by the combination of benzylidene phthalide with bromine (Gubriel, B. 17, 2527). Thick glistening prisms. Sparingly soluble in alcohol.

(Iso) - Benzylidene - phthalide C1311100, i.c. CII:C.C.H. OH . [91°]. Formed by reduction \co.ò

of nitro-benzylidene phthalide

Call CO CO Swith HI and P. Prepared by reduction of nitro-benzylidene-phthulide; yield, 47 p.o. of the phenyl-acetic soid employed to prepare the benzylidene-phthulide (Gabriel, B. 18, 3171). Flat colourless needles. Easily soluble in alcohol and benzene, sparingly in ligroin. By further reduction with HI and I at 200° it yields s-di-phenyl ethane c-carboxylic acid C.H., CH., C.H., CO.H. lly boiling with aqueous NaOl1 it is converted deoxybenzou · o · enrhoxylio Call (CO,11).CH, CO.C. II, Heated with alcoholic gives eisebenzylidene phthalimidine

C11:0.0.11, (Gabriel, B. 18, 2445). ∕со.хи

BENZYLIDENE-PATHAL-IMIDINE ≻ с ⊆ сн.с**.н' €**_nH_nON i.e. C_sH_s< Phthal.

imidyl-benzyl. [1833]. Yellow plates. Formed from the amide of deoxybenzoin-o carboxylie meid C, H, (CO, NH,), CO, CH, C, H, by solution in 41 SO, or by boiling with acetic acid (Gabriel, B. 18, 2433). I'. also Baome- and Nitrio errzyl-TOURE PRODUCEDED

Isobouzylidene-phthalimidine v. (l'y. 4)-Oxv-(Py. 2) PHENYL-ISOQUINGLINE.

BENZYLIDENE DI PIPERIDINE

C.H. CH(C.H. N). [81°]. Formed by heating piperidine with benzoic aldehyde. Colourless prisms. Very unstable, being decomposed even by boiling with wator. Dilute acids resolve it into benzoic aldohyde and piperidine (Laun, B. 17, 678).

BENZYLIDENE-RHODANIC ACID C10 11,NS2O i.e. C6113.CH:C(SH).CO.S.CN. [2009]. Formed by the action of benzoic aldehydo upon rhodanic acid in presence of dehydrating agents (Nencki, B. 17, 2278). Yellow needles, sol. water. By heating with baryta water it is split up into a sulphydro-einnanio acid C_eH_CH:C(SII).CO₂H and hydrogen sulpho-cyanide. Heated at 410° with cone. H₂SO₄ (4 pts.) it is converted into benzylidene-rhodanieoxy-sulphonic acid C, H,NS,O, (Ginsburg a.

BENZYLIDENE. RHODANIC.OXY.SULPHO. NIC ACID C1011,NS2O3. Formed by heating benzylidene rhodanic acid with cone. H2SO4 (4 pts.) at 110°. Needles. V. sol. water and alcohol. Very strong soid (Ginsburg a. Bond. zynski, B. 19, 119).

BENZYLIDENE ROSANILINE C27H13N2. From resamiline and benzoic aldehyde by heat or by shaking with SO, Aq (Schiff, A. 140, 111; Z. 1867, 176). - B'.11.PtCl.
BENZYLIDENE SELENIDE C.H.CHSe.

Bondzynski, B. 19, 119)

Seleno benzoic aldehyde. [70"]. From benzylidene chloride and alcohol K.So (Cole, B. 8, 1165). Yellow needles (fron water. Not attacked by NH_a. Yellow needles (from alcohol), insol.

BENZYLIDENE DI SKATOLE C. 11 2N2 i.e. PhC11(C,11,N), [142]. From skatole (2; pts.), benzoie allediyde (I pt.) and a little ZnCl. (Wenzing, A. 239, 241). Insol. water; v. sol. hot alcohol and other. Boiling HCl does not split off benzoic aldehyde.

BENZYLIDENE SULPHIDE v. Tuio-BENZO10 ALSKUYDE.

BENZYLIDENE THIO-BIURET C, 11, N, S, i.e. C,11_CH<N:C(SH) N1I(?). [237]. Formed by heating henzoic aldehyde with ammonium sulphocyanide at 137°-165° (Brodsky, M. 8, 27). Minute prisms (from alcohol); insol. water, sl. sol. cold alcohol; sol. dilute KOII. Boiling Boiling baryta-water forms benzoio aldehyde, barium

BENZYLIDENE o. TOLUIDINE C, 111, 18 i.e. C.H. CH:N.C.H. CH, [1:2]. Benzaldehyde o. toluide (314°). From e-toluidine and bonzois aldehyde (Etard, C. R. 95, 730). Resolved by boiling water into its generators. By passing through a tube heated to dull redness it is convorted into phenyl-indole C.H. CH CC. C.H.

(Pietet, B. 19, 1063). Benaylidene-p-toluldine

C.H., CII: N.C.H., CH., [1:4]. From benzeic alde-Nydo and p-toluidine at 100° (Sehiff, A. 140, 96; Kohler, A. 211, 359; Mazzara, G. 10, 370), Melts below 100°, but changes at 160° into an isomeride [120°-125°].—B'.H.PtCl.

DI.BENZYLIDENE TOLYLENE DIAMINE C., II., N. i.e. C. H.Me(N:CHPII), [1:2:4] [122-128°]. From benzoie aldehyde and tolylene-diamine at 100° (Schiff, A. 140, 98). Neutral erystalline mass; at 140°-150° it gives amarine.

BENZYLIDENE-DI-UREA C,H,2N,O, C.H., CH(NH.CO.NH.). Benzaldehyde di-ureide [1956]. Formed by adding benzoic aldehyde to an alcoholic solution of urea (Schiff, A. 151, 192). Crystalline powder, iasol, water and ether, sol. alcohol.

Di-boazylidene-tri-urea C. H., N.O., Powder, formed by heating urea with benzoic aldehyde.

Tri-bonzylideae-tetra-nrea Call N.O. [o. 240]. Powder, formed by heating benzylidene-di-urea with benzoic aldehyde.

BENZYLIDENE DI URETHANE v. Bunzyl IDENE-DI-CARRAMIC ACID.

BENZYL-INDOLE Callan i.c.

C.H. CII CII. [41-5]. From its carboxylic acid (q. v.) by heat. Yellowish needles (from alcohol). V. sol. benzene, light petroleum, chloroform and other. Turns pine wood mois-tened with HCl yellow. Picrate forms red needles.

BENZYL-INDOLE CARBOXYLIC ACID

 $C_{la}H_{la}NO_{2}$ i.e. $C_{s}\Pi_{4} < C_{NC,H_{2}}^{CH_{1}} > C.CO_{s}H_{s}$ [1952] with decomposition). Pyravie acid combines at 16° with benzyl-phenyl-hydrazine, forming *CH, C(CO H): N. NPhC H, whence 41Cl at 1005 forms benzyl-in-lol-carboxylic acid (Antrick, A. 227, 362).

Properties. - Colourless needles (from glacial acetic acid). Sl. sol. water, chloroform, and petroleum, sol. other and alcohol, v. sl. sol. Converted by heat into CO, and benzeue. benzyl indole.

BENZYL IODIDE C.H.I. i.e. C.H.CH.J. [34°], S.G. 72 1-73.
Formation. - 1. From benzyl alcohol in CS.

and iodide of phospharus. 2. Slowly formed by the action of cold III (S.G. 196) on beazyl chloride (Lieben, Z. [2] 6, 736).—3. From benzyl chloride and KI (V. Meyer, B. 10, 311; Kumpf, A. 224, 126), Zul,, or Pbl, (Brix, A. 225, 154).

Properties. - Crystals: decomposed by distillation. Gives benzyl nectate with AgOAc, and tribenzylamine with alcoholic NII, Silver nitrite gives benzoic aldehydo and acid (Van Renesse, B. 9, 1451; Brunner, B. 9, 1741). BENZYL - (pseudo) - ISATIN Callo NO. i.e.

C4H4 NC, H7 > CO. [131]. From benzyl-indole carboxylic acid and NaOCl in feeb'y alkaline solution, the insoluble chloride then produced being subsequently boiled with alcoholic NaOH (Antrick, A. 227, 365).

Properties. -Slender needles (from alcohol). Sl. sol. water, sol. ether. Shows the indophenine reaction with H.SO, and crude b uzene containing thiophene.

DI-BENZYL-KETONE C,II, O i.e. CO(CH.Ph). Di-phenyl-acetone. Mol. w. 210. [30]. (320?). Formod by the dry distillation of barium phonyl-acetate. Prisms. CrO, oxilises to benzoic and actio acids (l'opoft, B. 6, 560). Reduced by HI at 180° to di benzylmethane (Graebe, B. 7, 1623).

BENZYL-MALONIC ACID C₁₀|I₁₀O₁ i.e.

C. H. CH, CH(CO. H), Phenyl-isosuccinic acid. [1179].

Formation .- 1. By supenification of its ether. p. 25.

2. From bensylidene-malonic sold by sodiumamalgam.

Properties .- Triolinio crystals, sol. water, alcohol, and ether. Splits up at 180° into CO, nud B-phenyl-propionic acid.

* Ethyl other Et.A". (300°). S.G. § 1.08 (Conrad, A. 204, 174; B. 12, 752). Sodium benzyl-malonic ether is converted by iodine disselved in other into *C.11 CH CI(CO.Etl., which is converted by alcoholic KOil into ethoxy-benzyl-malonic ether (Bischoff a. Hausdörfer, A. 239, 110). Converted by alcoholic NII, into the amides CH.Ph.CH(CONII.), [225] and CH.Ph.CH(CO.Et)(CONII.) [98] (Bischoff a. Siebert, A. 239, 96).

Di-benzyl-malonic acid (C. 11 , Cll.) C(CO, ll)2 [1727], P. ; [1637] (B. a. H). Formed by suponifying the other (Perkin, C. J. 47, 821). Slender needles (from water) or thick prisms (from alcohol). V. e. sol. ether and alcohol, m. sol. hot water, sl. sol, hot ligroin. Gives di-benzylacetic acid on heating.

Ethyl ether (C.H. CH.) C(CO.Et). (250) at 40 mm. S.G. 30 - 1093. Thick yellow liquid. Formed by the action of benzyl chloride upon sodio-mulonic ether. By heating with alcoholic KOH it is converted into di-benzyl-acetie acid (Lellmann a. Schleich, B. 20, 439). Converted by treatment with alcoholic ammonia into CO Et.CH(C,H.).CO.NH. and (CO.NH.) CH.CH Ph, benzyl being split off (Bischoff a. Siebert, A. 239, 97).

TRI-BENZYL-MELAMINE

(C_H_CH_NH CN),. Formed spontaneously from benzyl-cynnamido by isomorio change (Strakosch, B. 5, 691). --B"311Cl. BENZYL-MERCAPTAN C:11,8 i.e.

C.H. CH. SH. Mol. w. 124. (195°). 8.0. 19 From benzyl chloride and KHS in alcohol (Marcker, A. 136, 75; 140 86) Pungent liquid with aliaceous odour.

Salts. -(C.H.S), Hg: needles. -O,H,SHgCl. (C.11.S),Pb

Benzoyl derivative C.H.CH.,SBz. [40°]. Colourless crystals (Otto a. Lüders, B. 13, 1285).

Ethylderivative C,H,SEt. (216°).

BENZYL-MESITYLENE C₁₄11, i.e. C₁11_C11_O₄11_Me₄, [36°]. (c. 302). V.D. 7·35. Prepared by boiling bonzyl chloride with mositylene in presence of Al_sCl_s (Louise, A. Ch. [6] 6, 176, C. R. 95, 11°3). Prisms; v. sol. alcohol, ether, and benzeno.

Ileactions. - 1. HI at 180° gives tolueno and mesitylene. -2. CrO, gives benzoyl-mesitylene. 3. IlNO, forms a tri-nitre dorivative, [185] and an acid [266].—4. Passage through a red-hot tubo forms two di methyl-anthracones, anthracere, and phenanthrene.

Di-beazyl-mesityleno . (Caii .CH.), Call Mo. [13:5], (355) at 120 mm. From lenzyl meaitylene, benzyl chloride, and Al₂Cl₄ (Louiso, A. Ch. [6] 6, 197). Minute prisms.
BENZYL-METHANE v. ETHYL-BENZENS.

Di-bearyl-mothago v. Di-Phentl-Propane. BENZYL MUSTARD OIL v. BRNZYL THIO-

BENZYL-METHYL-ACETIC ACID v. PHENYL-ISO-BUTYRIC ACID.

BENZYL METHYL ACETO ACETIC ACID v.

x x 2

BENZYL-DI-METHYL-AMINE C.H., N %.o. C4H, CH, NMe, Di-methyl-bensylamine. (184°). From benzyl chloride and alcoholic dimethylamine (Schotten, B. 15, 424; Jackson a. Wing, Am. 9, 78). Oil, miscible with alcohol and ether.

Salts. - *B'HCl. - *B'HNO2. - B'2H2PtCl2. $B'_2H_1FeCy_s$.— $B'_2H_2ZnCl_4$.

Methylo-chloride B'MeCl: white crystals, sol. water, v. sl. sol, Na, CO, Aq. -- (B'MeCl), PtCl, BENZYL-TETRA METHYL BENZENE

C.II., CIL, C. IIMo. [1:2:3:4:6]. [61°]. (c. 310°). From benzoyl-iso-durene and fuming III nt 250° (Essner a. Gossin, Bl. [2] 42, 170; A. Ch. [6] 1, 516).

BENZYL-METHYL-CARBINOL

C₄H₃,Cl1,Cl1(Ol1),Cl1, (215° i. V.). From benzyl methyl ketone and sodium-amalgam (Errera, G. 16, 315).

BENZYL METHYL-GLYOXIM C101112N2O, i.e. C.H., C11, C(NO11).C(NO11).C11, [181°]. Formed by the action of hydroxylamine hydrochloride on isouitroso-benzyl-acetone (Schramm, B. 16, 180). Small white needles. Sol. alcohol and other. Sublimable. Weak acid.

Di-acetyl-derivative C₁₀11,0(NOAc), — [80°], small white crystals (Schramm, B. 16, 2188î.

BENZYL METHYL-KETONE C,11,0 Call .. Cll .. CO.Cll .. Phenyl - acctone. (215°). S.G. 1010. Produced, together with nectono and di-benzyl-ketone, by distilling calcium acetute with calcium phenyl-acetate (Otto, J. pr. [2] 1, 144). Unites with NaIISO, heating with conc. 11,50, on the water-bath it is converted into the sulphonic acid Call (SOall).Cll2.CO.Cll2; but by heating quickly to a higher temperature it is split up into w toluene sulphonie acid C,11, CH, SO, H and

acetic acid (Krekeler, B. 19, 2625).
BENZYL-METHYL-KETONE \$\$LPHONIC ACID Call, (SO, 11).Cll, Cll, CO.CH, Formed by the action of funning sulphuric acid upon benzyl methyl ketone in the cold .- PbA',

(Krekeler, B. 19, 2625).

BENZYL-METHYL MALONIC ACID $\Gamma_0\Pi_0O_4$ i.e. $\Gamma_0\Pi_3\Pi_2\Pi_2$ CMe(CO₂H)₂₂ [135°], Colourless crystals. Prepared from the ether. On heating it gives CO, and phonyl-iso-butyrie

Discthyl-other A"Et., (300'), S.G. $\frac{190}{16}$ = 1 061. Prepared by the action of benzyl chloride on sodio-methyl-malouic ether or of methyl: iodide on sodio-henzyl-malonic ether (Conrad a. Bischoff, B. 13, 598; A. 204, 177).

BENZYL METHYL OXIDE C.H. CH.O.CH. (170'). From benzyl chloride and KOMe (Caliours, C. R. 80, 1317).

BENZYL METHYL PIPERIDINE C,H,N(C,H,)(CH,), (245°). Colourless fluid. Formed by dry distillation of the alkaline hydrate produced by the action of moist Ag.() on benzyl-piperidine-methylo-iodide.— (B HCl), PtCl, (Schotten, B. 15, 423).

DI - BENZYL - METHYL - (pseudo) - THIO -UREA C., H., N. S i.e. CH., S.C. NC., II. by heating di-benzyl-thio-urea with methyliodide At 100'. Oil. V. sol. alcohol and ether, insol. water.

Salts.—BHO: [122]; Cashy Soluble large rhombic four-sided tables.—BH-SO. [145°]; glistening needles; v. sol. water and alcohol.—B'HI: [99°]; octahedra; v. sol. warm alcohol, sl. sol. hot water.—B'H-Cl.PtOl.; sparingly soluble four sided prisms (Reimarus, B. 19, 2348).

(a) BENZYL NAPHTHALENE C, H, (c. BENZYL 150°). (c. 330°). S.G. LI 1.166. S. (alcohol) 3.3 at 78°; S. (ether) 50 at 15°. From naphthalene, benzyl chloride, and zincdust (Froté, C. R. 76, 639; Miquel, Bl. [2] 26, 2). Monoclinio prisms. Dilute HNO, produces phenyl (a) naphtliyl ketono [75°]

Sulphonic acid C, il 13 SO, H. - KA' aq: needles (from alcohol).

(B)-Benzyl-naphthalene C_i, H_{1i} (c. 315°). S.G. ? 1.176. S. (alcohol) 2.25 at 15°. Formed, together with the preceding, by heating naphthalene with benzyl chloride and Al,Cl. (Vincent a. Roux, Bl. [2] 40, 163). Monoclinic prisms (from alcohol); v. e. sol. benzene and chloroform. Nitrie acid produces phenyl (β) naplitly ketone [82°].

BENZYL-(a)-NAPHTHYLAMINE

C.11. CH. NII. C1. II.. [67°]. From naphthylamine and benzyl chloride (Froté a. Tommasi, Bl. [2] 20, 67).

BENZYL-NAPHTHYL-KETONE C,H140 i.e. C₁₁11, -CO -C11, C₁H₂, [57°]. Tables. Prepared by the action of Al₂Cl₃ on a mixture of phenylacetyl chloride and naphthalene. On reduction with HI it gives phenyl - naphthyl - ethane (Graebe a. Bungener, B. 12, 1078).

BENZYL . NAPHTHYL . METHANE v.

PHENYL-NAPHTHYL-ETHANE.

BENZYL-(B)-NAPHTHYL OXIDE C_s11_s.CH_s.O.C₁₀H_r, [99°]. White plates. Prepared by the action of benzyl chloride on sodium-(8) naphthol (Staedel, B. 14, 899; A. 217, 47).

BENZYL-NARCEINE v. NARCEINE,
BENZYL-NITRATE C. 11, C11, NO. 15

perhaps formed by the action of benzyl chloride on AgNO₄ (Brunner, B. 9, 1745).
BENZYL-NITRO-ARBUTIN v. p. 298.

BENZYL-NITRO-PHENYL-v. NITRO-PHENYL-BENZYL

BENZYL-ISO NITROSO MALONIC ACID C.11.C11.ON:C:(CO.11), From its other. The potassium salt on dry distillation gives KCN. potassium carbonate and benzyl alcohol.

Di-ethyl ether A"Et. Prepared by the action of benzyl chloride and solium ethylate on iso-mitroso-malonie ether (Courad a. Bischoff, B. 13, 599).

BENZYL - NITROSO - MALONYL - UREA v. Benzyl other of VIOLURIC ACID.

BENZYL OXALATE C, H₁, O, i.e. (C,11,CH₁), C,O₁, [81°]. From benzyl chloride and silver oxalate (Beilstein a. Kulilberg, A. 147. 341). Scales (from alcohol); may be distilled.

BENZYL OXAMATE C.H.NO.
i.e. NH.CO.CO.CH.Ph. [135°]. From

i.e. NH. CO.CO., CH.Ph. [135°]. From NH., CCl., CO., CH.Ph. and benzyl alcohol (Wal-

NH₂CCl₂CO₂CH₂H₁ in calculated a. Liebmann, B. 13, 507).

OXAMIDE C₁₆H₁₆N₂O₂ C.O.(MILCII,Ph). (216°). From oxalic ether and benzylamine; or by boiling benzylamine cyanide with HCl (Strakosch, B. 5, 694). Scales (from alcohol)

BENZYL-OXANTHRANOL v. OXANTHRANOS.

TETRA-RESTL OXY-AMMONIUM IODIDE | V. HYDROTYLANDER

BENZYL-OXY-BENZOIC ACID Oxy. BENETL-BENZOIC ACID

BENZYL-OXY-BUTYRIC Oxx. PHENYL-VALERIO ACID.

BENZYL-OXY-MALONIC ACID v. BENZYL-TARTRONIC ACID.

BENZYL-OXY-SULPHIDE DI-BENZYL SULPHOXIDE.

BENZYL . PHENANTHRENE Cz. 11,4 CH.Ph.C.,H. [1562]. From benzyl chloride, phenanthrene (q.v.) and zinc dust (Goldschuniedt, M. 2, 414). Needles (from benzene). CrO, gives

benzoie acid and phenauthraquinone.

p. BENZYL. PHENOL. C₁₃H₁₂O i.e.

C.H., C.H., C.H., OHI[1:4]. Oxy-di-phenyl-methane.

Mol. w. 184. [84°]. (325° 330°).

Formation.—1. From phenol, benzyl chlo-

ride and zinc dust (Paterno, G. 2, 2; 3, 121) .-2. From benzoyl anisol and III (Paterno, B. 5, 288; 6, 1202).—3. From PhOAc C.H. Cli,Cl alone or with Al,Cl, and saponification of the product (Porkin, jun., a. Hodgkinson, C. J. 37, 722, Reunic, C. J. 41, 228).-4. By heating phenol with benzyl alcohol and ZnCl, (Liebmann, B. 15, 152).-5. By diazotising pamido-di-pheuyl-methane and treating the produot with water (Basler, B. 16, 2719).

Properties.—Long needles or plates (from sleohol). Sol. KOHAq but not NHAq.

Reactions. - I. Distilling with P.O. benzene, phenol, and anthracene. - 2. By bromination and nitration, or by nitration and bromination it gives the same brome-nitro-derivative, when treated with HNO, in C.H.O. lorms C_sll₂(OH)(NO₂) Br [1:2:4:6]. Hence the brome-nitro-derivative C_sll₂(OH)(C₃H₂)(NO₄)Br is either [1:2:4:6] or [1:4:2:6]. But the oxidation of CaH, Cll, Call, OMe to CaH, CO.Call, OMe [1:4] shows that the arrangement is [1:2:1:6] 3. Phosphorus pentachloride forms (C13H11), PO4 [94°].-4. Chloro acctic acid and KOII convert benzyl-phenol into CH,Ph.C,H,O.CH,CO,H, [100°], while CH₄.CHCl.CO₂H and KOH give rise to CH₄Ph.C₂H₄.O.CHMe.CO₂H [102°] (Maxzara, G. 11, 437; 12, 262).

Methyl derivative C. H, CH, C, H, OMe. Benzyl-anisol. (305°). From anisol, benzyl chloride, and zine. Oxidised by alkaline KMnO. it gives the methyl derivative of p-henzoyl.

phenol.

Acetyl derivative Ph.CH₂.C_aH₄.OAc. (315°-320°). Benzoyl derivative PhCll, C. II. OBz.

Sulphonic acid C,H,.C,H,(80,H)(OH). Salts .- NH, A'aq: needles. - KA': feathery

orystals.—BaA',—C₁,11°,80°,Ba ac; minute crystals.—These salts are all spaningly soluble (Rennie, C. J. 41, 34; 49, 406). They, as well as the free acid, give a violet colour with Fe,Cl.

p-BENGYL-DIPHENYL C, H, i.e. C, H, c.e. C, H, CH, C, H, C, H, C, H, (S5°). (286°) at 100 mm. By heating benzyl chloride and diphonyl with zine dust at 100° two benzyl-diphenyls are produced together with traces of authracene. The phydrocarbon is less soluble and solidifies more easily than its isomeride (Goldschmiedt, M. 2, 433). Leaflets or needles, m. sol. alcohol, v. sol. benzene and ether. CrO, oxidises it, to phenyl-benzophenone and benzophenone p-car boxylic acid.

o-Bensyl-diphenyl (?). [54°]. (o. 285°) at 110 mm. Prepared as above. needles. CrO, oxidises it completely. Monoclinia

DI-benzyl-diphenyl OuH,(CH,Ph), From di-benzoyl-diphenyl and HI at 170° (Wolf, B. 14, 2032). Lamina (from alcohol).
BENZYL-PHENYL. v. Phenyl Benzyle.

BENZYL DIPHENYL AMINE P. DI THENYL-

u-DI-BENZYL . p. PHENYLENE - DIAMINE C.114(NH.).N(CH., C.114).1(1:4). Amido-di-beneylaniline. [90']. Obtained by reduction of p. nitro-di-benzyl-aniline with tin and HCl. Glistening colourless needles. V. sol. ether and hot alcohol, sl. sol. cold alcohol. With Fo.Cl. it gives a deep-red colouration, with Fc.Cl. and 11.S a blue insoluble up. By cone. HCl at 1700 it is completely resolved into benzyl chloride and p-phenylene-diamine.

Benzaldehyde compound

C. II. CII(OII).NII. C. II. N(CH., C. II.). [130°] Microcrystalline yellow pp. V. sol. henzene, sl. sol. ether, insol. alcohol (Matzudaira, B. 20, 1614).

BENZYL-PHOSPHINE C,11,1' i.e. C.H., CH., PH. (180°). From benzyl chloride, PH, l. and ZnO (Hofmann, B. 5, 100). Oil, volatile with steam. -B'IH: decomposed by water into its components.

Benzyl-triethyl phesphenium chlerlde PEt_a(CII,Ph)Cl. From benzylidene chloride, tri-cthyl-phosphine and alcohol (llofmann, A. Suppl. 1, 323).

Di henzyl phesphine (CH.Ph),Plf. [2050]. Prepared together with the preceding, and separated by stemm-distillation, not being volatile. Groups of needles (from alcohol); insol. acids.

Tri-henzyl-phesphine P(Cll Ph), Appears to be formed as a by-product in the action of benzyl chia, ide on PNa, (Letts a. Collie, Tr. E. 30, 181). Splits up on distillation into phosphorus, s-di-phenyl cthylene, dibenzyl, and toluene.

Tri-henzyl-phosphine exide (Cll.Ph).PO. [213']. Formed by heating l'II,I with henzylidene chloride at 130° and boiling the product with alcohol (Fleissner, B. 13, 1665). Formed also by the action of conc. Ba(OH), on tetra-henzyl-phosphonium acid sulphate; a weak solution of baryta giving P(C,II,),OH (Letts a Collie, Tr. E. 30, 181). Needles; insol. water, sol. nlcohol and er. $(C_{g_1}H_{g_1}P_C)_{_2}HgCl_{_2} - (C_{g_1}H_{g_1}P_C)_{_3}Fe_{_2}Cl_{_{g_2}}$ $(C_{g_1}H_{g_1}P_C)_{_3}CoCl_{_2} - (C_{g_2}H_{g_1}P_C)_{_3}PdCl_{_{g_2}}$ $(H_{g_1}P_C)_{_3}PdCl_{_{g_2}} - (C_{g_1}H_{g_1}P_C)_{_2}ZnI_{_{g_2}}$

(C_n,H_n,PO_j,PtCl_r, (C_n,H_n,PO_j,ZnI_r) (C_n,H_n,PO_j,PtCl_r, (C_n,H_n,PO_j,ZnI_r) (C_n,H_n,PO_j,Desphine sulptide (Cll_sPh),PS.

[206°]. Obtained by distilling the acid sulphate of tetra benzyl-phosphonium (Letts a. Collie). hong thin needles, m. sol. alcohol.

Tetra-henzyl-phosphenium hydroxide

P(C11), Oll. [over 200°]. Obtained hy adding Ba(O) to the sulphate. Rhomoledral plates; v. sol. water and alcohol; alkaline to litmus. Decomposed by heat into P(C, II,),O and toluone. Tetra-henzyl-phesphonium salts.

Chloride P(Cll.Ph),Cl. [224°]. benzyl coloride and l'Na, (1. a. C.). Crystallises from water with 2aq, and from chlorolorm with CHCl. Split up by heat into tri-benzyl-phos-phine, s-di-phenyl-ethylene, and HCl.

{P(CH,Ph),Cl},PtCl, Platinochloride

Sulphates. - {P(0,H,),} SO,: [220]. -

P(O,H.),SO,H.: [217]... Ozalate P(C,H.),C₂O,H.: needles. BENZYL-180.PHTHALIC ACID C₁,H₁,Q, i.e. C.H. CH2 C.H3 (CO.H)2. [243°]. From benzoxlbe plthalic acid by reduction with sodium-amalgam (Zincke, B. 9, 1765). Crystalline powder, v. sl. sol. water. BaA".—CaA"aq.

Benzyl-terephthalle acid C.H. .CII, C.II, (CO,H),... Obtained by reduction of benzoyl tereplithatic acid (Weber, J. 1878,

403). BaA". BENZYL PHTHALIMIDINE CHILON i.e. CH-CH1.C'H' CO NH Colourless

lates or scales. Formed by reduction of benzylidene-phthal-imidine by means of Hl.

Nitrosamine Call CII CII CII N(NO)

[93°]; yellow crystals, easily soluble in benzene, ligroin, and chloroform (Gabriel, B. 18, 1262). BENZYL-PHTHALIMIDE

C.H.G.O.:N.CH.C.H., Phthalyl-benzylamine. [116"]. Long needles. Obtained by heating potassium phthalimide with benzyl chleride at 170 '=180'. 11Cl at 200 'splits it up into phthalic acid and benzylamine (Gabriel, B. 20, 2227).

BENZYL-PIPERIDINE C, II, oN (C, II,). (245°). Colourless liquid. Insol. water. Prepared by the action of benzyl chloride on piperidine.

(B'HCl), PtCl,: sparingly soluble pp. Methylo-iodide B'Mel. [145°]. prisms. By moist AgO it gives an alkaline

prisms. by moise Argo it gives an altrameter hydrate which on dry-distillation yields methylbenzyl-piperidine (Schotten, B. 15, 423).

BENZYL PROPIONATE C₁₀II₁₂O₂ i.e.

C₂H₂CII₂O.CO.CII₂CII₂. (220°). S.G. (75 i - 0360. Decomposed by Na into sodium provionate and benzyl phenyl butyrate (Conrad a. Hodgkinson,

P - BENZYL - PYRROL C,H,:N.C,H,. (217° uncorr.). Colourless crystalline solid. Melts when touched with the hand. V. sol. alcohol and other, nearly insol. water (Ciamician a.

Silber, B. 20, 1369).

BENZYL-PYRRYLENE DI. METHYL. DI. KETONE C.H. (CO.CH.) NC.H., v-Benzyl-di. acetyl-pyrrol. [130]. Formed by heating v-benzyl pyrrol with Ac.O at 210. Colourless plates. Sol. ether and benzene, st. sol. water, nearly insol. petroleum-ether (Ciamician a. Silber, B. 20, 1370).

DENZYL QUINOLINIUM HYDROXIDE v. Benzylo-hydroxide of Quinning.

BENZYL ROSANILINES. From rosaniline and benzyl chloride (Dahl, D. P. J. 263, 393); ve ROSANILINE.

Methylo-iodide. From resaniline, MeI and MeOH (Holmann, B. 6, 263).

BENZYL SELENIDE (C.H., CH.), Sc. [46°]. From benzyl chloride and P.Se, (C. L. Jackson, A. 179, 1). Long needles or prisms (from alco-A. 113, 1). Long needees or prisms from accombin); faint odour, insol, water, v. sol, alcohol and ether. IINO, forms 'selenobenzyl nitrato' [88°]. ... {(C.H.), Se}, PtG|,
Benzyl discelentide (CH,Ph), Se, [90°].
Formed by boiling Na, Se, benzyl chloride, and alcohol for some hours (J.). Unotuous yellow

Unotuous yellow

scales (from alcohol). With MeI it forms (OH,Ph)SMe,I, [65°] from which may be obtained {(OH,Ph)SMe,Cl},PtCl, Conc. HNO, forms tolucne-exo-selinic acid, C,H,,CH,,SeO,H.

BENZYL SELENO.CYANIDE C.H., NSe i.e. C.H., CII., SeCy. [72°]. From benzyl chloride and potassium selenccyanide (Jackson, B. 8, 321). Prismatic needles with repulsive odour; insol. water, v. sol. het alochel. C.II.(NO.).CH. SeCy [123°]. BENZYL SELEN - UREA HNO, forms

CallioN, Se i.s. NII. CSe.NH.C11, Ph. [70°]. From benzylamine hydrochloride and alcoholic potassium seleno-cyanide (Spica, G. 7, 90). Sol. water, alcohol, and ether, gradually depositing Sc. Conc. HCl forms benzylamine, Se, and HCN.

u.di-benzyl-selenc-urea NH, CSc.N(CH,Ph), [150°]. From dibenzylamine hydroculoride and KSeCy. Thin prisms or needles; v. sol. hot water, alcohol, and ether. Conc. HCl forms Se,

CNH, and dibenzylamine.
TETRA-BKNZYL-SILICANE C28H28Si i.e. Si(CH₂Ph)₁. Silicon-t-tra-benzyl. [128]. (above 550). S.G. 22 1.078. Formed by the action of sodium upon a mixture of benzyl chloride and SiCl., with addition of a little acetic ether (Polis, B. 18, 1543; 19, 1023). Large monosymmetrical prisms, sol. ether, benzene, and

chloroform, sl. sol. alcohol. May be distilled. BENZYL SULPHIDE C., H., S i.e. (CH_Ph), S. [50°]. From benzyl chloride and alcohelic K₂S (Märcker, A. 136, 88). Thick trimetric tablets (from ether), a:b:c = 813:1: 515 (Forst, A. 178, 370; Bodewig). On distillation it gives s.di-plienyl-ethylene and its sulphide (Barbier, S. C. R. 78, 1772), tolucue, benzyl mercaptan, s-di-phenyl-acctylene sulphide SC,Pl₂, and thionessal C, Il, S. McI forms SMe₂1, benzyl iodide and (GH,Ph)SMe₂1; the latter gives riso to the compound {(CH,Ph)SMe,Cl},PtCl. Exhyl icdide at 100° forms similarly (C.H.) SEt.I whence {(C,II,)SEL,Cl},PtCl, (Sohöller, B. 7, 1271; of. Cahours, A. Ch. [5] 10, 21).
DI-BENZYL DI. SULPHIDE (C,4H,0H,),S,

Sulphobenzol. [70°]

Formation. -1. By the action of an alcoholic solution of KIIS or K,S on beuzylidone dichloride. -2. By the action of alcoholic KHS on (a) thicbenzoic aldehyde (Klinger, B. 15, 861).-3. By the oxidation of benzyl mercaptan by air or bromine (Märcker, A. 140, 86).-4. By the action of K,S, on benzyl eliforide in alcohol (M.).

Properties.—White plates. Gives a crystal-line pp. (C₁₄H₂₁S₂AgNO₂) with an alcoholic solution of AgNO₃.

BENZYL-SULPHINIC ACID v. Toluene Rio-SULPHINIC ACID.

BENZYL SULPHOCYANIDE

C.H.,CH.,S.CN. (41°] (B.); [38°] (H.); (c. 233°) (B.); (250°) (H.). From benzyl chloride and alcoholic potassium sulphosyanide (Henry, B. 2, 636; Barbaglia, B. 5, 689). Prisms (trom alcohol), insol. water; pungent smell. Combines with HBr, forming a compound decomposed by water. Fuming nitric acid forms C.H.(NO.).CHL.S.Cy.
DI-BENZYL-SULPHONE C., HI, SO. i.e.

(CH.Ph.),SO., [150].

Fornation.—1. Together with C,H,CH,SO,K
by the action of K,SO, on benzyl chloride
(Vogt a. Henninger, A. 165, 375).—2. By exida-

tion of di-benzyl sulphoxide with KMnO, and HOAc (Otto a. Luders, B. 13, 1284).-3. By the action of benzyl chloride on sodium benzene-sulphinate. 4. By oxidation of di-benzyl sulphide.

Proporties. Small needles. Insol. water, sol. alcohol, benzone, and acetic acid. By oxidising agents it is readily oxidised to benzoic and sulphuric acids (Otto, B. 13, 1277).

BENZYL-SULPHONIC ACID v. TOLUENE-620-SULPHONIC ACID.

DIBENZYL-SULPHONIC ACID v.

PHENYL-RYHANE SULPHONIC ACID. DI-BENZYL SULPHOXIDE Callaso i.e. (CH.Ph.)SO. [133°]. From di-benzyl sulphide and cold HNO, (S.C. I'3) (Märcker, A. 136, 89; Otto a. Ludwig, B. 13, 1284). Lamina (from water or alcohol)

BENZYL SULPHUROUS ACID v. TOLUENE

SULPHONIC ACID

BENZYL-TARTRONIC ACID C 1011,00, i.c. C, H, CH, C(OH); (CO H), [143°], Formed simultaneously with cinnamic acid by the action of KOH on benzyl chloro malonic ether. On heating it forms \$-phenyl-a-oxy-propionio acid (plienyl lactic acid [98°]) (Conrad, B. 13, 2160; A. 209, 215).

BENZYL-TEREPHTHALIC ACID v. BENZYL-

PHTHALIC ACID.

BENZYL-THIO CARBAMIDINE C.II. N.S. i.e. NII:C(NII).S.CII.Ph. Cyanamide benzylmercaptan. [72]. From benzyl chloride and thio-urea (Berntlisen a. Klinger, B. 12, 575). Slender needles, m. sel. water; decomposed by heat into benzyl mercaptan and di-eyan-di-amide. -B'HCl. [1683].--B',H,PtCl. BENZYL-TH10.CARBIMIDE

C.H., CH., N:CS. Benzyl mustard oil. Benzylamine is dissolved in CS, and the product boiled with alcohol and HgCl2 (Hofmann, Z. [2] 4, 690; B. 1, 201). Oil, swelling like water-cress.

BENZYL-THIO GLYCOLLIC ACID v. THIO.

GLYCOLIAC ACID

BENZYL-THIO-UREA C. II, N.S i.e. (CH_Ph)NH.CS.NH, [1019]. From potassium sulphocyanide and benzylamine hydrochlorida (Paterno a. Spica, G. 5, 388; B. 9, 81). Sol.

water and alcohol. Benzoyl derivative C.H.NH.CS.NHBz. [145']. From benzoyl sulphocyanide and benzyl-

amine (Miquel, A. Ch. [5] 11, 313).

s.Di . benzyl . thio . urea (Ph.CH, NH) CS. [114°]. From alcoholic benzylamine and CS, (Strakoseli, B. 5, 692). Four sided plates, insol. water, sol. alcohol and other. Converted by HgO into di-benzyl-urea. Alkyl iodides give the following derivatives:

PhCH, NILCS NMe.CH, Ph. C,H,NH.CS.NMeC,H,J,H,PtCl,... C,H,NH.CS.NMeC,H,B (99)... PhCH,NH.CS.NEt.CH,Ph... (C,H,NH.CS.NEtC,H,),H,PtCl,--C,H,NH.CS.NEtC,11,H1 [93°]. C.H.NH.CS.NEtC.H.H.SO.

PhCH₂NH.CS.NPr.CH₂Ph.— PhCH₂NH.CS.N(C₃H₁₁).CH₂Ph (Reimarus, B. 19, 2348).

u-Di-bensyl-thio-nrea (PhCH₂), N.CS.NH₂. 7°]. From potassinm sulphocyanide and dibenzylamine hydrochloride (P. a. S.). Largo needles, m. sol. water.

BENZYL-THYMOL C, H,O &. C.H.MePr(CH.Ph)(OH). (255°) at 8 mm. Formod, together with di-benzyl-thymol by C.H.MePr(CH,Ph)(OH). heating benzyl chloride with thymol and sincdust (Mazzara, G. 11, 346). Oil, insol. aqueona alkalis, sol. alcohol and ether. Fe₂Cl₂ gives a red colour on heating.

Acetyl derivative C, HnAco. (245°) at 8 ftm.

Di-benzyl thymol C₂₄H₂₆O i.e. C₂HMePr(CH.Ph)₂OH. [76°]. Prepared as above. Silky lamine, sol. ether and HOAc, insol. water and aqueous alkalis. Fe Cl. gives a red colour on heating.

Acetyl derivative C, H, Aco. [0.84], Methyl derivative C, H, MeO. [909] Benzoyl derivativo CallaBzO. [c. 78°]. BENZYL . TOLUENE v. PHENYL - TOLYL-

METHANE

DI.BENZYL-TOLUENE C., II., i.e. CH., C., II., i.e. CH., C., II., CH., Ph.). A product of the action of benzyl chloride on tolucue in presence of zine-dust (Weber a. Zincke, B. 7, 1154).

BENZYL p-TOLUIDINE PhCH_NH.C.H.Me. (313°). From benzylidene-p-toluidine (Kohler,

A. 241, 3590.

Di-benzyl-p-toluidine C_nH_nN i.e. (Ph.CH_s)_sN.C_sH_sMe. [55°]. From henzyl chloride and p-tolnidine (Cunnizzaro, A. Suppl. 4, 80). Slender needles, m. sol, cold alcohol. Weak base.

BENZYL-TOLYL. P. TOLYL-BENZYL-.

BENZYL - TOLYL - METHANE v. PHENYL-TOLYL-UTHANK.

BENZYL-TOLYL OXIDE v. BENZYL ALCOHOL-

BENZYL-UREA C. II. N.O. i.e. NH. CO.NH. CH. Ph. [147°]. Formed, together with di-benzyl-urea, by beating benzyl oliloride with potassium cyanate in alcoholic solutions (Cannizzaro, G. 2, 41). Also from bonzyl eyanate and alce! olic NII, (Letts, C. J. 25, 448) or from benzylamine chloride and potassimu cyanate (Paterno a. Spica, G. 5, 388; B. 9, 81). needles (from alcohol); m. sol. water. A it splits up into NII, and s-di-benzyl urea.

s Di-benzyl-urea (CII.Ph.NII).CO. [167°]. Formation.-1. From benzyl chloride and KNCO or urea .- 2. From benzyl urea by heating. 3. By heating benzyl alcohol with urea mitrate (Campisi a. Amato, G. I, 39; B. 4, 412). 4. From s di benzyl thio urea, HgO, and alcohol (Strakosch, 3, 5, 692).

Properties. Needles, insol. water, v. sol. alcohol. Weak base.

u Di benzyl nrea (CH, Ph)2N.CO.NII2. [125°] From di benzyl-amine nydrochloride and KNCO (Paterno a. Spice, G. 5, 388; B. 9, 81). Thick prisms; sl. sol. cold water.

BENZYL-URETHANE v. BENZYL-CARBAMIC

BENZYL m. XYLENE C, II, i.c. C.H.CH., C.H., Mc, Phenyl xylyl methane (296° i. V.). From mxylene, benzyl chloride, and zine-dust or Cu (Zincke, B. 5, 799; 9, 1761). Oxidation gives benzoyl iso phthalio acid. Ap pears also to be formed by reducing plienyl xylyl ketone with HI (Söllscher, B. 15, 1682).

Benzyl-p-xylene. (295°). From p-xylene, benzyl chloride and zinc-dust (Z.).

BENZYL XYLYL KETONE

 C_aH_1 , CH_2 , $CO.C_aH_2$ (CH_2)₂. [1:2:4]. Dimathuldecrybensom (above 250°). Finid. Formed by the action of Al₂Ol₂ on a mixture of m-xylene and phenyl-acetyl-o'nloride. On oxidation it gives di-methyl-benzoic acid (Söllscher, B. 15, 1681).

BERBAMINE C, H., O, N. [156°]. Occurs in the root of Berberis vulgaris, together with berberine, oxy-acanthine, and at least one other alkaleid. Small plates, containing 2uq. Easily sol, other. The hydrochloride forms entill plates, the nitrate needles. -- B', II, Cl, PtCl, 5 or 6 aq: yellew crystalline pp., el. eol. celd water (Heese, B. 19, 3193).

BERBERIC ACID C.H.O.aq. An acid fermed by fusing berberine with KOII (Hlasiwetz a. Gilm, J. 1864, 407). Needles; v. sol. alcohol and other, m. sol. water. Fe Cl, gives a green colour turned red by ammenium tartrate. Redutes hot Felding's solution und silver solution.

BERBERINE C., II., NO, 41 aq. [120]. S. 22 at 21°. S. (alcohel) 1 ut 15°. Occurs in the root of the barberry, Berberis vulgaris, together with oxy-acanthine (q. v.), berbamino and another alkaloid (Buchner, A. 21, 228; Hesse, B. 19, 3190). Occurs also in Colomboroot (Cocculus pulmatus) (Floitmann, A. 59, 160; Bödeker, A. 66, 381; 69, 40); in Menispermum fenestratum (Perrins, G. J. 15, 339); in Abeccouta bark from Coelocline polycarpa (Stenhouse, Ph. 14, 455; C. J. 20, 187; Daniel, A. 105, 360); in Launtice thalietroides (Mayer, J. Ph. [3], 46, 496); in Xanthorylon clava Herculis (Chevallier a. Pelletnn, Berz. J. 7, 266; Perrius, A. Suppl. 2, 171); in bark of Geoffroyea inermis (Gastell, J. 1866, 180); in Coptis trifolia (Groes, J. 1871, 914); and in the root of Evodia glanca (Martin, Ph. [3] 13, 337).

Preparation .- 1. The finely powdered root of Hydrastis canadensis is extracted with alceliol; H.SO, is added to the cooled extract, and the pp. decomposed by NII₃. The operation is repented a second time (Lloyd, Ph. [5] 10, 125; of. Merril, Am. J. Pharm. 35, 97; Procter, C. N. 9, 112). 2. Barberry root is exhausted with boiling water; the extract ovaporated, and treated with 93 p.o. alcohol. The berberine is purified by crystallisation from water or ulcohol

Properties. Silky yellow needles; tastes bliter; sl. sol. cold water and alcohol. insol. ether. Turned brown by anumonia. On adding iodine in potassium iodide to a solution of berberine hydrochloride the periodide is ppd. It crystallises from alcohol in red needles, but on adding water, green plates separate.

Bactions.—1. Zipc and dilute acids form hydro-berberine. 2. Potash fusion produces two acids, C.H.O., and C.H.O., (Hlasiwetz a. Gilm, J. 1861, 406). —3. Nitric acid oxidises it to berberonic acid.—4. KMpO, in presence of KHO forms hemipic acid, [162] (E. Schmidt a. Schilbach, Jr. Ph. [3] 25, 161).

Salts.—Gleitmann 4. 59, 160). Hoper 4.

Salts. (Fleitmann, A. 59, 160; Henry, A. 115, 132; Perrins, C. J. 15, 339; Illasiwetz, A. Suppl. 2, 191). — B'HCl: elender yellow needles. -- B HCl laq. -- B'HCl 2aq. S.G. 194 1.397 (Clarke, Am. 2, 175). - B'211.11gCl. B'H.Cl.; small needles. S.G. V 1758 (C.).
B'HAuCl.; maroon - coloured needles.— Methylo-iodide B Mol: needles (Bern-heimer, G. 13, 315).

Ethylo iodide B'Et1: needles. Hydroberberine C, II , NO. Obtnined by reducing berberine in acid colution with zino (Illasiwetz a. Gilm, A. Suppl. 2, 191). Granules er needlee (from alcohol). Reconverted into berberine by HNO.

Snlts. -B'HCl. - B'2H.PtCl. - B'2H2SO. -B'H.SO. - B'.(H.SO.), 4aq (?). - B'HI. B'HNŌ,.

Methylo-iodide B'Me1: trimetric crystals, a:b:c=1.033:1:1.789. -B'McOll (Bernlicimer, G. 13, 312).

Ethylo-iodide B'EtI: prisms.

BERBERONIC ACID v. PYRIDINE-TRI-CAR-HOXYLIO ACID.

BERGAMOT, OIL OF. An arematic escential oil expressed from the rind of an orunge, Citrus bergamia. 1ts S.G. is 87. It contnine a etearoptene, a terpene, and a terpene hydrate (?). By rectilication a liquid (183°) may be got, which absorbs HCl (Soubeiran a. Capitaine, J. Ph. 26, 68, 509). The etenroptene (Bergaptene) is deposited after long keeping. It is eolid, [206'], but volatilises without decomposition. It may be (C,11,0)x (Mulder, A, 31, 70; Olmie, A. 31, 346).

BERGENITE C, II, 0, aq. [130°]. [a]0 = -51' 36'. Obtained from Siberian saxitrage (Bergenia siberica) by extracting with het water, ppg.the tannins with lend acctate, and evaporating to crystallisation (Merelli, C. R. 93, 616). Trimetric prisms. Tustes bitter. V. el. eol. gold alcohol and water.

Acetyl derivative CaH, AcOs: amorphoue,

v. col. water, alcohol, and ether,

Triacetyl derivative C, II, Ac, O, Penta-acetyl derivative Callaco BERLIN BLUE = Prussian Blue v. Ferrocyanide of iron under Cyanides.

BERONIC ACID v. PYRIDINE DI-CARBOXYLIC

BERYLLIUM. Be. (Glucinum.) At. w. 9-08. Mel. w. unknown. S.G. 30 (after compression) 1-85 (llumpidge, Pr. 39, 1). S.H. (100°) 4702; (200°) 540; (100°) 6172; (500°) 6206 (Humpidge, Pr. 39, 1). S.H. (20°) 397; (73°) 448; (157°) 5519; (257°) 581 (Meyer's calculation, B. 13, 1780, from data of Nilson and Pettersson who worked with metal containing about 95 p.c. Be; B, 13, 1151). S.V.S. 4.92. Crystallises in hexagenal, holehedral, forms; a.c. = 1:1:5801 (Regger a. Flink, B. 17, 819).

Occurrence. Only in combination; in beryl

(3BcO.Al.O.,6SiO.) and come other silicates, also in chrysoberyl Al.O.,BeO. Beryllium exido was recognised as a distinct body in 1797 by Vanquehn, the metal was obtained by Wöhler in 1827, but approximately pure beryllium was not prepared until 1885, in which year Hum. pidge obtained specimens containing 99.2 p.c.

Be, 1 Fe, and 7 BeO.

Preparation.—Wöhler (P. 13, 577) obtained an impure metal by the action of K on fused Be(I. Debray (O. R. 88, 784) obtained purer specimens by using Na and a special form of apparatus. Nilson a. Pettersson, by decomposing BeCl, by Na in closed iron crucibles heated in a wind-inrnace (B. 11, 381), and sifting the crystals through Pt gauze (B. 13, 1455), obtained epecimene containing 94.4 p.c. Be (BeO = 4.89, Fe = .70). Humpidge (Pr. 38, 188; 89, 1) purified BeO by solution in (NH,) CO, Aq and decomposing the solution by steam; he mixed the BeO thus obtained with pure charcoal and starch paste and heated in Cl in a glazed porcelain tube; the BeCl, thus obtained was placed in an iron boat, and this in an iron tube surrounded by another tube of hard glass; another iron boat contained Au, the BeCl, was heated in a stream of II, and the BeCl, was then vaporised (in H) over the molten Na. crystals of Be were washed in dilute NaOHAq, to remove BeO, then in water, and dried.

Properties. Steel-coloured, hard, hexagonal, holohedral (Brögger a. Flink, B. 17, 849) crystals. Unchanged in ordinary air; scarcely changed by heating in air. Scarcely acted on by O or S at red heat, but burns in Cl to BeCl, (Nilson a. Pettersson, B. 11, 381). Burns in O-H flame (Humpidge, T. 171, 601). Dissolves slewly in acids, also in aqueous alkalis, with evolution of H. Many of the properties ascribed to Bo by Wöhler (P. 13, 577), and Debray (A. Ch. (3) 41, 5), were the results of experiments with very impure material. Emission-spectrum characterised by the lines 3320:5, 2649:4, 2493:2, and 2177.7, of which 3320.5 is the most prominent (Hartley, C. J. 43, 316). The atomic weight of Be has been determined (i) by analyses, and determinations of V.D., of BeCl, and BeBr₂; (ii) by determinations of S.H. of Be; (iii) by analyses of various compounds, especially recently of the pure crystallised sulphate by Nilson a. Pettersson (B. 13, 1 51) [for older analyses v. Berzelius, P. 8, 187; Awdejew, P. 56, 101; Klatzo, J. pr. 106, 227]; (iv) by the application of the periodic law. There has been much investigation and disoussion concerning the value to be given to the atomic weight of Be; some chemists insisted that 13.65 is the true value, and that the oxide is Re₂O₂. The determinations of the V.D. of BeCl₁ and BeBr₂, the eareful measurement of the S.H. of almost pure Be, and the consideration of the physical and chemical relations of Be and its compounds to other elements carried. out on the lines suggested by the periodic law, have finally established the value 9:08 9:1 for the atomic weight of Be. The S.H. of Be in-creasee rather rapidly as the temperature increases, and approaches a constant value, equal to about 62, between 400° and 500° (Hum, pidgo, Pr. 39, 1). The relation between S.H. and temperature is expressed by the empirical formula K, = 3756 + 00106t - 00100111t' (Humpidge, Pr. 38, 188). The product of S.H. into At. w. $(.62 \times 9.1 = 5.64)$ is lower than the mean value of this quantity for the solid elements, but is nearly the same as that obtained for B, C, and Si. The atom of Be is divalent in the gaseous molecules BeCl, and BeBr,; these are the only compounds of Be at present known in the gaseons etate. A comparison of the spec-

hand, and of Mg. Ca. Ba. Sr. on the other, show that Be is rather to be classed with the latter than with the former elements; the value 9-1 for the atomic weight of Be is thus confirmed (v. Hartley, C. N. 48, 195). Beryllium is a markedly positive, or metallie, element; it does not exhibit allotropy. It is eliemically related to Mg, Ca, Sr, und Ba, in much the same way that Li is related to Na, K, Ce, and Rb. BcO resembles MgO, but is distinctly less hasie, e.g. it dissolves in KOHAq, and does not combine directly with H₂O; compare also [BeO 11', H²SO Aq] = 16,096, with [MgO H; H SO Aq] = 31,216. Analogies also exist between Be and Al; e.g. the existence of many basic salts; Be, however, does not form an alum; BcCl₂ does not combine with NaCl and KCl as Al Cl₄ does. In dilute acid solutions Be is electronegative to Mg but positive to Al; in cauetic alkali solutions the electrochemical order is + Al, Mg, Be -(Humpidge, Tr. 174, 601). Be forms only one eeries of compounds, BeCl., BeSO, Be2NO, &c.; it exhibits a marked tendency to form basic salts, e.g., BeSO, BeO.3H₂O, BeCO, 3BeO.5H₂O, &c. Brauner (B. 14, 53) enums up the chemical relatione of Bo in the three statements
(1) Li:Be - Be:B. (2) Li:Na - Be:Mg - B:Al.
(3) Li:Mg - Be:Al - B:Si. The chemical relations of Be will be more fully discussed in the art. Maonesicm merals; v. also the remarks on Group 11. in art. CLASSIFICATION. The following are the principal papers bearing on the At. w. of Be: Reynolds, P. M. [5] 3, 38; ibid. Pr. 35, 218; Humpidge, Pr. 35, 358; 38, 188; 39, 1; Hartley, C. J. 13, 316; ibid. Pr. 36, 462; Carnelley, Pr. 29, 190; ibid. B. 17, 1357; Branner, B. 11, 872; 14, 53; Meyer, B. 11, 577; 13, 1780; Nilson a. Pettersson, B. 11, 381, 906; 13, 1451, 2035; 17, 987.

Reactions.—As most of the reactions eaid to characteric Be have been obtained by experimenting with material far from pure, that following statements must be accepted as provisional only.—I. Hydrochloric acid, whether gaseous or aqueous, reacts readily to produce 1scl..—2. Sulphuric acid dissolves 1se, forming BeSO Aq.—3. Nitric acid acts very slowly even when hot and concentrated.—4. Be dissolves in warm KOHAq or NaOHAq.

Combinations.—1. Be combines readily with Cl. Br. and I to form BeCl., BeBr., and BeG. respectively.—2. At also combines very readily with Si; and, according to the observations of Wöhler, made, however, with very impure material, with P. Se, and S. later experiments seem to show that Be and S do not combine when heated together (Nilson a. Pettersson, B. Il, 331).—3. An alloy of Be and Fo was described by Stromeyer as a white solid, less malleable than iron, obtained by strongly heating, BeO with Fe and charcoal.

Detection, -1. Caustic potash or soda pos, BeoILO soluble in excess, but reppd. on diluting and boiling.—2. Amnonium carbonate pos, the carbonate easily soluble in excess; this reaction distinguishes salts of Be from salts of Al.—3. Be salts give no colour when heated with Co.2NO.

the only compounds of Be at present known in the gaseons etate. A comparison of the spectrum of Be with those of In and Al on the one BeO is separated from Al.O. by the action of

(NH_{...})₂CO₂ (v. Hofmeister, J. pr. 76, 1); or by converting the alumina into potash alum (v. Scheffer, A. 109, 144).

Beryllium, Alloys of. Little or actling is known; v. Beryllium, Combinations, No. 3.

Beryllium, Bromide of. Bellr., Mol. w. 168-6. (abt. 600°) (Carmelley, B. 17, 1357); sublimes readily at 150°. V. D. 90 (Humpidgo, Pr. 38, 188).

Preparation. -1. By heating Be in Br vapour (Wöhler, P. 13, 577). -2. By heating BeO mixed with chargoal and made into u pasto with starch, in dry Br (Humpidge, T. 174, 601). Crystals of hydrated BeBr, are obtained by dissolving freshly 1pd. BeO.H.O in HBrAq, and evaporating (Berthemot, A. Ch. [2] 44, 391).

Properties. Long white needles; very deliquescent. Heated in air, partly sublines, and

is partly decomposed into BeO and Br.

Beryllium, Chlorido of. BeCl. Mol. w. 80, at low temps. 169 | BeCl. [about 600] but sublimes considerably lower (Carnelley, C. J. 37, 26; v. also ibid. B. 17, 1357). V. D. (685° to 1500') 40°96 (mean of 4); V. D. (520') 60°4 (Nilson u. Pettersson, B. 17, 987; J. pr. [2] 31, 1).

Preparation. - 1. By heating Bo in Cl. - 2. By heating Bo O and C in Cl, and subliming in HCl. - 3. By heating Bo in perfectly dry HCl, and subliming in the same (Nilson u. Pettersson, B. 17, 987).

Properties.—Snow-white crystalline mass; easily fused and volatilised; melts to a colour-less refractive liquid; may be sublimed unchanged in pure dry N or CO₂; but is easily decomposed into lic0 and Cl if a little air is present (Nilson a. Pettersson, B. 17, 987.—Is practically a non-conductor of electricity (Nilson a. Pettersson, B. 11, 382; Humpidge, T. 171, 601).

Reactions.—Dissolves in water visit production of much heat; on evaporation, an expedience 1, if the solution is placed over 11-SO, under a bellipar, crystals of BeCL-H1O separate out (Awdsjow, T, 56, 101; Atterberg, RL, $\{2, 21, 358\}$.

Constantains. 1. With chlorides of various keary metals, to form double compounds; (specially fact), 31 from double fact), 32 from double fact), 33 from double fact), 34 from double fact), 35 from fact), 35 from fact), 36 from fact), 36 from fact), 37 from fac

Beryllium, Fluoride of. Hydrated Bet) dissolves easily in HFAq; on evaporation, a transparent gun-like, mass sobtained which becomes opaque at 100°, but remains soluble in water; this is probably BoF, (Berzelius). Tho doublo compounds BeF_2KF, and BeF_KF, are doscribed by Marignae (A. Ch. [4] 30, 45), the former obtained by adding much KFAq, the latter by adding little KFAq, to a solution of BeO.H O in HFAq, and evaporating. The compounds 2NaF_BeF_q, NaF_BeF_q, and 2NH_F_BeF_q are also described by Berzelius.

Beryllium, Hydrated exide of, v. Beryllium, avdroumes or.

Beryllium, Hydrexides of. A compound of Be, H, and O, agreeing in composition with the formula BsO,H, (=BeO.H,O) is obtained as a [2] 26, 227) says that the pp. by KOHAq has the composition BcO.H.O only when heated to 150°-180°: he describes a gelatinous hydrate. BeO.H.O. obtained by the action of NHAq on BeSO, Aq, washing with cold water and drying in dry air at 15°-20°; the composition of this hydrate is constant up to 200°. The compound Bett. Il, is not re-formed by the action of water on BeO (for more details of this action v. Van Benunelen, l.c.); it seems better to regard it us a hydrated oxide, BeO.H.O, than as a hydroxids Be(OII). Other hydrates of BeO are said to be obtained by drying the pp. by NII,Aq over H.SO, or merely in air, but the composition of these bodies is variably (v. Schalfgotsch, P. 50, 183; Atterberg, B. 7, 473; Van Bemmelen, J. pr. [2] 26, 227). The gelatinous BeO.H.O is ensily soluble in ucids, ulso in NaOllAq und KOHAq, and in (NII.) CO.Aq. By boiling the solution in KOHAq a ppt. of 3BeO.411.0 (Atterberg, B. 7, 173) is obtained (but v. Van Bemmelea, J. pr. [2] 26, 227). BeO.H., acts as a hasic hydroxide towards acids; Thomson gives the following values for its heats of neutralisation, solid ReO₂H, being used in each case (Th. 1, 363); [BeO IF, IFSO Aq] = 16,096; [BeO IF, 211CAq] = 13,644. These values are much less than those for the alkaline earth hydroxides (about 31,000 for H.SO, Aq).

Beryllium, Iodide of. Bel. Mol. w. nn-known, as V.D. has not been determined. Described as colourless needles obtained by testing together Be and I (Wöhler, P. 13, 577; Debray, A. Ch. [3] 41, 5). Easily decomposed by action of hot air into BeO and I (De-

ray, Lc.).

Boryllium, Oxide of. BeO. Mol. w, unknown. S.G. 3:016. S.H. (0° to 100°) :2471 (Nilson a. Pettersson, B. I.I., 1454). S.V.S. 8:3.

Preparation, "Bery is fined with NaKCO, in graphite crucibles in a wind-furnace; the fused mass is leated for some time with excess of 11.80, Aq, water is added and SiO, removed by filtration, the liquid is svaporated until a crust begins to form and is then allowed to stand for 21 hours or more; potash alum and K.SO, separate out; the mother liquor is again evaporated and a second crop of alum crystals is obtained and removed; the mother liquor is now poured into a-warm cone. aqueous solution of ammonium earbounde, the pp. (of Al₂O₂ &c) is desested in contact with the liquid for four or five days and then filtered, the insoluble portion is again heated with (NH₂)₂CO₂Aq and the liquid is filtered off. The solution in (NI₁)₂CO₄Aq contains BeO, free, or almost free, from Al₂O₄, CaO, Fc₂O₃ &c.; the BeO may be obtained by boiling the solution (Nilson a. Petlersson, B. 11, 383), or by activating, boiling off CO₂, and ppg. by NH₂Aq (Ilofmeistor, J. pr. 76, 3); in either ease, the ppd. BeO.zIf₂O₃Aq and then reppd. There are various other ways of

preparing BeO.zH.O from beryl (v. especially Joy, J. pr. 92, 232; Scheffer, A. 109, 146; Berzslius, P. 8, 187; Debray, A. Ch. [8] 44, 15).

Properties .- A white, loose, infusible powder; insoluble in, and unacted on by, water; soluble, in acids and in molten KOH. According to Ebelmen (A. 80, 213) BoO is obtained in hexagonal crystals (a:c=1:1:587) by cooling a solution of the oxide in molten borio acid; Debray obtained similar crystals of BeO by etrongly licating ammonium-beryllium oarbonato (A. Ch. [3] 44, 15). H. Rose described crystale of BeO obtained by heating the ordinary oxide in a porcelain oven (Ph. C. 1848, 486); S.G. of thsee crystale = 3.02.

Reactions and Combinations .- 1. With most acids to form salts, e.g. BeSO, Bc.2NO, &c.; tho oxido becomes less easily soluble in acids by heating .-- 2. Decomposes molten potassium carbonate with evolution of CO,; on addition of water BeO remains dissolved in the KOHAq. -3. Does not combine directly with water, but varions hydrates, of which BeO.11.0 is the most important, are obtained by the action of , NII, Aq on solutione of Bo ealts (v. BERYLLIUM,

Hypnoxibus or).

Beryllium, Oxychleride of,

Bo₂OCl₂=BeCl₂BcO. Said to be formed by evaporating an aqueous colution of BeCl...

Beryllium, Phosphide of. Described by Wöhler as a greyish powder obtained by heating Bo in vapour of P; existence very deubtful as Wöhler's Be was very impure.

Beryllium, Salts of. Salts obtained by roplacing II of acids by Be: they are generally obtained by the action of aqueoue acids on BeO.H.O. Most of the commonor salts—salphate, nitrate, oxalate, chlorido-are solublo in water and have a sweetish taste; the carbonate and phosphate are insoluble in water. When heated, the salte of Be are completely uccomposed, except the acid be non-volatile. The chief salts of Bo are the following (they are described under the headings Carnonares, Na-TRATES, &c.) carbonates, chromates, molyblates, nitrates, perchlorate, periodates, phosphates, selenate, selenites, silicates, sulphates. The following salts probably exist, but few if any definite facts concerning them are known :bromate, chlorate, hypophosphite, iodate, phosphite, tungstate, vanadates.

Beryllium, Selsnids of. Exictonce very doubtful.

Bsryllinm, Silicids of. Be readily combines with Si; when Be is prepared in porcelain vessels a portion of the SiO, is reduced and as much ae 20 p.o. Si may combine with the Be to form a hard, brittle mass. It is doubtful whether any definite compound of Bo and Si has been obtained.

Beryllium, Sulphide of. Described by Wöhler (P. 13, 577) as a greyish fused mass, which evolves H2S by action of acids; obtained by heating Be in S vapour. But oxistence is extremely doubtful; according to Freiny (A. Ch. [3] 38, 326) no snlphide of Bs is produced by heating BeO in S, or CS, vapour. According to Nilson a. Pettersson (B. 11, 381), Bs and S lo not combine when heated together.

M. M. P. M.

BERYLLIUM, ORGANIC DERIVATIVES OF, Beryllium ethide BeEt, (185°-188°). From HgEt, and Be at 130°. Takes fire in air (Cahours, J. 1873, 520).

Beryllium propide BePr. Does not take fire in air.

BETA .- Compounds beginning with beta- or het- are described as \$ compounds under the word to which this prefix has been added.

BETAINE $C_3H_HNO_4$ i.e. Me_3N , $< \frac{CH}{O} > CO$.

Internal anhydride of the methylo-hydroxide of di-methyl amido-acetic acid. 8.16 at 25°. In the hydrated condition C.H, NO, aq, it may be represented by the formula Me, N(OII).CII, CO, II.

Occurrence. -1. In the juice of beet root (Beta vulgaris), and in beet-root molasses (Scheibler, Z. 6, 505; B. 3, 155; Liebreich, Z. 6, 506; B. 3, 161). The unripe root contains 25 p.o.; the ripe root only I p.o. The betaine ie not present in the root as such, but is liberated by treatment with HCl or baryta. 2. In man-gold wurzel (Scheibler, Z. [2] 5, 539). 3. In cotton seed (Ritthausen, J. pr. [2] 30, 32). -4. In the leaves and branches of Lycium burbarum (Marmé a. Husemann, A. Suppl. 2, 383; 3, 245; Ar. Ph. [3] 6, 216). -5. In putrefying flesh (Gautier, Bl. [2] 48, 13).

Formation.-1. From tri-methyl-amine and chloro acetic acid (Liebreich, B. 2, 13). -2, By oxidation of neurine Me,N(OII) CIL,CIL,OII.
3. Glycocoll (1 mol.) is dissolved in KOHAq and mixed with Mel (3 mols.) and McOll; the liquid being kept alkalino (Griess, B. 8, 1406). 4. Silver glycocoll and Mel gives the iodido. Me,N1.CH_.CO.11.

Preparation. -1. Diluted molasses are beiled for twelve hours with haryta; excess of baryta is removed from the filtrate by CO2; the liquid is evan tited to a treacle and exhausted with alcohol; the alcoholic solution is treated with alcoholio ZuCl,; the pp. is recrystallised from water, and decomposed by baryta; the barium ie exactly removed from the filtrate by H.SO. and betaine hydrochlorido crystallises on evaporation (Liebreich, B. 3, 161; cf. Schoibler, B. 2, 292; Frühling a. Schulz, B. 10, 1070).

Properties.—Large orystals (containing aq) (from alcohol). Ppd. as plates by adding ether to an alcoholic solution. Deliquescent. Over II,SO, the crystals become C,H,NO, Sweet taste; heutral to litmus; inactive. Decom-Sweet posed by heat, giving off odours of NMe, and of hurnt sugar. Not affected by CrO, or III. Fusion with potash gives off NMe, Iodino in KI pps. brown needles of a periodide.

Salts. -B'fici or Me, NCl.Cll, CO.H: mono-Salts. B'HCI o'r Me₃NCI.GH₂CU₂H; mono-clinio tables, v. sol. water. B'HAuCl₄; plates or thin needles. • B'H2PCl₂ 2aq (R₄).— B'₄H₂PCl₄ 4aq (L₄).—(B'HCl₂)H₄Cl₄. • B'ZnCl₄. -B'Kl 2aq [139²] (Korner a. Menozzi, G. 18. 351).—B'Kl [226²].—B'4H₂(Bil₄)₂ (Kraut, A. 210, 318).—B'₂H₂SO₄. Methyl ether.—Iodide INM₂, CH₄CO₄Me. Recombilious alwacoll and Maj (Kraut, A. 199).

From silver glycocoll and Mel (Kraut, A. 182, 160).

BETH A BARRA COLOUR C₂₁H₂O₃. [135°]. A dye extracted from a West African wood (Sadler a Rowland, Am. 3, 22). When dried at 100° it contains 3aq in the molecule.

RETORGIN G.H., O. 4.4. C.H. Me.(OH). [1:4:3:5]. [163°]. (6. 279°). \$\beta\$-Orcin. Di-oxy. p-xylene. Di-methyl-resorcin.

Formation.—1. By boiling (3)-picroerythrin with baryta (Stenhouse, A. 68, 101; Lamparter. A. 134, 248; Menschutkin, Bl. 2, 428). -2. From amido xylenol, C, H, Me, (NII,) (OH) [1:4:3:5] by the diazo- reaction (Kostanccki, B. 19, 2321).

Preparation.—The lichen Usnea barbata is thoroughly extracted with cold water (10 pts.) and CuO (1 pt.), the extract is mixed with IICl. A pp. of usnic und barbatic acids is formed. This mixture (I pt.) is boiled with water (40 pts.) and CaO (1 pt.) for four hours. An insoluble basic calcie usnute is formed while the barbatic acid splits up into CO, and betorein. Air must be excluded, for betorein exidises very readily. The littrate is at once neutralised with HCl. acidified strongly with acetic acid, evaporated (to 5 pts.) filtered from some tarry matter, and evaporated further to crystallisation. Recrystallised first from benzene, then from water. Yield $\frac{1}{10}$ per cent. (Stenhouse a. Groves, C. J. 87, 896).

Properties. - Less soluble in water than orein. Gives a more crimson colour with hypochlorites than orein does. Fe Cl, gives a green colour. Ammoniacal solutions turn red in air. Boiled with NaOH and chloroform it forms a red, non-fluorescent solution.

BETULIC ACID C, 111, O, [195°]. From betulin and CrO, in HOAe (Huusmann, A. 182, 378). White powder, v. sl. sol. water, v. sol. alcohol.—Ph₂(C₁₀H₂₁O₆)₂.

BETULIN C. HanO. [251°]; [258° eor.] (Hausmann, A. 182, 369). S. (alcohol) 7 at 15°; 4.2 at 78°. Occurs in the bark of the birch Lowitz, Crell. Chem. Ann. 1788, i. 302; Hünefold, J. pr. 7, 53; Hess, J. pr. 16, 161; Stühelin a. Hofstetter, A. 51, 79; Paterno a. Spica, G.

Preparation. - Tho bark is extracted with 96 p.c. alcohol, the alcohol evaporated and the residue after washing with water and with sodasolution is crystallised from benzene or naphtha. the crystals are finally decolourised with animal charcoal and recrystallised from alcohol (Franchimout, B. 12, 7).

Needles; may be sublimed. Insol. water, v. sl. sol. CS, sl. sol. alcohol and other. At 130 it gives un Anhydrido C, II, O. On distillation alone with powdered zine, P.O. or P.S., hydrocarbons are produced of doubtful constitution.

Acctyl derivative CasH su(()Ac)2 [2162]. BETULIN AMARIC ACID C, II, 20,6. From betulin and fuming HNO, (Hausmann, A. 182, 374). Crystals, v. sl. sol. water, v. e. sol. alcohol and other. At 110° it gives the anhy-

alcohol and other. At 110° it gives the anny dride C₃₁I₁₀O₁₁ [181°].

Salts. — K₁C₃₁I₁₀O₁₂. — CaBaC₃II₁₀O₁₂. — Ca₂C₃₁I₄O₁₄. — Pb₁C₃₁I₁₀O₁₂. — Cu₂C₃₁I₁₀O₁₄. — Cu₂C₃₁I₁₀O₁₄. — Ethyl ether Et₁C₃₂I₁₀O₁₄. [117°].

BETULORETIC ACID C. II. O. [91°]. white resin found on young hirch shoots and teaves (Kosmann, J. Ph. [3] 26, 197). Insol. water, v. sol. alcohol and ether. Oxidised by HINO, to pierie acid. -AgC3.Ha,O3: flocement pp.

BEZOAR. - A concretion found in the stomach or intestines of a variety of goat, Capra ægragus, or of the gazelle, Antilope Dorces. They sontain eliagic and lithopellic acids. BICHROMATES, same as DICHROMATES, q. S.

under Chromium, acids or.

BICUHYBA FAT .- The fat of Myristica bicuhyba coneists chicfly of the glycerides of myristic and oleic acide; it also contains emall quantities of resine and free fatty acid (myrietic acid), and a very small amount of an ethereal oil (Noerdlinger, B. 18, 2617).

BIEBRICH SCARLET v. p. 368

BILE .- A liquid secreted by the liver. It is viscid, of green or brown colour, and has a bitter taste. S.G. about 1.02. Faintly alkaline. Possesses an caudsifying power like soap. composition varies. Ox-bile contains sodium glycocholute and taurocholate, cholesterin, urea, fats, salts of acetic und propionic acids, glyceryl tri acetate, glyccryl tri-propionate, pigments, mucus, KCl, phosphates of Na, Ca, and Mg, and traces of iron, manganese, and eilica. Human bile is of a similar composition.

Latschinoff (B. 18, 3039) has shown that saponified ox gall contains, in addition to cholic acid (which is derived from the glyco- and tauro-cholic acids), a new acid to which he gives the name cholcic ucid. The latter acid, according to this investigator, occurs in two formsanhydrous C., II, O, and hydrated C., II, O, II, aq. Mylius (B. 19, 369) has found that choice acid by putrefactive fermentation is reduced to a body (the 'desoxycholic acid' of M.) which L. considers as identical with his so-called 'hydrated choleic acid.' As however L. (B. 20, 1043) was unable to convert his 'anhydrous choleic acid' into the 'hydrated choleic acid' by any other method than by boiling with acctic acid and M. (B. 20, 1968) was unable to effect the conversion even in this wuy, there appears to be little doubt that these so called 'anhydrous' and hydrated choleic acids' are quite distinct ucids (the conversion by AcOH is probably erroneons), the latter being identical with the 'desoxycholic acid' of M. Henco the 'anhydrons 'acid will be described as cholcic acid, the 'hydrated' as deoxycholic acid. Both these acids according to L. give dehydrocholeic acid on gentle exidation, and cholanic acid on more vigorons oxidation.

To cholic acid L. assigns the formula C23H12O3. but M. (B. 19, 369, 2000; 20, 1968) has shown by a long series of careful analyses that, almost beyond a doubt, it is represented by the formula Call to O, originally proposed for it by Strecker. On oxidation it first gives dehydrocholic acid C2, II4, O3 and then bilianic acid C2, II4, O4 (but no cholanic acid, which when obtained from cholic acid by earlier investigators, was due to the presence of cholcic acid).

Pig's bile contains sodium hyoglycocholate and hyotanrocholate instead of glycocholate and taurocholate; it also contains glycero-phosphoric acid and neurine derived from the decomposition of lecithin. In other respects it resembles ox bile. The various constituents of bile are separately described.

l'ettenkofer's test. Bile, or an aqueous solution of a salt of bile, is mixed with two-thirds of its volume of H.SO, and a drop of a 10 p.c. solution of sugar is added. On warming to 75° a crimson colour is produced. The reaction is

given by glycocholic, taurocholic, hyoglycocholic, hyotaurocholic, and hy cholic acid (Pettenkofer, A. 52, 92; cf. Neukomm, A. 116, 80; Strassburg, Pfluger's Arch., 4, 461). The test may be the strained by an analysis acid. The submodified by using phosphoric acid. The sub-stance to be tested, together with very little cane sugar, is dissolved in 3 drops of a mixture of syrupy phosphoric acid (5 vols.) and water (1 vol.) and the tube is then dipped into beiling water. A crimson colour soon appears (Kolbe, J. pr. [2] 27, 421). A red colour is produced by many other substances than those mentioned. hence it is necessary to confirm it by observing the absorption spectrum which contains three bands: one extending from midway between o and D to D, the second midway between D and E, and the third between B and F (Heynsins a. Campbell, Pflüger's Arch., 4, 497; cf. Schenk, Fr. 12, 119).

BILE COLOURING MATTERS v. PIOMENTS. ANIMAL

BILIANIC ACID

O2,HacO, probably C20Has(CO)2(CO.H)4. Formed by further oxidation of dehydrocholio

acid C₂₀H₂₁(CO)(CHO)₂CO₂H.

Preparation. -- Cholio acid (20 pts.) in fine powder is added to a mixture of K.Cr.O. (40 pts.) and H.SO4 (60 pts.) in water (160 pts.), finally warming on the water-bath till the reaction is complete. It is isolated by conversion into the acid potassium salt, which is sparingly soluble in alcohol, and then into the di-ethylether (Mylius, B. 20, 1981; cf. Clève, Bl. [2] 35, 373; Latschinoff, B. 19, 480; Bl. [2] 46, 818).

Properties .- Flat neodles (containing laq).

Tri-basic ketonio seid.

Salts.—A"₂Bu₃6aq or 8aq: tables or prisms. -A"'IIBa 2aq: hexugonal tables, sl. sol. water and alcohol.— $\Lambda g_1 \Lambda'''$.

Diethyl ether A"HEt,: [193°]; long flat needles; sol. alcohol, less sol. cther. —
A''',Et,Ba. — A''',Et,Pb.
Tri-cthyl ether A'''Et,: [127°]; satiny

tablets, or thick pillars.

Di-oxim C₁₈11₂₁(C:NOH)₂(CO₂H)₂. Formod by warming a slightly alkaline solution of bilianic acid with hydroxylamine. Glistening tables. Sol. dilute alcohol, nearly insol. water and absolute alcohol. Dissolves in alkalis, forming acid or neutral salts.

Di-phenylhydrazide

 $C_{10}H_{31}(\hat{C};N_a\Pi\hat{P}h)_a(CO|\Pi)_a$: colourless needles. Nearly insol. acetic acid and hot alcohol, insol. water. Dissolves in alkalis.

iso-Bilianie Acid C. [I_mO_a (?). [234°-237°]. Flat needles. Formed in small quantity, together with bilianic acid, by oxidation of cholic soid with K2Cr2O, and 112SO.

Salts.—A"H.K: silky rhombic plates, al. sol. water and alcohol.—A"Am: amorphous pp. - A"Ba 6aq: amorphous, sl. sol. wuter, insol. alcohol.

Methyl ether A"Mo,: [982]; neodles (Latschinoff, B. 19, 1530).

BILIC ACID C. H.O. [about 190°]. pared by careful oxidation of cholic acid with chromic acid mixturo (Egger, B. 12, 1958). White needles. Sol. hot water and alcohol, sl. sol. ether. Dibasic acid. 1t gives Pettenkofer's reaction with sugar and H2SO,. By oxidation it is converted into cholesteric acid (O, H, O,).

BINARY THEORY OF SALTS, The name salt was given in ancient times to the solid residue obtained by boiling sea-water; it was then extended to include all solid substances easily soluble in water and obtainable by evaporuting watery liquids. At a later time the possession of a taste more or less resembling that of sea-salt was regarded as a characteristic property of all salts. When the composition of the bodies called salts began to be studied, a threefold division was made into acid salts, alkaline salts, and neutral salts (v. Acro, Alkali, Salt). Lavoisier's discovery of the hature of oxygon led to the definition of acids as highly oxygenated compounds; und Davy's decomposition of soda, potash, lime, and baryta, showed that these alkaline salts were also compounds of exygen. But as neutral salts were formed by the mutual action of un acid and an alkali, it followed that they too were exygenated compounds. neutral salt, or we may say simply a salt, for the qualifying word neutral had been dropped by this time, was then regarded as constituted of two parts, an acid or electro-negativo part, and a basic or electro-positive purt. From this time dualistic views prevailed; overy compound, said Berzelius, must be constituted of two parts, which may themselves be simple or complex, and of these parts one is positively and the other negatively electrified. Such a salt as sulplinte of soda, for instance, was regarded as constructed of positive soda and negative sulphuric acid, rather than as formed by the mutual interaction of the elements sodium, oxygen, and sulphur. When, chiefly as a result of the work of Davy and Dulong, acids had come to be regarded as composed of the positive elemont hydrogen united with a negative element or group of elements, and salts were said to be formed by putting metals in the place of the hydrogenof acids, the conception of a salt as a binary structure still remained. One part of every salt was a positivo element, a metal; the other part was a negative radiole, either a nonmetal or a group of non-metallic elements. In some such way as this area the binary

theory of salts, a theory which is based on the notion of every salt being a definite structure, and which conceives it possible to place all salts in one class, regard being hud for classificatory purposes rather to the composition than to the properties of salt (v. Classification and Salts).

BIRCH BARK. Contains betulin (q. v.) and a kind of tannin which is turned green by Fe,Cl.

(Stahelin a. Hofstetter, A. 52, 79).

B18MUTH. B2. At, w. 208. Mol. w. probably 208 (r. Biltz a. Meyer, Z. P. C. 4, 249). [268°8] (Rudberg, P. 71, 462; Riemsdyk, C. N. 20, 82). (1096°, 1450°) (Carnelley a. Williams, C. J. 85, 565), S.G. 2 9 759 (Schröder, P. 106, 226). S.G. liquid 10:055 (Roberts a. Wrightson, A. Ch. [5] 30, 274). S.G. is lowered by greut prossure. S.H. (20°-48°) •0305 (Kopp, T. 155, 71); (9°-102°) •02979 (Bede, Mém. B. 1855-56, 28); (liquid 280°-380°) 0363 (Person, A. Ch. [3] 24, 129). C.E. (12°-41°) 001333 (Kopp, A. 81, 1); (0°-100°) 001316 (Mutthicssen, Fr. 15, 220; v. also Fizeau, P. 135, 372; 138, 267). It. of fusion at 266°8° 12,640 (Person, A. Ch. [3] 24, 129). T. C. (Ag = 100) 1.8 (Wiedemans

a. Franz, P. 89, 497). E. C. at0° (Hg at 0° = 1) holes in the crust, and pouring off the sell 18076 (Lorenz, W. 13, 422, 582). Cryst. form, molten metal; the crustle is found to be lined hexagonal, atc=1:13035; isomorphous with with orystals of Bi. Te, As, Sb. S.V.S. abt. 21 3. H.C. [Bi', O'] abt. 95,500 (Woods, P. M. [4] 4, 370). Emission-spectrum characterised by very many lines; in arc spectrum the predominant lines are 4722-1, 4119, 3595-3, 3510-4, 3396-2, 2593, 2524, 2400 8, 2277 (Liveing a. Dewar, T. 174, 187, v. also, regarding spectrum of Bi, Hartley a. Adency, T. 1884, 63).

Occurrence. Uncombined with other eloments, in Saxony and other parts of Germany, in Norway, Spain, California, and in Cornwall and Comberland, &c. Also as Bi₂O₄ (Bismuth ochre), Bi₂S₂ (Bismuth glance), Bi₂Ye₃, Bi₂Cu₂S₄, Bi Pb S, &c., &c. Bismath has been known for many centuries; Basil Valentine (15th century) seems to have been the first to recogniso it as a definite metallic body Bergmann gave the earliest fairly accurate account of its reactions.

Formation .- Native hismuth is melted in iron tubes, and the metal is run off from gangue &c. into pots. By remelting with 10 of its weight of KNO, at as low a temperature as possible, until the nitre forms a solid slag on the surface, approximately pure Bi is obtained.

Preparation. Approximately pure metal is disselved in the minimum quantity of HNO, Aq, much water is added, the pp. of basic nitrate is washed, boiled twice with pure KOHAq, er NaOHAq (Herapath, D. P. J. 169, 40), dissolved in as little HNO Aq as possible, and water is again added. The pp. is washed, dried, mixed with black flux (obtained by heating cream of tartar in a closed vessel) and heated at about 270° 280° in a closed crucible. The reduced metal is washed in dilute HClAq, and in water, and dried. Traces of As, Sb, or Fe, which yet remain in the metal are removed by partially oxidising and melting under the small quantity of Bi₂O₃ formed: this may be effected (1) by adding a little pure KNO₄ melting in an open porcelain crucible, keeping molton for some the metal solidifies, and pouring off the still molten metal from the more solid oxides on the surface; (2) by melting under Na,CO, containing 2 5 p.c. KClO, and proceeding as in (1) (Türach, J. pr. [2] 14, 309); (3) by strongly heating with I part cream of tarter, then running the molten melal (which contains K) into a orneible containing charcoal, heating for a little, running into an open porcelain vessel, strongly heating in air for some lime, and finally pouring off the molten metal from the slag on the surface (Mehu, D. P. J. 211, 187). Löwe (Fr. 22, 498) recommends ppn. of Bi3NO₃ in HNO, Aq by KOII, solution of pp. in excess of KOH in presence of glycerine, addition of grape sugar, filtration from Ag and Cu, and boiling; pure Bi is ppd. Bi is oblained in well-formed crystals by melting the commercial metal with a little KNO, in a crucible until a small quantity taken out appears yellow on the surface (indicative that foreign metals are oxidised), removing the seum of oxides from the surface, covering the molten mass with pieces of charcoal (to prevent oxidation), allowing to cool until a firm crust has formed, piercing two is easily exidised in moist air.

Properties .- Very lustrous; white with slightly reddish tings; very easily crystallises; brittle; diamagnetic, but not eo when molten (Faraday, P. Suppl. 3, 1; Weber, P. 73, 241; 87, 145; Reich, P. 97, 283; Plücker, P. 72, 339; 76, 576; 81, 133). Fer thermo-electric behaviour v. Svanberg, C. R. 31, 250; Franz, P. 83, 374; Matteucci, C. R. 40, 541. Bi expands as it solidifies (fer method of demonstrating this v. Böttger, D. P. J. 212, 441). May be distilled at a high temperature (over 1100') in an atmosphere of H. The atom of Bi is trivalent in the gaseous molecule BiCl₃. Bi forms numerous alloys most of which molt at lew temperatures, and expand on selidification (v. BISMUTH, ALLOYS er). The atomic weight of Bi has been determined (1) by finding V.D. of, and determining Cl in, BiCl, (Dumas, A. Ch. [3] 55, 129 a. 176); (2) by oxidising Bi to Bi O, by means of HNO, (Schneider, P. 82, 303; J. pr. [2; 30, 237; Löwe, Fr. 22, 498; Marignac, A. Ch. [6] 1, 289); (3) by converting Bi.O. into sulphate (Marignae, i. c.). The exact value to be given to the at. w. of Bi is still doubtful; it is certainly not greater than 208. Bi is metallic in its chemical functions; it shows a marked tendency to form basic rather than nermal salts; many of these basic salts may be represented as containing the group BiO e.g. BiO.NO, (BiO) SO, &c.; several exychlorides and oxybromides are known; no hydride of Bi has yet been obtained; the exides of Bi are salt-forming in their reactions with acids, none of them is an anhydride, but maist BigO, dissolves in very cone, boiling KOHAq probably with formation of compounds in which Bi acts as part of the negative radicle (v. Bismuthic oxide under Bismuth, oxides or). Bi shows distinct analogies to As and Sb, also to the other members of Group V., in its chemical relations; fer fuller discussion v. BISMUTH. OHEMICAL RELATIONS OF. Bi sults are used in medicine; the alleys are used in printing, soldering, &c.

Reactions. -1. Very superficially exidised in ordinary air : heated in air er oxygen burns to Bi.O. - 2. Decomposes steam at a red heat .-3. Combines directly with several *elements*, especially O, Cl, Br, I, S, Se, and To: [Bi,Cl³] = 90,630 (Th. 2, 410); $[Bi^2,O^1] = abt$. 95,000 (Woods, P. M. [4] 4, 370).—3. Scarcely acted on hy hydrochloric acid dilute or cone, -4. With hot cone. sulphuric acid a basic sulphate is formed.—5. Quickly dissolved by nitric acid with formation of Bi.3NO₃, -6. Oxidised, but slowly and partially, by fusion with potassium nitrate or chlorate.

Estimation.—Generally as Bi₂O₃, after ppn. from a solution free from HCl and chlorides by excess of ammonium carbonate, and warming for some time; the pp. is washed, dried, and heated whereby Bi₂O₅ is produced. Also by adding much water to a solution in as little HCl as possible, warming, collecting BiOCl, and drying at 100°-110° (traces of Bi remain nuppd.); the BioCl may be reduced by heating with KCN, and the Bi weighed. Ppn, as Bi-S, and weighing is not to be recommended, as Bi₂S₄ is easily exidised in moist air. Velumetric

methods of astimating Bi, none of which however is altogether satisfactory, have been based on (1) ppn. of Bi(IO2), from acetic soid solutions by a measured mass (excess) of HIO, Aq, and determination of residual H1O, (Buisson a. Ferray, M. S. [3] 3, 900); (2) ppn. of Bi chromate from nearly neutral solutions by K.CrO, Aq. or K,Cr,O,Aq; (3) pptn. of Bi phosphate by standardised Na,HPO,Aq; (4) ppn. of Bi oxalate, conversion into basic oxalate by boiling water, and titration with K.Mn.O.Aq; (5) ppn. of double oxalate of Bi and K by standardised K.C.O.Aq and determination of residual K.C.O. by K,Mn,O,Aq (Puttison Muir, C. J. 29, 483; 32, 674; 33, 70; (with Robbs) C. J. 41, 1).

Chemical Relations of Bismuth .- Bi is the highest known member of Group V.; this group contains the following elements:-

Taken as a whole these elements are negative and their compounds with I and O react as acids. As the group is ascended the negative characters become less marked until Bi is reached; omitting N, the oxides M.O. of the even series members, so far as known, are salt forming in their reactions with acids; but in the odd series members these oxides, on the whole, are saltforming only in their reactions with bases, until Series 9 is reached, when the character of the oxides M.O. becomes decidedly basic. oxides M.O., on the whole, are acid-forming, but the acidic functions of Bi.O. are very feedde. Salts formed from acids by replacement of H are obtained in the cases of V, Di, Er, and Bi; but most of the salts of vanadiam, and many of these of Bi, seem to contain groups of the form M.O. acting as the more positive part of the salt; several salts of the normal type, e.g. Bi.3NO,, are however known and several normal Er and Dr salts have been prepared. Vanadium is characterised by the great number of complex compounds into which it enters, sometimes as part of the positive, sometimes as part of the negalive, group of the salt. Considering the compositions of the linkoid, and oxyhaloid compounds, we find that, so far as investigation has gone, compounds of the form MX, and MOX,, where Y a halogen element, exist when M is any member of the group except Bi or As (N is omitted as there is much doubt concerning the composition of its halaid compounds), and in the case of As compounds of AsF, with KF, &c., seem to exist; the compounds BiBr (C.H.), and BiCl (C.H.), are known as solids (Michaelis, B. 20, 52). The haloid compounds liiX, are less easily axidised than the corresponding compounds of P. As, or Sb. The basic character of the existence of many salts in which Bi acts as the metallic element, the stability of the haloid compounds BiX, and the non-existeneo of BiX, the non-existence of any compound in which Bi,S, acts as the negative radiole; these, among other properties, show that Bi must be classed as the distincti-metallic element of Group V. But the feelig seidie functions of Bi.O. towards strong alledis, the readiness with which so calle I basic salts of bismuth are formed, the fact that Bio, and

Bi.O. form no corresponding salts, the existence of several complex oxyhaloid compounds; these, among other properties, show that the general non-metallic character of Group V. to some oxtent belongs to Bi.

Bismuth, Alloys of, Bismuth alloys with many metals when melted with them; these alloys are characterised by low melting points, and, many of them, by the expansion which they undergo as they cool after being melted. The most technically important alloys are: -Newton's metal; 8 parts Bi, 5 Pb, and 3 Zu, M.P. = 91°5: Rose's metal; 5 Fi. 3 Fb. 2 Su, M.P. = 91°6: Wood's metal; 15 Fi. 3 Fb. 4 Su, 3 Cd, M.P. 682: Fusible metal; 2 parts Bi, 1 Pb, 1 Sn, M.P. 9327; this alloy expands from 32' to 95', contracts gradually to 131° when its volume is less than at 32°, then expands to 1747, after which its expansion is uniform.

Amalgams of Bi are easily formed at ordinary temperatures. Alloys with copper are formed below the melting point of Ca; an alloy of 2 parts Bi with 1 part Cn begins to expand after solidification (Marx, S. 58, 470). An alloy of 3 parts Bi with 1 part iron is magnetic. Alloys of Bi and pulladium are hard as steel; with 1 part spongy platinum Ri forms an easily fusible alloy which separates into Pt and Bi when fused at a low red heat. By Joes not alloy with zinc: on mixing melted Zn and Bi two layers are formed, one containing a little Bi and much Zn, the other much Bi and little Zn.

Bismuth, Bromides of. Only one bromide, BiBr, has been obtained with certainty; but many facts point to the existence of a lower bromide, probably BiBr.

Thibboomide, BiBr, (Bi muthous bromide,) Mol. w. unknown, but probably as represented ly formula. (210°-215°) (Pattison Muir, C. J. 29, 144). (Untween 454° a. 498°) (Carnelley a. Williams, C. J. 33, 283).

Formation. - 1. By heating powdered Bi in CO, charged with Er vapour. - 2. By adding powdered Bi to a solution of Br in dry ether,

and evaporating in racuo.

Preparation, -1, parts Br are allowed to flow, in small successive quantities, into 1 part powdered Bi in a small retort with the beak tilted upwards; when the mass is cool, the refort is very gently warned for some days, and from time to time a few drops of Brane poured futo the retort; the bromide forms in yellow ery-tals a little distance above the heated nerge.

Properties and Beactions. Golden yellow erystals; S.G. 26, 54; very deliquescent; soluble in dry ether; decomposed by water to BiOBr and ItlirAq; partially reduced to birby heating in hy lingen; heated with Bi O forms BiOBr; by retion of nitrogen occider olds, and by heating starch with HNO, An the oxybramide BisOcker, is produced; unchanged when heated in CO, or SO,; reacts with anomonia, when he ated in that gar, to form (1) Billir, 3NB, which is a strawyellow powder, soluble in HClAq and yielding BiBr. SNH Cl.11.O by evaporation over IL SO .; (2) probably Biltr, 2NH, an olive-green solid; (3) an ash grey, crystalline, infusible, solid, pro-battly BiN, Er. The commontal 2BiBr, 5NH, is

obtained (along with Bi), as a greyish-green powder, by heating Bi.O., Br. to dull redness in dry NH. (Pattison Muir, C. J. 29, 144; 16, 27). A solution of BiBr, in saturated KClAq deposits orystals of BiCl, Br, K, 11H,O (Atkinson, C. J. 43, 292). Does not combine with Cl.

DIBROHIDE. - I'robably BiBr, or Bi, Br, In proparing BiBr, dark grey crystals are formed if there is a deficit of Br; these contain Br nearly agreeing with the formula BiBr_s; on heating they give Bi and BiBr_s. Weber (P. 107, 599) obtained a brown mass - probably a lower bromide than BiBr,-by heating BiBr, with Bi; Macivor (C. N. 30, 190) obtained a dark groy solid, melting at 198°-200°, by heating Bi and Br.

Blsmuth, Chlorides of. Two chlorides are known, BiCl, and Bi,Cl,: all attempts to form a chloride with more Cl than BiCl, have failed. Both may be obtained by the direct combination of Bi and Cl; Bi₂Cl, is separated into BiCl, Bi, and Cl, by heating; BiCl, is reduced to Bi,Cl, by heating with Bi, or with Hg.Cl.

Tractioning. BiCl, (Bismuthous chloride).
Mol. w. 314. [227] (Pattison Muir, C. J. 29, 144). (427'-439°) (Carnelley a. Williams, C. J.

83, 281).

Formation .- 1. By heating I part powdered Bi with 2 parts HgCl, in a retort.-2. By dissolving Bi₂O₃ in HClAq, evaporating to dryness, hoating in air and then in a retort .- 3. By

heating Bi.O. in dry Cl.

Preparation.-Powdered Bi is heated in a ourrent of dry Cl, in a retort with the beak tilted upwards and furnished with an exit tubs passing into cone. H,SO4; when a light yellow liquid has been formed, the stream of Cl is slackened and the retort is very gently heated for some time; crystals of BiCl, sublime on to the upper parts of the vessel. The crystals may be distilled into small tubes in a current oldry N; the tubes are at once scale

Properties and Reactions. - White, very deliquescent, crystals, melting in Cl in a pale yellow liquid; S.G. 12 4:56; soluble in dry alcohol. Hented in air between two watch glasses part sublimes and an oxychloride Bi₂O₂Cl₃ or Bi₄O₃Cl₄ -remains. The same oxy-obloride is obtained by the action of nitrogen oxides (from starch and HNO, Aq) on BiCl, Heated in hydrogen Bi Cl, and Bi are formed, at a higher temperature all the Cl is removed. Decomposed by water to BiOCl and MCIAq; the amount of change depends on the relative massee of BiCl, HCl, and H.O, and on the time; when the reacting bodies are mixed in the ratio BiCl_s: 26HCl: 19,000 H₂O a little BiCl₂ remains unchanged even after 14 days action (v. Pattison Muir, C. J. 35, 311; Ostwald, [2] 12. 2641. Heated with sulphur BiSCI is formed (r. Bismurn surmocuromor). Not acted on by Cro.Cl.; scarcely acted on by So.; does not combine with Br (P. M., C. J. 89, 83).

Combinations. - With ammonia to form (1) 2HiCl₂NH₂, a red, fusible, crystalline, solid; (2) BiCl₂2NH₂, a greenish solid; (3) BiCl. 3NH, a white, volatile solid (Déhérain, G. R. 54, 724). These ammonio chlorides by

and y home 1 to 5. With possessum chloride forms BiCl, 2KCl.2 H.O; also with sodium torms BIC4, 2M.O.; also with sodium chloride forms corresponding salt with BH.O.; as solution of BiCl, in hydrochloric acid when evaporated gives needles of BiCl, 2HCl (Jacquelain, A. Ch. [2] 62, 363).

DICHLORIDE. - Probably Bi.Cl., Mol. w. nn.

Formation .- 1. By gently heating BiCl, in H; but the product is mixed with Bi and BiCl. 2. By heating BiCl, with Bi (Weber, P. 107 596). - 3. By heating BiCl, xNII, Cl in H to 3005 (Schneider, P. 96, 130).

Preparation .- A very intimats mixture of 2 parts H₂Cl with 1 part extremely finely powdered Bi is heated to 230°-250° in a closed tube for soms time; the mixture melts to a dark brownish black liquid, and Hg (with a little Bi) collects at the bottom of the tubs; the sides of the tube are tapped from time to time to make the Hg settle; after cooling the Bi_sCl_s solidifies over the Hg, it is removed as quickly as possible to another tube—which is at once closed—and again mislted; this process is repeated several times; nearly pure Bi.Cl., containing a very little Hg and Bi, is finally obtained (Schucider, P. 96, 130).

Properties and Reactions .- Black, or nearly black, extremely deliquescent, solid; with water forms BiOCl; with potash gives Bi₂O, which is quickly oxidised to Bi,O3,...H,O; with dilute mineral acids gives Bi salts and Bi; heated to

about 300° gives BiCl, and Bi.

Bismuth, Cyanides and Ferrocyanides of,

C. CYANIDES.

Biemuth, Fluoride of. Only one fluoride of Bi has as yet been prepared (l'attison Muir, lloffmeister and Robbs, C. J. 39, 33), BiF, Mol. w. unknown.

Preparation.—I. Bi₂O₁ is added in small successive quantities to HFAq heated in a Pt dish until the oxide ceases to be dissolved; the liquid is decanted and evaporated at 100°; the residue, BiF, 3HF, is warmed at about 110°-120" in a closed Pt crucible until dry, and is then heated (in the closed crucible) so long as HF is evolved. - 2. Excess of saturated KFAq is added to a solution of Bi(NO₃)₃ in the minimum quantity of dilute HNO, Aq, the pp. is thoroughly washed with boiling water, dried at 100°, and heated to dull rodness in a closed Pt cruciblo.

Properties. - Grey, heavy, crystalline, solid. S.G. 7 5:33. Unacted on by water; incoluble in alcohol; scarcely changed or volatilised by heating to redness in open Pt dish; not oxidised by heating in nitrous oxides (from etarch and HNO,Aq); dissolved, with decomposition, by hot IICl, HNO, or H.SO, Aq. Combinss with IIF to form BiF, 3HF (v. supra) which is a erystalline, gevish white, deliquescent colid, decouposed by boiling water to BiOF (v. Brs-

MUTH OXYFLUORIDE).

Bismuth, Haloid Compounds of BiF,; BiCl, BiCl, BiBr, (? BiBr,); BiB, The V. D. of BiCl, only has been determined; the other formulæ ars probably molecular. Bi₂Cl₄ and Bi₄Br₄ are docomposed by heat to Bi and Bi₃; the others are unchanged when heated out of air; heated in air all sxcept BiF, are more or form zBiCl, which z varies from 1 to 2, the least, extent. Bil, is a very stable cor

pound (v. Brestute, Francisco or : Caronides or : mosmore or; and topion or).

. Mismuth, Hydrated oxides, or hydroxides of, v. BISMUTH, OXIDES OF

Bismuth, Iodide of. Bil. Mol. w. unknown: probably as represented by formula.

Formation.—1. By heating an intimate mixture of 1 part Bi.S. with 14 parts I, in a large, loosely covered thank, and then heating the sublimed Bil at 100 to remove I (Schneider, P. 99, 470). -2. By dropping Bi3NO, in dilute HNO, Aq into cone, KIAq, dissolving the loown pp. in fairly cone. HIAq, and ppg. Bil, by as little water as possible (R namelsberg, P. 48, 166), drying pp. at 100', and removing free I by one or two washings with absolute alcohol, -3. By the action of HIAq on Bi,O,

Preparation. -An intimate mixture of 1 part Bi with 2 parts I is gently heated in a flack with a long neck passing into another flusk; the sublimate is finely powdered and again heated in the same way as before; this is repeated once or twice; and finally the mass is distilled in a fairly rapid current of dry CO, (Weber, P. 14,

113 (slightly modified).

Properties and Reactions. (Pattison Muir, Hoffmeister a. Rabbs, C. J. 39, 33.) Dark grey, metal-like, histrous, crystal-quobably hexagonal, Nickles, C. R. 50, 872); S.G. 545, S. (alcohol at 20°) c. 3.5. Unchanged in air; bested in air a very little Bi Ox is formed. Unchanged by heating in kalvojen, or with sulphur, or in sulphur dioxide. Slowly changed to BiO1 by a farge quantity of cold water, more quickly by boiling water. Very partially converted into BiOI by heating in N oxides (from starch and HNO, Aq). Bil, is much more stable than either BiCl, or BiBr.

Combination .- With 111 to form Bil, 111.411.0 (Arppe, P. 11, 248). With MI (M. Na, K. NII.), and ML, (M . Ca, Ba, Mg, Zu), to form der de compounds isomorphous with the corresponding compounds of Sbl,: obtained by direct combination of the iodides, or by acting on Bi with I in presence of the iodide MI or MI,; they are all deliquescent, and are easily resolved by water into their component iodides (Nickles, C. R. 51,

1097; Linan. P. 111, 210).

Bismuth, Oxides and hydrated oxides ef. Four oxides are known, Bi₂O₂, Bi₂O₃, Bi₂O₄, Bi.O.; as none has been gasified, the V.D. and hence the molecular weight of none is known. These oxides all react with acids to form the same series of salts, BiX, where $X = NO_{s}$, $\frac{SO_{s}}{2}$, $\frac{CO_{s}}{2}$, &c.; if much water is present, basic salts, usually of the form BiOX, are produced; in the reactions of Bi₂O₂ with acids S.H. (12²-97²) 0509 (Regnault, A. Ch. [3] 1,129). Bi is separated as metal; in the reactions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₂ and Bi₂O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₂O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ and Bi₃O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821); 803 Bi₂O₃ expressions of *8.0. +8.21 (Herapath, P.M. 64, 821 but no salts have been certainly obtained in which the acid radicle is composed of Bi and O. Bi₂O₂ is easily oxidised to Bi₂O₂; Bi₂O₄ and Bi₂O₅ are deoxidised to Bi O, by heating in air or oxygen to about 320' and 250° respectively; Bi,O, is nuchanged when heated in air or oxygen. Bi.O, is not hydrated by contact with

water; Bi,O, and Bi,O, are hydrated in moist

air, in contact with water they are partially and

slowly deoxidised to hydrates of Bi.O.

The more important papers on the oxides at hydrated oxides of Bi are as follows:—1. On oxid containing less O than Bi₂O₂:—Thomson (Proc. Glasgow Phil. Soc. 1841-42, 4); Heintz (P. 63, 55, 559); Schneider (P. 88, 49; 97, 480); Arppo (P. 64, 237); Vogel (Kastner's Archiv, 23, 86); Berzelius (Lehrbuch, 2, 574 [5th ed.]); Schiff (A. 119, 331); Pattison Mnir (C. J. 32, 128). -2. On oxides containing more O than Bi,O,: --Jacquelain [1838] (J. pr. 14, 1); Heints [1844] (P. 63, 559); Arppe [1815] (P. 64, 287); Böttger [1858] (J. pr. 73, 494); Schröder [1862] (A. 121, 201); Boedeker (1862 (A. 123, 61); Wernicke (1870) (P. 111, 109); C. Hoffmann, [1884] (A. 223, 110); l'attison Mair (1876 to 1886) (C. J. 29, 144; 31, 21; 32, 128; ibid. 39, 21 [with Hoffmeister a. Robbs]; 51, 77 [with Carnegiej); Hasebrock [1887] (B. 20, 213). - 8. On Bi.O.: - Boundorff (P. 41, 305); Fuchs (S. 67, 429); Stromeyer (P. 26, 553); Liebig (Mag. Pharm, 35, 114); Pattison Muir (I.c.).

Hypobismurnous exide. Bi O2 (Bismuth subscide. Bismuth diswide. Black oxide of

bismuth).

Propuration. - A mixture of 1 part SnCl. and 25 part: Bio, is dissolved in as little fairly cone, HClAq as possible, the solution is poured into an excess of KOHAq (about I KOH in 10Aq) in a stoppered flask so that the flask is nearly tilled with the liquid; the stopper is placed in the flask, and the black pp. is allowed to settle; the pp. is washed with cold KO11Aq (in air-free water) less concentrated than that used in the ppn, the flask being each time nearly tilled with the liquid, and then with airfree water; it is then quickly dried by pressing between filter paper, and placed over Haso, in v.c.uo (Schneider, P. 88, 45).

Properties and Reactions .- Black, crystalline, powder; begins to oxidise in air at about 180 ; at ... heat quickly oxidised to Bi₂O₂; oxidised to Bi₂O₄xH₄O and Bi₂O₅xH₄O by boiling with KOHAq and Be; oxidised to Bi₂O₄xH₄O by K.Mn.O.Aq; who moist, Bi₂O₄ is rapidly oxidised in air to Bi₂O₂211.O; oxidised by contact with a very little IINO, Aq, decomposed to Bi, 6NO, and Bi by more IINO, Aq, dissolved entirely by a considerable quantity of the same acid; decomposed by HClAq or H,SO,Aq to BiCla or Bi sulphate, and Bi; deoxidised, to Bi, by heating in H or CO; decomposed by boiling KOIIAq with formation of Bi (Schneider, *Lc.; l'attison Muir, Lc.). No hydrate of Bi2O, has been definitely obtained. Solution of Bi,O. in turtaric acid is said to give Bi,S, by reaction with H.S (Schneider), v. Bisaum pisukemps.

Bismurnous oxide. Bi,O, (Bismuth trioxide),

Formation .- 1. By heating Bi in air or O .-2. By ppg. Bi nitrate or chloride solution by excess of alkali and boiling; thus prepared always contains some oxynitrate or oxychloride.

Preparation.—1. Basic Bi nitrate, obtained by ppg. solution of Bi in HNO,Aq by large excess of water, is heated in a Pt dish with. constant agitation until oxides of N are no longer evolved. If this oxide is fused with KOH it orystallises on cooling in rhomble

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prisms, a:5c = \$165:1:1064 (Nordenskjöld, P.)
114, 512).—2. BiOCl (q.v.) is shaken for some
sime with very conc. KOHAq, until the change
to Bl₂O₂ and KCl is complete; the Bi₂O₂ is washed with cold water until quito free from KOH and KCl and dried; the oxide may be thus obtained in distinct crystals.

Properties and Reactions .- Heavy, yellowishwhite, solid; unchanged by heating in air or oxygen. Dissolves in acids to form Bi salts (v. Bishittii, salts or). HFAq heated with Bi₂O₂ dissolves part of it as BiF₂3HF and converts the rest into BiOF.2HF; fIXAq (X = Cl or Br) added to Bi2O2 little by little converts the whole of the Bi into B:OX, on addition of more HXAq the BiOX dissolves as BiX,; with a little HIAq Bil, is alone formed, if very dilute 111Aq is added and the temperature is raised BiOl only is produced, with considerable excess of fairly cono. fllAq Bil, is formed and dissolved. Ifented in chloring, BiCl, is formed; with broming BiBr, sud BinO Br, are produced. Heated in carbon monoride reduction begins at about 2007, and in hydrogen at about 240". Bi.O. is not hydrated in moist air; nor is it altered by contact with water. Unchanged by heating in nitrogen. Oxidised to BigO, and BigO, by action of chloring in presence of much hot KOHAq; scarcely exidised by K.Mn.O.Aq (v. further Hyromsmornic, and Bismurine, oxner).

HYDRATED RISMUTHOUS OXIDE. Bi_O, x11,0; x=1, 2, and 3. The hydrate with 211,0 is obtained by dissolving Bi,O,xII,O in conc. 11ClAq, ppg. by KOHAq, and drying over 11_8O, in vacuo: the hydrate with fLO is obtained by dissolving Bi,O,xf1,O in conc. H₂SO₁, reducing by SO₂, ppg. by KOlfAq, and drying as before (P. M., C. J. 32, 131). Bi₃O₂3H O is very difficult to obtain quite free from oxy salts and Bi₂O₄; pps. formed by adding KOHAq to Bi₂O₅ in BCt, HNO₂, ox I₂, SO₄, and washing with cold water, always contain basic chloride, &o.; if washed with hot water they contain Bi O. Nearly pure Bi O. 3ff. O is obtained by dissolving Bi O. in the minimum of HNO,Aq, pouring into excess of cone. NII,Aq. ; washing with cold water until the washings contain no nitrates, then repeatedly agitating with very dilute Na₂CO₂Aq (to decompose traces of basic nitrates), again washing repeatedly with cold water, and drying in vacuo over H.SO. The hydrates of Bi₂O, are white solids, easily dehydrated by heat, partially oven by contact with hot water; BiOs does not directly combine with water. The hydrates behave towards acids and oxidisers similarly to Bi,Ox. None of these hydrates shows the slightest indientions of acidio functions. Thomsen gives the thornal values [Bir, Or, 311'O] = 137,710; [BiO'1P, HClAq] = 14, \$100, with formation of BiOCl + 211 OA4 (Th. 2, 244).

HYPORISMUTHIC OXIDE, AND HYPERATES, Bi2O4; Bi.O. 11.0; Bi.O. 211.0 (Arppe, Schröder, Böttger, Wernicke, Pattison Muir). Preparation of Bi₂O_x - Bi₂O_x is suspended in KOHAq, 8.6. abt. 1.85, the liquid is kept nearly boiling, and Cl is passed in until the solid is dark chocolatered and quite homogeneous to the eye; the solid is washed with hot water until the washings are neutral to litimus, kept in contact with dilute HNO, An (1 cone, acid to abt. 20 water) until the Bi, O, Cla, and BiBr, and a little Bi, O, Br,

colour of the solid has become brownish-yellow (12-16 hours) (to dissolve any Bi₂O₂ and reduce any Bi₂O₃), washed free from soid, and boiled with conc. NaClOAq (to reoxidise any traces of Bi,O.) until a heavy, yellow-brown, powder is formed which settles quickly; this powder is washed with hot water until quite free from alkali and Cl, and dried at 180°.

Hudrates.- If the drying is conducted over H.SO, the hydrate Bi_O, 1120 is obtained. If Bi.O.x11.O (v. inf.a) is treated with warm fINO Aq until the colour is orange-yellow, washed, and dried over fl SO,, the hydrato Bi.O. 21f.O is obtained. These hydrates are also formed, the first by the action of ordinary air on Bi.O4 and the second (with 2H2O) by the action of moist air on Bi₂O₄; they part with their water of hydration at about 150°.

Properties and Reactions. -BigO, is a brownish-yellow solid; S.G. 300 506; devaidised (to Bi,O₂) and dissolved by fairly conc. HNO₂Aq, or HClAq, more slowly by conc. H.SO₄; slightly deoxidised by contact with water in direct sunlight, oxidised to Bi,O, xf1,O by Clin presonee of hot conc. KOHAq; is not oxidised by ezonised O at 100' 140'; heated in Cl gives BiCl, and a little Bi,O,Cl,; heated in Brgives BiBr, and considevable quantity of Bi₁₁O₁₂Br₂. Heated in CO reduction begins at abt. 105° and the change to Bi,O, is complete at about 215°-250°; with H the corresponding temperatures are abt. 200° and 265', respectively; heated in air or in O the temperatures are abt. 240° and 320', respectively. Neither of the hydrates exhibits any decided acidic functions.

Bismetthe oxide, and Hydrate, Bi.O. (Bismath peroxide). Bi.O.,11,0 (Bismath acid). S.O. Bi.O. 5-917 (Brainer a, Watts, P.M. 1881, 62). S.V.S. 42. S.G. Bi.O.,11,0 5-75.

Preparation. Bi O₃₁ or BiO.11₃₁ or BiOCl, is suspended in about 10 parts of cone. KOllAq, S.G. about 138, the liquid is kept nearly boiling and Cl is passed in until a dark-rod howogeneous colid is formed; this solid is washed with hot water until the washings do not change the colour of red litmus paper and overy trace of chloride is removed, it is then warmed for a very short time with a little cone. HNO,Aq until its colour is scarlet, washed repeatedly and quickly with dilute HNO, Aq, each quantity of acid being more dilute than the preceding, and then with cold water until every trace of acid is removed. If the solid is now dried over 11.SO, Bi.O.H.O is obtained; if this is dried at 120' Bi.O. remains (Pattison Muir, C. J. 39, 22).

Properties and Reactions. A red, heavy powder; combines with water to form Bi,O, H,O in contact with much water is slowly deoxidised with production of hydrates of Bi₂O₄ and Bi₂O₅; also depaidised by hot dilute HNO₂Aq, giving first Bi₂O₄211₂O₅ and then hydrates of Bi₂O₄. Deoxidised to Bi₂O₄ by heating in current of air or oxyger at about 250', and to Bi O, hy heating in the same gases to about 305°; reduction in CO begins at about 70°, in ff at about 100°; reduction to BiO, is complete in H current at about 215°, and to Bi,O, at about 255°. Reacts with Cl, and Br, to give BiCl, and a little respectively. Does not exhibit any decided acidic functions; Bi₂O₂H₂O₃ however, dissolves in about 100 parts of boiling KOHAq so concentrated that solidification begins the moment the lamp is removed; on cooling, dissolving in as little water as possible, and nearly neutralising by HClAq (or by exposure to air) yellowishwhite solids are obtained from which all potash is removed only by very long-continued washing with boiling water. The solids dried at 10d contain a little water, Bi, and generally rather more O than is required by Bi₂O₂. Solutions of Bi₂O₂.H₂O in very cone, boiling KOHAq, therefore, probably contain compounds of the form xBi₂O₅yK O. In the preparation of Bi₂O₅ the very cone. KOllAq dissolves a little of the Bi₂O₅ as this is formed; on nearly neutralising with HClAq a white pp. is obtained, which, after longcontinued washing with boiling water, consists of Bi O. 311 O (Pattison Mair a. Carnegie, C. J. In the preparation of Bi.O, a portion of the potash is very obstinately retained; the whole of the potash can scarcely be removed by washing with boiling water; compounds of the form xBi,O,yK,O are probably formed, but every attempt to isolate these bodies has failed.

Bismuth, Oxyhaloid compounds of. Oxybromides, oxychiorides, oxylobides, and an oxythuoride, of hismath have been prepared. Althe haloid compounds BIX, where X=Cl, Br, or f, are oxidised by heating in air; only a very little BiOl is produced by long continued heating Bil,; BiBr, gives Bi,O, Br, as BcCl, gives Bi,O, Cl,. The same oxyhaloid compounds are formed by the reaction between N oxides, (from starch and hot HNO, Ap and hot BiX, the oxidation is carried furthest in the case of HiBr, in this case the whole or nearly the whole of the haloid compounds towards oxidisers are BiF, and BiI.

Oxygremmes. Three oxybromides are known; BiOBr, Bi_DO₁₄Br., and Bi₈O₄Br_o.

Bismuthyl bromide, BiOBr, is produced by the action of water on BiBr₃; or by heating Bi₂O₃ in HBrAq (BiBr₄; is formed in solution) and adding Bi₂O₃ ith HBrAq (BiBr₄; is formed in solution) and adding Bi₂O₃ little by little. It is a white amorphous powder; S.G. \(\frac{27}{27}, \frac{6}{7}; insoluble in water; unchanged when heated to redness; mixed with charcoal and heated in dry Cl, BiCl₄ is formed; reacts with cold HClAq to form BiCl₄ and BiBr₄, with cold HBrAq to form BiH₄, and HBrAq, and with lot HFAq to form BiBr₄, BiOF and BiF₂-8HF. BiOBr heated in NH₄ is reduced to Bi, and a little zBiBr₂, yNH₄ is formed, x probably = 2 and y probably = 5 (Pattison Muir, C. J. 29, 144).

The cryptomic Bi₃O₄ with excess of Br for some

The oxybroni le Bi₁₀O₃Br is produced by heating dry Bi₁O₃ with excess of Br for some hours and removing uncombined Br by warning in free contact with air. It is a cream-coloured, non-deliquescent, amorphous powder: unchanged by heating in air; unacted on by water; dissolved by warm HClAq and HNO₂Aq (P. M., C. J. 31, 24).

The oxphromide Bi,O,Br, is produced (1) by by water, or by heating in air; by long continued slowly subliming BiBr, in contact with a little heating to bright redness in air a very little air; (2) by passing N oxides toltained by Bi,O, is formed; reach with HClAq, HBrAq, heating starch and HNO,Aq) into melted BiBr,; and HFAq, similarly to bismuthyl chloride

in the first reaction only a little of the BiBr, is oxidised, in the second reaction most of the BiBr, is oxidised. In either case the product is washed with water and dried at 100°. This oxybromide is a grey, lustrous, crystalline, powder; unchanged by water, or by heating to redness; soluble in HClAq and cone. HNO,Aq; slowly reduced by 11, tinully giving B1; heated in, dry NH, B1 remains, and a grayish-green sublimate of 2BiBr, 5NH, is formed (P. M. C. J. 30, 12; 31, 24; 32, 40).

Oxyemonous. Three exychloroles, BiQCl, Bi;O,Cl, Bi;O,Cl, (or Bi;O,Cl, are known.

Bismuthyl chloride, BiOCl, is formed by adding water to BiCl, in a little HClAq; or by pouring Bi.3NO, in HNO, An into dilute NaClAq; or by reacting on excess of BiO₈ with very dilute HClAq; or by digesting a solution of BiCl₈ in HClAq with excess of BiO₉. The pp. is washed with cold water and dried at 100°. This compound is a white, listrons, exystalline, powder (known commercially as 'pearl white') S.G. 20° T·2. Reacts with cold HBrAq to give BiCl₈ and Bil₁; with hot HFAq to form BiCl₈ BiCl₉ and Bil₁; with hot HFAq to form BiCl₈ BiCl₉, and Bi₁; Alfe. Reduced to BiCl₈ by heating with charcoal in dry Cl (disquedain, J. pr. 14, 1; Appe, P. 54, 237; Ocston, P. 140, 128; Heintz, P. 63, 55; Pattison Muir, G. J. 39, 37).

The oxychloride Bi.O.Cl, is said to be produced by heating BiOCl to reduces (Arppe).

The oxicidoride Bi O.Cl, is formed in small quantities by slowly subliming BiCl, in contact with a little air, and in large quantities by passing N oxides (by heating starch with HNO,Aq) into melted BiCl. The analytical numbers agree fairly with Bi O.Cl, and also with Bi O.Cl., The compound is a yellowish-white, hard, crystalline, solid; unchanged in air, or by water or by heating to reduesa; soluble in het HClAq or HNO,Aq; boiled with NuOHAA, Bi O. and NaC Aq are ferned (P. M., C. J. 32, 10).

Over layouter. Only one is known. Hismathyl fluoride, BiOF, is obtained by heating BiO, with large excess of HFAq so long as any reaction occurs, boiling the residue with water until every trace of acid is removed, and drying at 100°; if the washing is conducted with cold water until nearly neutral, BiOF, 2HF remains; when this is strongly heated in a closed Pt erneible, BiOF is obtained in cry talline form. The liquid obtained by boiling BiO, with HFAq evaporated at 100° gives BiF₂3HF; when this saft is boiled with water it is slowly decomposed to BiOF, BiOF is a heavy, white crystalline, powders, S.G. 73, 7555. With HF it forms the double compound BiOF, 2HF. Reacts with cold HClAq to form BiCl, and HFAq; with HBAq to form BiR1, and HFAq; With HBA to form BiR1, and

Oxyronnes. Bismuthyl isoide, BiOl, is produced by boiling BiL, with H.O in small quantities, by subliming BiL, in air; or, also in small quantities only, by reacting with N oxides (from starch and hot HNO.Aq) on hot BiL. BiOl is a heavy, red, cry-failine powder, unchanged by water, or by heating in air; by long-continued heating to bright reduces in air a very little BiO, is formed; reacts with HClAq, HBrAq, distributed by the contract of the

Another oxylodide, probably 8BiOI.4Bi₁O₂, is obtained as a yellow powder by pouring a dilute solution of BiSNO₂ into KIAq mixed with NaC₂H₂O Aq (Fletcher a Cooper, Ph. 1882). Bismuth, Oxysulphids of, According to Hermann (J. pr. 75, 452) the compound

Bi,O,S, is formed by heating 1 part 8 with 8:55 parts Bi₂O₃ to low reduces in a retort; S.G. 6.3. A compound of Bi, S, and O, occurs as Karclinite (said to be Bi,O,S).

Bismuth, Phosphide of. No definite compound has been isolated. Berzelius (Lehrbuch, 2, 582 [5th ed.]) rays the two elements do not unite directly, but that a phosphide is formed

by leading PH, into Bi.3NO, solution.

Bismuth, Salts of. These compounds are obtained in a few cases by the reaction between Bi and an acid, e.g. Bi.3NO, but more generally by using Bi,O,x11.O in place of Bi, or by double decomposition from Bi.3NO, in 11NO Aq or BiCl, in HClAq. Bi. muth salts are insoluted in water; they are decomposed by water with production of so-called basic salts; the salts of Bi may indeed be arranged in two classes, normal and basic, us types of which may be taken the nitrates Bi3NO, and BiO.NO, respectively. Many of these basic sults are most simply regarded as derived from acids by replacement of 11 by BiO; they are often called bismuthyl salts; other basic salts, however, at present at any rate, are best represented as compounds of acid-forming oxides with Bi₄O₂. All the basic nitrates for instance, and many of these salts are known, belong to the general form xBi₂O₂yN₂O₃H₂O₅. The salts obtained by reactions between acids and the oxide: Bi₂O₄ and Bi₂O₅ are the same as there which aro formed when Bi₂O₂ is used. The more important salts of Bi are the nitrates and sul-phates, also bromate, chlorate, oralates, phase phates, tartrates, &c. (v. Nithaues, Schemains, do de).

Bismuth, Scionide of. Bi, Sep. Black, lustrous, metal-like, powder; S.G. 6-82. Obtained by passing H.Scinto Bi.3NO, in as little HNO, Au as possible, or by heating together I part Se and 1.8 parts Bi, and repeatedly melting the product in contact with Sc. Insoluble in solutions: of alkahs or alkali-sulphides; decomposed by HNO Aq; gives up So when deated. Combines with Fict, (v. infra) (Berzelius; Schneider, 1). 94, 628).

Bismuth, Scleaochlorido of. BiSeCl.

- Bi Se, BiCl, Permed by adding powdered neadle shaped, crystals. Heated in CO, is separated into Bi,Sc, and BiCl, (Schneider, I.c.); heating 1, S, and Bi,S., placed in alternate layers in a large crucible (P. 110, 147).

Bismuth, Sulphides of. One well-marked sulphide of Bi, Fi,S., is known; another, Bi,S., corresponding to the crystal Bi, O, 11,0 is sometimes called bismuthic acid:

Pags, corresponding to the oxide Bi,Og, probably exists. Attempts to prepare a sulphide with more S than Bi,S, have failed (Pattison Muir, C. J. 33, 192). Sulphide of bismuth does not react with more positive sulphides as a saltforming compound (comp. Schneider, Z. [2] 5, 680 with P. M., C. J. 33, 192).

History, Taisurunde, Bi₄S₆, Occurs native as bismuth adance. S.G. 6:5. Rhombic forms, a:b=1: 9881; isomorphous with As₂S₂ and

and bromide (Schneider, F. pr. 74, 424; P. M., Sh.S., Obtained by heating a minture of 1 part H. a. R., I.c.).

Another oxygodide, probably BBiOI.4Bi₂O_s, is formed, and then repeatedly heating this is obtained as a yellow powder by pouring a with a little S: also by passing H₂S into an adilute solution of BiSNO_s into KIAq mixed with obtained is heated with an alkali-sulphide solution to 200° the Bi,S, is said to become crystulline. Steel-grey, crystalline, lustrous, solid: strongly heated it is separated into Bi and S; unneted on by alkali or alkali-sulphide solutions.

BISMUTH DISULPHIDE. 2Bi.S. H.O. Said to be ppd. by H.S from alkaline solutions of Bi,O, Schneider (P. 97, 480) dissolved 8 grams Bi tartrate in the necessary quantity of KOHAq, added air free water to make up to 1500 c.o., then 2 grams SuCl, in KOHAq, and passed in air-free Il S mutil the liquid became colourless. He washed the black pp. with KOHAq and then with H.O (air-free), and dried at 100 '. A black powder; becomes lustrous by compression; decomposed by heating into Bi and Bi2S1, by HClAq into Bi, BiCla and H.S.

Bismuth, Sulphoehloride of, v. Bismurii, THIS DALOID COMPOUNDS OF.

Bismuth, Sulphoiodido of, v. Bismurii, Tuto-HALODI COMPOUNDS OF.

Bismuth, Sulphoeyanids of, Bi(SCN), v. CYANHOES.

Bismuth, Telluride of. No definite comapproximately Bi₂S_x2Bi₂Te₃, occurs native. According to Berzelius (*Lehrbuch*, 2, 581 (5th ed.)) the two elements may be melted together in all proportions.

Bismuth, Thiotraloid compounds of. Only two are known : BiSCl, and BiSL. The former is obtained by the direct reaction between BiCl, and S, but Bil, and S do not react together; when BiBr, and S are heated together there are indications of the formation of a thic-compound, but none has yet been isolated (v. P.M., H. a. R., C. J. 39, 21). The thio compounds are much less stable than the corresponding oxy- compounds; c. Bismuth, extraloid compounds or.

BISMUTH THIOCULORIDE (sulphochloride) BiSCl. Obtained by heating S with BiCl, or by adding powdered Bi, Sa to molten 2NH, Cl. BiCl., and washing the product with very dilute IIClAq, Small, metal-like, grevish, needles; easily docomposed into its constituents, e.g. by heating in CO, by II, by HClAq or HNO, Aq, and by alkalis (Schneider, P. 93, 464).

BISMUTH THIO-TOPHOE (sulphe-fielde), BiSI, is said to be formed, as long needles, by strongly

bodies obtained by dissolving Bi₂O₂ in much motten KOII, or by saturating with Cl conc. KOHAq holding Bi O, in suspension, have been described as bismuthates. But later experiments have shown that these bodies cannot be isolated although they probably exist in presence of much potash. The acidic functions of Bi.O. 11.O are extremely feeble, v. BISMUTRIO oxide under Biskutu, oxides of, p. 515. M. M. P. M.

BISHUTH, ORGANIC DERIVATIVES. Bismuth mono-methyl compounds.

Di-chloride BiMeCl. [242]. Obtained by adding BiMe, to an acetic acid solution of BiCl White plates. M. sol. alcohol and acetic acid, insol. ether (Marquardt, B. 20, 1520).

Di-bromide BiMeBr. (214". Formed by mixing ethereal solutions of BiMe, and BiBe. Yellow powder. Sl. sol. alcohol, benzene, and

acetic acid, insol. ether (M.).

Di-todida BiMel. [225]. Formed by heating BiMe, with methyl iodide at 200 (M.). Glistening red crystals. Sol. alcohol, sl. sal. acetic acid, insol. other.

Oxide BiMeO. Formed by adding NII, to an alcoholic solution of the double compound of BiMeBr, and BiBr, which is obtained as a yellow erystalline pp. on mixing ethereal solutions of BiMe, (1 mol.) and BiFr, (2 mols.) (Marquardt, B. 20, 1522). White powd r. Insol. water. Dis solves in NaOII and in dilute HNO. Ignites in tha air if gently warmed.

Bismuth di-methyl campaunds,

Chloride BiMe Cl. [116]. Formed by passing chlorine into a solution of BiMe, in etroleum ether, cooled in a freezing mixture. petroleum-ether, coorea in a White micro-crystalline powder. V. sol. alcohol, insol. ether (Marquardt, B. 20, 1519).

action of water upon the double compound of Billr, and BiMe, Br, which is pard, as an oily liquid by mixing othereal solutions of equal mols, of BiBr, and BiMe, Crystalline solid. Ignites in the air spontaneously. Discolves in aqueous NaOH. Decomposed by aqueous HCl with evolution of CII, (Marquardt, B. 20, 1523).

Bismuth tri - methide BiMe. Tri-netical. bismuthine. (110). S.G. 23 at 182. Obtained by slowly adding an othereal solution of biamuth bromide (2 mols.) to an othereal solution of zinc methide (rather more than 3 mols.). Molille refractive liquid, of impleasant pungent odour. In the air it formes and rapidly oxidises, when heated in the air it explodes violently. Volatile with steam, but decomposes on long boiling with water. Dilute II SO or IINO, have little action upon it, but cone. HCl decomposes it with evolution of Cil, and production of BiCl, It does not combine with alkyl haloids or with halogens. The latter replace Me forming BiMe Cl. &c. (Marquardt, B. 20, 1517).

Bismuth mone ethyl campounds.

alcoholic HgCl, to alcoholic BiEt, (q. v.).

Iodide BiEtl. From the chloride and KI. Golden, six-sided plates.

Oxide BiEtO. From the iodide and potash. Yellow amorphous powder, takes fire in air.

Nitrate BiEt, NO. J. From the oxide and

HNO. Crystalline tufts.

Bismuth di-ethyl hromide BiEt Br. Formed by dropping bromine into a co 'ed solution of , up by heat into NII, and eyal, ric neid. Dis-BiEt, in petroleum ether. White powder. V. sol. sloohol, insol. ether. Iguites in the air on warming (Marquardt, B. 20, 1520).

Bismuth tri - ethide Bilit,. Tri . cth ft. bismuthine. (107°) at 79 mm. S.G. 1 82.

Preparation .- 1. An ailoy of Bi and K is made by strongly heating bismuth (5 pts.) with eream of tartar (4 pts.). This alloy is treated with and area. 2. Boiling conc. HClAq forms NIL

adding an ethereal solution of bismuth bromide (2 mols.) to an ethereal solution of ZnEt, (rather

more than 3 mois.) (Marquardt, B. 20, 1519).

Broperties. - Stinking oil. Is cannot be distilled at ordinary atmospheric pressure, for on heating to 150° it detountes violently; volatile with steam. Funnes and takes fire in air. V. sol. alcohol, ether, and acetic acid. Its ethercal solition exposed to air deposits Bi.O.H O. Forms unstable compounds with non-include. Billt S.Bi S, is a yellow solid, insoluble in water, soluble in yellow ammonium sulphue. Bilit, throws down calomet from alcobolic HgCl, but when alcoholic HgCl, is added to alcoholic BiEta crystals of BiEtCl, may be got: BiEta, 2HaCl, 2HBEtCl, (Dünhanpt, 4. 92, 371).

Bismuth-tri-phenyl Bi(C.H.), Tri-phenyl-bismuthine, [82], 8.0, 15851 at 20°, Formed by heating bromo-bendene containing some acetia ether with an alloy of sodium and leannith (10 p.c. Na). Colcurless needles or taldets. Sol. hot alcohol, sparingly in eddl, v. sol, ether and petroleum ether. By boiling with cone, HCl it is completely decomposed into benzeme and bismuth chloride. It combines with Cl. forming the chloride (C.H.), BiCl, which crystallises in thick prisms, (110%, sol. hot alcohol, al. sol. ether and cold alcohol, v. sol. benzone, not decomposed by cone. HCl. The bramids (C.H.), BiBr, forms long prisms, [119], v. sol. benzene, al. sol. alcohol and other (Michaelia a. Weits, R. 20, 54),

BITTER ALMOND OIL r. ALMONDS and Вихлов админутог.

BIURET CHINO, i.e. NII, CONH.CONH Allophagaide. Mol. v., 103. [1904]. 8, 125 at 105. France on 1.

Facilities. 1. Unca is heated at 1502, 1702 until the melted muss becomes party and ceases to give of Nala. The pro-fact is extracted with hot water (Wiedemann, P. 74, 67; Hofmann, R. 4, 262). 2. thy positive the vapour of cyanic acid into melted area (banck, A, 121, 331). -3. Urea is treated with elderine till the muse becomes pasty (Boppert a. Dogiel, Z. [2] 3, 691; B. 4, 175). 4. By the action of NII, on all aphania ether (H. a. D.). 5. By he ting famido-dievanic acid' (p. 165) with H SO, (1 pt.) and water (2 pt.) at 60 · 70' (Boneman, B. 8, 708). 6. By the action of NII, upon tri bromo acetylurea (Bueyer, A. 130, 154). 7. By heating urea Chloride BiEtCh. Prepared by adding with PCl, at 100° (Westle, D. 10, 1743). 8, By mixing drinte solutions of uren and potassio evanute, acidifying with acctic acid, evaporating, craining a little H₂SO₂ and extracting with alcohol (Drechsel, J. pr. 128, 472). 9. By electrolysis of a solution of NH₂, using carbon electroles (Millot, Bl. [2] 46, 244).

Properties, "Long modiles containing ag); or long anhydrous lamina (fros calcohol). Split solves unchanged in cold conc. 11,80,. Its solution is not poel, by saits of lead or silver or by tannin. A little CuSO, Iollowed Ly KOH

gives a deep violet solution.

Reactions .. - 1. At 120° it absorbs HCl form. ine B'zHCl which at 160°-170° in a current of HCl gives H.O. CO, guanidine, cyanurio acid, Etl (Breed, A. 82, 106). -2. Obtained by slowly urea, and guanidine. -3. Boiling baryta-water

BIURET. 518

(Fenten, C. J. 35, 14).—5. HNO, gives CO, and N₂O in equal volumes (Franchimont, R. R. C. 6, 216). - 6. COCl2 at 60° forms carbonyl-di. biuret (C.H.N.O.), CO (E. Schmidt, J. pr. [2] 5, 47); a crystalline powder, v. sl. sol. cold water, insol alcohol and other. This body is converted by boiling baryta-water into urea and cyandric acid; and by COCl, at 140° into cyanurie acid; Hg(NO,), gives, in hot dilute solution, a pp. of C.H.N.Ö. Higo.

Salts. BullCl; decomposed by water .-Ag₂C₁H₁N₂O₂: ppd. by adding AgNO₂ (2 mols.) and NILAq to a saturated aqueous solution of hinret (1 mol.); sol. HNO3 and NII3Aq (Bonné a. Goldenberg, B. 7, 287). - Cyanurate B'C, N, II, O,: needles; formed on crystallising biuret from water. It has probably been mistaken for urea cyanurale from which it differs in yielding with baryta water, barium cyanmute and biaret, in giving off 3 atoms of nitrogen as ummonia when heated with barium hydrate (urea cyanurate yields 2 atoms), in giving off 118 p.c. nitrogen with sodium hypobromite, (while urea cyanurate gives 115 p.c., both results corresponding to 2 atoms nitrogen) (Herzig, M. 2, 111).

Biuret dievanamide Call, N.O. i.e. NH(CO.NH.C(NH).NH_)2. From neetyl-nrea (2 pls.) and guanidine carbonate (5 pts.) at 110° 150° (Rusinski, J. pr. [2] 27, 157). Amor-

phons substance, v. sol, acids and fixed alkalis, insol. NH, Aq. Does not give the binret reaction with CuSO,

BIXIN C_{pt}II_{It}O_p. [1760]. A colouring matter contained in annatto seeds (Bixa orelland) (Preisser, A. 52, 382; Girardin, J. Ph. (3) 21, 171; Bolley a. Mylius, Bl. [2] 3, 230; Stein, J. pr. 102, 175).

Preparation. - Annutta (1,500 g.) 🕭 digested at 80" with alcohol (2,500 g. of 80 p.c.) with mblition of Na CO, (150 g.); after filtration, the residue is again digested with alcohol (1,500 g. of 60 p.c.). The mixed filtrates are ppd, by adding half their bulk of water and cone. Na CO Aq; the ppd. sodium-bixin is dissolved in alcohol (60 p.c.) and re-ppd, with Na, CO, Aq. The sodium bixin is then decomposed by Hell (Eti, 17, 11, 861; 7, 416).

Properties. Minute red leaflets; insol, water, sl. sol. ideoliol, benzene, CS, and acetic acid, v. c. sol. ether. Conc. H.SO, forms a bright blue solution, whence water gives a dark green pp. It reduces cold Pehling's solution.

Reactions,-1. Distillation with zine dust gives 24 xylene, rthyl tolnene, and an oil C11111 (270 280). Reduced by sodium-amalgam to $\mathbf{C}_{p}\Pi_{\mathbf{m}}\Omega$.

C. H. NaO, 2aq: lustrous red erys-Salts. tals, v. sol. water, insol. alcohol, and ether. C., I., K.(J. 2aq. C., II., K.(J. 2aq. BLEACHING-POWDER P. HYPOCHLORITES,

under Cheorine, oxyacins or.

BLOOD. In vertebrates, the blood is a somewhat viscous, and to the maked eye homogeneous, red liquid. The blood which leaves the lung or gills is of a bright scarlet colour, and that in the systemic veins of a purplish

forms CO, NH, and urea.—4. With NaClO it | ing with oxygen, becomes of the bright arterial evolves 1 of its nitrogen, with NaBrO it evolves 3 | searlet colour. This difference in tint is due to the amount of oxygen present in combination with the red pigment hæmoglobin; in the lungs a loose combination called oxy-hamoglobin is formed which is scarlet; in the tissues this oxygen is given up, and the blood returning to the heart is of the purplish colour due to havinoglobin.

Specific Gravity. Roy (Proc. Physiol. Soc. 1884) has introduced a method for ascertaining the specific gravity of living blood. A drop of blood is introduced into a mixture of glycerin and water of known specific gravity; if the drop tends to rise or sink, it is assumed that it is of lower or higher specific gravity than the liquid in which it is placed. The average specific gravity of human blood thus found was 1.060. Defibrinated luman blood has an average specific gravity of 1.055. Pflüger (Pflüger's Archiv, i. 75) found the specific gravity of dog's blood to be 1060; and Gschleidlen that of rabbit's blood 1:018.

Characters. .. Blood is always feebly alkaline in reaction (Külnne, Virchow's Archiv, 33, 65; Liebreich, B. 1, 48; Schäfer, Journal of Physiology, 3, 292). Under the microscope the blood is seen not to be a homogeneous red liquid, but to consist of a nearly colourless liquid, the plasma or liquor sanguinis, holding in suspension large numbers of solid bodies, the corpuscles. These corpuscles are of two kinds, the coloured and the colourless.

Red corpuscles. These owe their colour to hæmoglobin, and are much more numerous than the white corpuscles. They vary in size and structure in different groups of the vertebrate sub-kingdom. In Mammalin, with the exception of the Camelidie, they are biconcave, circular dises; they have no nucleus except during embryonic life; and thry have a tendency to run into ronleanx when the blood is at rest, but if it is disturbed they readily become separated. the Camel tribe they have an elliptical outline. Their average diameter in manimuls is '007-'008 millimetro and about one fourth of that in thickness; there are very slight variations in different classes of mammals. In birds, reptiles, amphibians, and fishes, the red corpuscles are biconvex, oval discs, with a nucleus; they are largest in the amphibia. C. Schmidt gives the specific gravity of red blood corpuscles as 1.089, Welcker as 1.105.

According to C. Schmidt, 1,000 parts of moist red corpuscles contain-

> 688 parts. Solids { Organie . Mineral . 303.88 ,, 8.12 ,,

According to Hoppe Seyler and Jüdell (Med. Chem. Unterschungen, Heft iii. p. 386) 100 parts of dried corpuscles contain -

•	Human	i blood	Dog's	Googe's
	I.	11.	b ood	h'ood
Proteids	12.24	5.10	12.55	36.41
Hæn: əgləbin	86.79	94.30	86.50	62.65
Lecithin	0.72	0.35	0.59	0.46
Cholesterin	0.25	0.25	0.36	0.48

The nuclei of the red corpuscles consist mainly, according to Lauder Brunton, of nuclein, a substance very akin in its properties to mucin hue, which ou exposure to the air, or on slink. (Journal of Anat. and Physiology, 2nd series,

BLOOD.

vol. 8, p. 91). The mineral constituents of the red corpuscles have been investigated by C. Schmidt, and the following tables contract these of the red corpuscles with these of the plasma in man.

1000 parts of moist Mineral matter (exclusi	corp	uscle:	s yiel	d— ich	
is contained in the he	emo	globie	i,	,	8:120
Chlorine	,	٠,			1.686
Sulphuric anhydride					0.066
Phesphorus pentoxide		•			1.134
Petassium					3.328
Sodium					1.052
Calcium phosphate					0·11·t
Magnesium phosphate				,	0.073
1000 parts of plasma	a vie	1.1 -			
Mineral matter .					8:550
Chlorino , ,					3.640
Sulphuric anhydride					0.115
Phosphorus pentoxido					0.191
Potassium					0.323
Sodium				,	3:311
Calcium phosphate				,	0.311
Magnesium phosidiate		,			0.222

The remarkable difference in the distribution of potassium and sodium seen in the above does not, however, hold for most animals as the following table shows (Gangee, Physiological Chemistry, p. 122).

	Black	Cells	
	к.	Na.	Ct.
Man	40.89	9.71	21.00
Dog	6.07	36.17	2t.83
Cut	7.85	35.02	27.59
Sheep	14 57	35.07	27.21
Cont	37:11	14.98	31.73
	Liquor 8	mornis.	
	K.	N.E.	Ci,
Man	5.19	37.74	40-68
Dog	3.25	39.68	37:31
Cat	5-17	37.64	41.70
Sheep	6:56	38.56	40.89
Goat	8:55	37.89	40:41

Probably the only gaseous constituent of the red blood corpuscles not in a state of chemical combination is carbonic acid.

Bloot tablets. Besides the red corpuscles, a number of colourless dies of 002-003 millimetre diameter are also seen: they are also called hematoblasts (Blutplättchen of Bizzazero). By some they are supposed to be stages in the development of red corpuscles; by others to take part in the formation of fibrin. Wooldridge considers them to be identical with the protein he calls fibrinogen A, which can be precipitated from liquor sunguinis by exposure to cold.

Coburless corpuscles or leucocytes. These are animal cells, and consist of nuclented masses of protoplasm, more or less granular, and exhibiting during life contractility, the movements so produced being called annabold. They are not constant in size, but in near they average about 0.01 millimetre; they are comewhat larger in the lower vertilerate groups. In mammals there is on the average one white or colouriess corpuscle to 330 or 350 red ones.

Our knowledge of the chemical constituents of the white corpuscles is meagre; they are lighter than the red ones; the great mass of the protoplasm is undoubtedly proteid in nature; and the nucleus consists mainly of nuclein

(Miescher, Med. Chem. Untersuch., Heft iv., p. 441). By miero-chemical investigation, the presence of glycogen can eften be demonstrated by iodine, and of fat granules by osmic soid.

Coagulation of blood. Within a few minntes after having been shed, blood passes first into the state of a soft red jelly, which gradually acquires greater consistence; and by the contraction of one of its constituents expresses a fluid, the serum, in which the clot or crassamentum ultimately floats.

Coagulation is due to the separation from the blood plasma of a solid proteid substance.

mentum ultimately floats.

Cosquistion is due to the separation from the blood plasma of a solid proteid substance called fibrin. The clot consists of fibrin entangling the corpuscles. By stirring blood, or whipping it with twigs immediately after it is slied, fibrin free from corpuscles adheros to the twigs as a yellowish stringy mass. Under the microscope, coagulation is seen to consist of the separation of fine tilaments from the plasma, which start from or entangle the blood plates, and corpuscles.

Congulation of the blood is hastened by exposure to a temperature rather above that of the body; by contact with foreign matter, or by agitation; and by dilation with not more than twice its volume of water. Congulation is hindered or prevented by exposure to a low temperature; by contact with living tissues, or by the addition of large quantities of neutral salts such as solume chloride, sodium sulphate, or magnesium sulphate. When these precautions are taken, the corposcles sink, and the plasma can be drawn off; in the last case mixed however with salt solution, the inhibitory influence of which on chagulation can be removed by diluting the mixture with water; librin is then formed. Many theories have been held with regard

Many theories have been held with regard to the cause of the congulation of the blood. Nearly up to the end of the last century, the clot was believed to be simply a mass of udheren corpuscles. Hewson, 1772 (r. Hewson's works, edited by Galliver, Sydenham Soc.), was the first to show that it was really due to the separation of some substance from the plasma. Buchunan (London Medical Gazette, vol. 18) showed that squeezed blood-clot had the power of hastening the coagulation of the liquor pericardii; and as this power was especially shown by the baffy coat, he supposed that it was due to the white corposeles; he compared the action of the e corpuseles to that of remet on milk. Denis (Memoire sur le sang, 1859, p. 32), by saturating the liquor sauguinis with sodium chloride, obtained a proteid pp., which after being rediscolved in water underwent coagulation. To this precuesor of fibrin be gave the name plasmine. A. Schmidt (Archie f. Anat. u. Physiol. 1861, 545) separated plasmine into it: two constituents, both prescids of the globulin ciase, to which he give the names fibrinogen and fibrinoplactin or pauc lobulin (now called scrum globalin). He thought leth of these substances were necessary for congulation, and that they united to form fibrin under the influence of a ferment. This fibrin-ferment he prepared from serum, by ppg. it with the serum proteids ly means of absolute alcohol; after leaving the pp. some months under alcohol, the proteids were by this means rendered insoluble, while the ferment could be extracted with water. Gamgee (Journal of Physiology, 1879) obtained fibrin-ferment by extracting blood-clot with 8 p.c. sodium chloride solution. The extract contained a small quantity of a globulin-like proteid, and had very marked power in inducing congulation. Hammarston (Plitiper's Archivel 14, 211, 17, 413; 18, 38; 19, 563) modified Schmidt's theory by showing that paraglobulin is not necessary for the formation of fibrin, but that fibrinogen is the only true fibrin precursor which under the influence of the fibrin fermeant is converted into fibrin. The presence of paraglobulin, however, lastens congulation perhaps by its combining with alkaline carbonates which otherwise would impede the action of the ferment; other proteid; such as casein, or even salts such as calcium chloride, will, however, take its place.

The source of the fibrin-ferment seems to be the white corposeles. Ranschenbach has pointed out (U,b,τ) the Weckselu irkangen zwischen Protoplasma and Blatplasma, Inang. Diss. Dorpat, 1883) that hencogytes are of two kinds: a barcocytes which are acted upon and disintegrated by the plasma when the blood is shed, two of the products of such action being paraglobulin and fibrin ferment; and β leucocytes which remain unaltered.

The latest theory of the congulation of the blood is that of L. C. Wooldridge (Beitrage zur Physiologie, Leipzig, 1887, 221). He injects eptone into the circulation of an animal, and kills it by bleeding; the blood remains uncoagulated for many hours, and the corpuseles are removed by centrifugalising; if a substance containing legithin be then added to the peptone plasma, congulation occurs. By cooling peptono plasma, a pp. is produced; this consists of little rounded discs similar to blood tablets; this is called fibringen a; after its removal from the plusma congulation does not occur; the furniz-tion of fibrin is supposed to be due to the locithin contained in filtringgen a combining with librinogen b (Hammarsten and Schmidt's fibrinogen).

Human blood yields from 2/2 to 2/8 parts of fibrin per 1,000.

Serum. This is the phason, minus the elements of fibrin. It contains three classes of constituents; proteids, extractives, and salts. The preteids consist of globulin and albamain. Owing to the disintegration of the white corpuscles the globulin is rather more abundant than in the phasma. The following table of Hammarsten's (Urluce's Archir, 1878) represents the percentage of these substances in the serum of some of the commoner manuals:

			Total Solida	Total Potente	Seraja Globa in	Serum Albertan
Beran	fro	nı Horse	86597	. P.C. (1966	2007
-		()x	6:1:65	7:1:93	4/1/11	34.429
4	**	Mon	9.207	7 619	34.43	4.51.6
99	*1	Rabbit	7:525	6.57	1	4, 136

The globulin appears to be a single substance; it is coagulated by heat at 75° C.: by fractional heat-coagulation, however, the serum aliminin can be differentiated in some unimals into two, in some into three proteids (Halliburton, Journal of Physiology, 5, 152; Kander, Arch.f. exp. Path.

10. Phirmac. 20, 411); in the cold-blooded however, are anhuals the total quantity of proteids in the

serum is much lower, the serum globulis is always greatly in excess of the serum albumin, and the latter substance is not differentiable into several proteids by fractional heat-coagulation (Halliburton, Journal of Physiology, 8, 319).

The extractives of serum ars organic substances present in small quantities, which are extracted by various liquids, especially by alcohol or ether. There is about 0.2 p.c. of fats and cholesterin; about 0.8 to 0.12 p.c. of glucosa (ravy, Cromian Lectures on Diabetes, London, 1878); urea 0.02 to 0.01 p.o.; and creating, exanthine, hypoxanthine, uric acid, and hippuric acid in still smaller quantities. A yellow pigment is found dissolved in varying quantities in the serum of most animals; Hammarsten (Maty's Jainberricht, 1878, 129), MacMunn (Pr. 31, 231) and others have described this asa biliary pigment; Krakon'org (Sitzungsb. der Jenaischen Gesellsch. f. Mad., 1885), and Halliburton (Journ. of Physiology, 8, 321) have described it as a lipochrome.

The salts of serum, mount to 0.7 to 0.9 ns.

The salts of serum Amount to 0.7 to 0.9 p.c. Gases of the blood. From the blood as a whole, or from the plasma, colonied corpusolos, or serum, a mixture of carbonic acid, oxygen and nitrogen can be separated. Oxygen present in much larger quantities than could be held in simple solution in the blood, and is, in fact, held in feeble combination with hemoglobin; only a small part being in solution in the liquor sanguinis. Carbon dioxide is partly in a state of chemical combination, but objetly in a state of simple solution. It is contained in great part in the liquor sanguinis, but in part also in the corpuscles. The nitrogen is held in simple solution in the liquor sanguinis. Arlerial blood of the dog yields for every 100 volumes 58:3 vols, of mixed gases composed of 23-2 vols. of oxygen, 34-3 vols. of earlionic anhydride, and 1'8 vols, of nitrogen. The maximum amount of oxygen observed has been 25:4 vols. (Pdiger, Controllof, f. d. m. l. Wissensch, 1868). In venous blood, the nitrogen is the same as in arterial, the oxygen is less in amount (from 8 to 12 vols, per 100 of blood), and the CO2 greater (from 40 50 vids, per 100 of blood).

Lymph. This is the name applied to that portion of the blood that transades through the walls of the blood-vessels, and after supplying the tissues with nutritivo materials and receiving the products of their combastion returns to the large value by means of the lymphatic vessels.

Lymph is a transparent Equid, which during dige-tion is more or less milky, owing to the suspension in it of fatty in trees absorbed from the alimentary canal. Its specific gravity varies between 1012 and 1022, and its reaction is alkaline.

alkaling.

Under the microscope, the lymph is sen to contain colourless carpusales. In a time varying from 3 to 20 minutes after it has left the versels, lymph undergoes congulation, fibrin being formed. The amount of fibrin which separates is between 0-4 and 0-8 per 1000, being less than that which separates from blood. Lymph is, in fact, simply while blood plasma; urea and carbonic acid, however, are rather more abundant in lymph than in blood.

Asamia. The chief change is a reduction in the number of red corpuscles, and the diminution of the amount of hemoglobin they contain. In severe cases there is also a reduction in the solid constituents of the plasma.

Leucocythamia. This is associated with great increase in the number of white corpuscles, which may become nearly as unmerous as the red. The blood is poor in hemoglohin, and rich in hypexanthine and lactic neid (Scherer). Charcot found in the blood, spleen, and liver of patients suffering from leucocythemia colourless elongated crystals, which he and Vulpian were inclined to consider as proteid in nature; while they were regarded by Salkowski as consisting of a mucin-like substance. Schreiner states that they consist of the phosphate of a base, to the hydrochloride of which he gives the formula C₂H₃N.HCl (A. 194, 68).

Gout. In this disease urio acid accumulates in the blood, probably owing to non-climination by the urine; there is also a large quantity of oxalic acid (Gurrod, Malico-Chirurg, Trans., 31, 83; 37, 51).

Rheumatism. The fibrin is much increased in amount: the same is, however, true for other inflammatory conditions. There is no excess of uric acid in the blood. Lactic acid is said to be the materies morbi by some, but this has never been satisfactorily demonstrated.

Fevers.—In various zymotic fevers, and septic diseases, the presence of different forms of bactoria has been described, or in some cases only presumed to exist in the blood. The best known of these are the spirillum of relapsing fever, the bacillus anthracis of splenic fever, and the bacillus anthracis of splenic fever, and the bacillus malarie of Klebs and Grudeli of intermittent fever. Pigment granules of a dark colour are also said to occur in the blood of ague patients. It is probably a derivative of hemoglobin (Marchiafava).

Diseases of the liver. In jaundice, bilirubin and in some cases the bile salts also accumulate in the blood; in a cute yellow atrophy the blood contains lencine and tyrosine.

Diabetes mellitus. In this disease the most marked feature is an increase in the amount of glucose in the blood. The peculiar odour of the breath in diabetics is stated to be due to acctone, and death is often said to result from acctonemia. It is probable that are down down not exist free in the blood, but is derived from the splitting up of accto-acctic other.

In some eases of diabetes a lipernic (fat in the blood) condition has been described. But there is no doubt that this may occur without evidence of disease, and also in other diseases than diabetes.

Bright's disease. In addition to an ambinic condition, there is an increase in the autount of nrea in the blood. The convulsions and coma that are apt to superview then the elimination of urea is defective have been designated evidences of uramic poisoning. It is probably however, that in these cases it is not ure itself which is the poison, but probably some arbestance or substances andecedent to urea. Frerichs' theory that the poison is ammonium sarbonate is now given up as untenable.

The Blood of Invertebrate Animals.

Our knowledge concerning the blood of invertebrate animals is much less complete than that of the vertebrates. In certain marine animals the circulating fluid is chiefly sen water in which a number of corpuscles are suspended (c.g. echinoderms) : in other invertebrates such as crustacea the blood is a highly organised fluid, and rich in profeid constituents, but even in these the amount of saline matters varies with the habitut, being much more abundant in marine than in fresh-water animals. There is never any distinction into blood proper and lymph in invertebrate animals; hence the name hamolymph is sometimes given to their encalating fluid; the term hydrolymph is applied in those cases in which the blood is chiefly water, and contains but few or anic constituents.

Hemoglobin is contained in the blood of many invertebrates (Lankester, Pyläger's Archie, 4, 315), chiefly worms, but also in a few crustaceans, insects, moliuses, leeches, and echinoderms. With the exception of four worms and two molluses, however, it does not occur in special corposeles as in the blood of vertebrates, but dissolved in the liquor sanguinis, colourless corpuseles only being found in the blood. In other invertebrates this red pigment is replaced by others, which apparently have a similar respiratory function; the most important of these other re piratory proteids are (1) brenocyanin, a libre pigment occurring in various crustaceuns, arachnids, and molluses (Frederica, C. R. 87, 996). This contains copper as one of its constituent characts; when oxidised it is blue, when reduced it is colourless. (2) Chlorocruorin, a green plament, closely related to humo lobin, found in the Idood of certain worms (Laukester, Journ. of Anat. and Physiol., 2, 114). 33411.cm rythrin, a purplish red pigment found in a few gephyrean worms (Krukenberg, Firyl. page. Studien, 1ste Reihe, Ste Abth. p. 82). In all these cases, the pigment is dissolved in the blood pla ma, which has thus a respiratory in addition to a nutritive function. In addition to these pigments, others occur which have apparently no respiratory function; thus chlorophyll appears in the blood plasma of many maths and butterflies (Poulton, Pr. 38, 269; tetronerythrin, a red lipochromo in the blood plasms of ceronin crustages (Halliburton. Journ. of Physiol., 6, 300). Various coloured granules are described in the corporcles of holothurians and sea urchius (Geddes, v. Gamgee's Physiol. Chem., 131), and the blood of the limpet is de cribed by Krakenberg as being of an orange colour.

The blood of most invertebrates is alkaline in reaction, the only known exception being that of mostly and butterflies which is acid (Poulton).

With regard to the coagulation in the blood of invertebrates Haliftenten (Journ. of Physiol., 6, 390) was able, in the case of the crustacea, to separate crustacean filtrin, and to show that as in vertebrate blood in was formed from a previously solube fibring; a nuder the influence of a ferment. Crustacean fibrinogen and fibrin differ but little from that of vertebrate blood and the fibrin learnest is identical with that

obtained by Schmidt from vertebrate blood. The congulation of crustacean blood is also hindered by cold, or admixture with neutral salts. The coagulum formed when the blood of cephalopods is shed is stated by Fredericq to be only a plasmodium of cells.

(The foregoing article has only discussed blood from a general point of view. The various constituents will be described under their proper headings; the proteids including hæmoglobin and the other blood pigments will be described under the heading Proteids.] W. D. ff.

BLOWPIPE v. ANALYBIN. BOHEIC ACID C, If 10 Ob. 1100°1. ()ccurs (to the extent of 2 p.c.), together with quercitaunic acid, in black ten (Theu bohen) (Rochleder, A. 63, 202). Yellow amorphous resin, v. sol, water and alcohol; ppd, by alcoholic or ammoniacal leaf acetate. BaA"aq. - PbA"aq. -PbA"PbO.

BOILINO-POINTS v. PHYSICAL METHODS; SOC. TREEMAL.

BOLDIN C₃₀H₅₂O₈. A glucoside which may be extracted by boiling alcohol from the leaves of Boldon fragrans, in which it occurs to the extent of 3 p.c. It is a syrup, volatile with steam, and decomposed by hot dilute HCl into glucose, McCl, and an oil C18H2O3 (?) sol. aleohol, insol. water (Chapoteunt, C. R. 98, 1052). According to Bourgoin a. Verne (Bl. [2] 18, 481) the leaves of Bolden contain an alkaloid,

BOLETUS P. AGAIGUES.

BONE OIL (Anderson, Tr. E. 16, 4; 20, ii. 217; 21, i. 219; 21, iv. 571; A. 70, 32; 80, 44; 94, 358; 105, 335). The following substances have been isolated from the tar obtained in the dry-distillation of bones:

By-products.

Methylanauo

Methyl-pyridine

Di-methyl-pyridino

Anilino

Pyridino

Quinoline

Propionitrile

Ethyl benzena

Naphthalene

Valeramide

Phenol

Tolucue

Chief Constituents.

Butyronitrile Valeronitrile Hexmitrile Isohexonitrilo Deconitrile Palmitonitrile Stearonitrilo Pyrrol. Methylpyrrol Dimethylpyrrol Hydrocarbons:

C.H. (Dihydra m-ethyl tolucne?) $C_{\alpha}H_{1a}$

C_{irth} (Dihydro m-methyl-cumene ?)

Weidel a. Cimuician (B. 13, 65) consider that the aitriles are formed by the action of NII, at the high temperature on the fatty acids contained in the hones; the pyridine bases they believe to be formed by the combination of the acrolein (from the glycerin in the fats) with ammonia, methylamine, &c., whilst pyrrol and its homologues are products of the decomposition of gelatin.

BORATES. Salts of boric acid r. Borox. OTTACHS OF, p. 528.

BORAX v. Borate of sodium : under Boron, SETACIDS OF, p. 529.

BORIC ACID v. Boron, oxyacids or, p. 528.

BORIC ANHYDRIDE v. Boson, oxide of, p. 521.

BORIDES. Compounds of boron with one other more positive element.—Very few of these compounds exist; manganese forms a crystalline borido probably Mn,B2; platinum easily combines with boron to form PtB (?); and aluminium and boron appear to form a kind of alloy, the proportion of the elements in which varies within very wido limits: two definite borides of Al are also known (v. Iaidion, fron, Manganese, Palladium, Platinum, Alominium, BORIDES OF)

BORNÉENE. A mixture of terpones (q. v.), exuding from Dryobalanops camphora, holding

borneol in solution (Gerliardt, Traite, 3, 628, 641).

BORNEOL C10 H 10 O i.e. C10 II 1, OII. Borneo camphor, Tetra-hydride of (1, 4, 6)-methyl-propyl-phenol (2). Mol. w. 154, [1989] (P.); [2079] (W.) (2122). R $_{\odot}$ 75:30 in a 22:5 p.o. alcoholic solution (Kanonnikoff).

Occurrence. - In Dryobalanops camphora, being extracted from hollow cavities in the trunk of old trees (Pelouzo, A. 40, 326). In the essential oil of valorian (Gerhardt, A. 45, 34; Bruylants, B. 11, 451). To the extent of 4 or 5 p.c. in oil of rosemary (Bruylauts, J. 1879, 944; Wober, A. 238, 89).

Formation .- 1. By heating camphor with alcoholic KOH (Berthelot, A. Ch. [3] 56, 78). -2. By the action of sodium on camphor (Baubigny, Z. [2] 3,71; Haller, C. R. 105, 227).

Preparation from camphor .- (Jackson a. Menke, Am. 5, 270; 6, 401; Knchlor a. Spitzer, M. 5, 50; B. 15, 16, 2730; Immenderif, B. 17. 1036). Camphor (50g.) is dissolved in alcohol (500 c.c. of 95 p.c.), and sodium (60g.) added slowly. Towards the end of the operation water (50 c.o.) is ndded (O. Wallach, A. 230, 225).

Properties .- Regular crystals; very readily sublines in plates. Smells like camphor, but more peppery. flas a burning taste. nleoholic solution is dextrorotatory; artificial borucol has a somewhat higher rotatory power (c. 43°) than the natural horneol $([a]_0 = 33^\circ)$ (Biot; Kachler). V. sl. sol. water, scl. alcohol and other. Lighter than water. The rate of etherification of borneol resembles that of primary alcohols (Menschutkin, J. R. 13, 162).

Reactions.—1. P.O. converts it into one or more terpence (borneene).—2. Boiling HNO. (S.G. 142) gives camphor and its oxidation products. 3. Behaves as an nlcohol with regard to PCl .- 4. HClO converts it into camphor.

Sodium borneol C_{1,1}H₁.ONa: six-sided plates (from benzene). Combines with CO₂, forming C10H1,O(CO2Na) (Kachler a. Spitzer, M. 2, 235).

Bromide.-Bromine added to an ethereal solution of borfeed forms crystals of a mixture of bromides (? CiallysOBr. mid (CiaHisO)zBrz).

Hydrobromide. (C.II.O).HBr. Crystal-line pp. got by passing fIBr into a solution of borneol in light petroloum. The compound is unstable and is decomposed both by water and by alcohol.

Hydriodide. (C10H10) Hl. Prepared similarly.

Methyl derivative C. H1: OMo. (194° cor.). From sodium-borneol and MeI (Baubigny, Z. 1868, 299).

Ethyl derivative O₁₆H₁₇OEt (202°) (B.). Formyl derivative O₁₆H₁₇O.CHO. (225° 230°). In oil of valerian (Bruylants).

Acetyl derivative O,H,OAe. K. a. S.); (227°) (M.). Occurs in oil of valerian (Bruylants, B. 11, 456); and may be formed by the action of Ac.O upon borneol (Montgoltier, A. Ch. [5] 14, 50), or of AgOAc upon bornyl chloride (Kachler a. Spitzer, A. 200, 352). On standing it becomes crystalline [24]. Fusion with NaOH gives NaOAc and borneol.

Isovaleryl derivative C, H, O(C, H,O).

(255°-260°).

Stearyl derivative C10H1.O(C18H2.O). From borncol and stearic acid at 200 (Berthelot, A. 112, 366). Oil.

Benzoyt derivative C10 II 11 OBz. Oil. Leeverctatory berneel C₁₀H₁₇OH, [35°], (210°). (Perrot, A. 105, 67). $[a]_0 = -33^{\circ}$. Occurs in the alcohol produced by fermentation of the sugar of madder-root (Jeanjean, A. 101, 95). Small regular crystals; sl. sol. water, rotating upon it.

HNO, forms hevorotat ry camphor.

Levorotatory berneol C₁₀H₁OII. [204°]. S.G. 1.02. From Ngai camphor (Hanbury, J. 1874, 537).

Laworotatory berneel $C_{10}H_{11}OH$. [201°]. [a]₀ = -37° 21′ in alcohol of 82 p.c. at 22°. From thymene picrate and boiling NaOHAq (Lextreit, J. Ph. [5] 13, 265). HNO, converts it into a hevorotatory camphor [176:] (204°).

Leveretatory berneel C, 11, OII. Formed. together with ordinary borncol, by the action of Na on dextro or lavorotatory camphor (Montgolfier, A. Ch. [5] 14, 21; C. R. 89, 101).

Inactive bornecl C₁₀II₁₀O. [1992]. Among the products of the distillation of colophene (q. v.). Also from its acetyl derivative.

Properties. — Differs from dextrorotatory borneol only in being inactive. The crystal's float on water, but when pressed into reolid; eake they sink (unlike camphor). Oxidised by HNO, to inactive camphor (Armstrong a. Tilden, J. 35, 752). Heated with a large quantity of HCl it forms C10H1, HCl.

Acctyl derivative C, II, OAc. (2150). From terebene and HOAc at 100 (Bouchardat a. Lafont, Bl. [2] 45, 164; C. R. 102, 171).

According to Haller there are two true borneols $[a]_j = +$ or -37, and the others are molecular compounds of these (v. Campon and

BORNYLAMINE C10 H111 N probably CH₂ $\alpha_0 = -18^{\circ} 35' 41''$, [160°]. [160°]. (200°). V.D. 5.5 (for

Formation .- I. By reduction of camphoroxim in alcoholic solution by means of sodiume 2. By saponification of its formyl derivative obtained by heating camphor with ammonium formate at 220 '-210'.

Properties.-Crystalline solid, having as odour resembling both camphor and piperidine. In its physical properties it greatly resembles camphor. Very volatile with steam. Sublinable even at the ordinary temperature. V. sol. alco. hol, ether, &c., nearly insol. water. Alkaline reaction to litmus. Takes up CO, from the air. Primary base. Gives the carbamine reaction. Leevorotatory.

Salts.—B'HCl: easily soluble white needles [c. 280°].—B',Cl,H,PtCl,: golden-yellow plates, v. sol. hot water or alcohol.—B'H,SO,; easily soluble rhombio tables. -B', H, Cl, H, Cl

Formyl derivative C.H. NHCHO; [61°]; colourless glistening plates.

Acetyl derivative Coll..NHAc: [1410] colourless plates.

Benzoyl derivative C10II,.NIBz: [131°]; colonriess plates; insol. water and cold ligroin (Leuchart a. Bach, B. 20, 104).

BORNYL BROMIDE C, 11, Br. [75°]. From borneol and HBr (Kachler, A. 197, 98).

BORNYL CARBAMATE Call, NO. i.e. CteH,O.CONH., [115°]. From sodium borneol in toluene by the action of eyanogen (Haller, C. R. 93, 1511; 91, 869). Monoclinic needles (containing aq.). Sl. sul. hot water; sublimes partially at 100°. Dextropolatory based in the sublimes are sublimes. Dextrorotatory. Benzoia aldehyde and HCl form CHPh(NH.CO.O.C, 1111)

[187°]. Cf. Centol.

BORNYL CARBONATE (C_oH₁₇),CO., [215°].

Extracted by boiling alcohol from the residue left in the preparation of eynno-burneol from sodium borneol and cyanogen (Haller, C. R. 91, 86). White plates or hexagonal tables, insol. water and alkalis, sl. sol, cold alcohol, sol, ether, May be sublimed. The rotatory power varies with that of the borneol from which it is prepared. Boiling alcoholic KOH gives K.CO, and borneot.

BORNYL-CARBONIC ACID C1111, O.CO211. Borneol-carboxylic acid. From sodium borneol and CO₂ (Baubigny, Z. 1868, 299; Kachler a. Spitzer, M. 2, 236; C. C. 1881, 359). -NaA': crystalline, v. sol. water; slowly decomposed by water with separation of borneol.

BORNYL CHLORIDE C. II, Cl. [1570 Formation. -From borneof (1 pt.) and HCI

(9 pts.) at 100° (Berthelot, A. 112, 366).

Preparation. - From PCl, (60g.), light petrofeum, and herneol (15g. added in portions of 6g.), The product is shaken with water and the petrolemn allowed to evaporate in the cold, where bornyl chloride separates (Wallach, A. 230, 231; Kachler, 4, 197, 93; B. 11, 460).

Properties. Crystals. V. sol. light petroleum, m. sol. alcohol. Lievorotatory

Resections. 1. Converted into HCl and camphene C₁₀H₁₆ [52] (c. 160°), by heating with water (40 pts.) at 95° (Kachler, A. 197, 96); better by warming with audine (W.). A little borneol is also formed by the action of water on bornyl chloride (Kachler, A. 200, 312; Rihan, A. Ch. (5, 6, 382). 2. Sodigm acting on a solution in benzene forms gamphene CmlI, and hydro-

camphene C_1,11,... BORNYL-METHYL-UREA

C. II. NII.CO.NIIMe. 200 Formed by the action of methyl evanuate upon bornylamine in chereal solution. Plates. V. sol. ether and hot water (Lenchart a. Bach, B. 20, 108).

BORNYL OXIDE C., H ., O i.e. (C., H.,), O (?). (285°-290°). Occurs in the escential oil of valerian (Bruylants, B. 11, 456). Not attacked by melted KOIL

BORNYL PHENYL-CARBAMATE

oc< Bornyl - phenyl - urethane. [133°]. Formed by the action of phenyl cyanate Isomeric with camphylamine. upon borneol. Needles. Sparingly sol. cold ligroin and alcohol, v. sol. other solvents | molten mass is stirred with an iron rod, and (Leuchart, B. 20, 115).

BORNYL-PHENYL-THIOUREA

C₁₀H₁₇.NH.CS.NHC,H,. [170']. Formed by the action of phenyl mustard-oil upon bornylamine in ethereal schrion. Colourless needles. Nearly insol. ligroin (Lenchart a. Bach, B. 20, 1091.

BORNYL-PHENYL-UREA

CeHt, NH.CO.NHC II. (218). Formed by the action of phenyl symmetre upon bornylamine in ethereal solution. Silvery plates or line medles. Sparingly soluble in other and cold alcohol, easily in hot alcohol, insoluble in water (Leuchart a. Bach, B. 20, 168).

BORNYL UREA Coll CNILCONIL 164 Formed by boiling bornylamine hydrochloride with potassium cyanate. Colourless needles, Easily soluble in hot water and alcohol (Lau-

chart a. Buch, B. 20, 108).

BORNESITE C 11, Os. [175]. [a]0 - 32°.

Methyl-dambore. A volatile substance occurring in the caoutehous of Bornea. Sublimes at 205", Sweet taste; does not ferment. After boiling with dilute acids it reduces Felding's solution. At 120 it is split up by funding 111 into Mel and da abose (Giracel, U. R. 73, 126),

BoxOFLUORIDES, v. under Boxon, perforibe

or, j. 526.

BORON. B. At. w. 10-97. Mol. w. unknown, as V.D. bus not been determined, S.G. amorphous not determined, but greater than 18t eryst. 253 268 (Wöhler, A. 141, 268; Hanpe, A. 183, 75). S.H. about 37 at 250', probably about 5 at 1000' (v. post; p. 525). Grystallises in dimetric forms, a:c = 1: 5762 (Sella, P. 100, 616); but crystals probably contained C and Al (v, post). S.V.S. about 4-1. Combines directly with D and Cl with production of much heat: [B',O'] - 317,200; [B,Cl'] - 104,000 (Troost a. Hautefeuille, 41 (2), [5] 9, 70). Chief lines in emission spectrum are 2496 2, 2197, 3150 1 (Hartley, T. 175, 49).

Occurrence .- Not us boron; chiefly us horax and horie acid in volcanie districts, also as borate of Mg with MgCl, (Banwite), as bornto of Ca with Ca silicate (Datolike), Ac. Borax, or tincal, has been known in commerce for many centuries; borie acid was prepared from borax in 1702 by Homberg; the element was obtained by Clay-Lassac and Thémard in 1808 by de-oxidising baric and by petassium, which metal had been obtained by Davy the year before.

Formation, . 1. By reduction of B,O, by K Formation. 1. By reduction of B₂O₃ by K (Gay Lussac a. Thénard, G. J. 30, 363), -2. By reduction of BF₂KF by K (Bergelius, P. 2, 113), 3. By electrolysis of fined BO₄ (Pavy, G. A. 36, 440), -4. By reduction of BO₄ by H at a red leat (Dunna, A. Ch. 3), 37(6), -5. By fusing dry horax with anouphous P (Pragemborf, C.C. 1991), and 1998. 1801, 865). - 6. by heating BF, KF or BF, NaF with Mg (Wöhler a. Deville, A. Ch. [3] 52, 62; Genther, J. Z. 2, 200). 7. By heating B.O. with Mg and treating the product with HClAq (Jones, C. J. 35, 42).

Preparation. Amountous, 10 parts fuscil B.O. in coarse powder are mixed with 6 parts Na in small pieces, the mixture is placed in an iron crucible bested to full reduces, 4 to 5 parts of fused NaCl are added, and the crucible is

the contents of the crueible, while still hot, are poured into water containing a little HCl. The NaCl, berax, and B₂O₂, dissolve, and the boron remains. The boron is washed with very dilute HClAq, then with alcohol, and then with ether; it is then dried at a very gentle heat (Wöhler a. Deville, A. 101, 113 a. 347; 105, 67).

CHYSTALLINE. Amorphous B is pressed as tightly as possible into a small Hessian crucible. a hole is then made in the mass and a rod of At (4-6 grams) is placed in the hole: the crucible is covered and placed in another, larger, covered, crucible; the space between is filled with powdered charcoal, and the crucibles are heated to 1500° or 1600° for 11 to 2 hours; after cooling the mass is treated with dilute HClAq which dissolves Al, and BN formed in the process (Wöhler a. Deville, A. 105, 67). According to Hampe (A. 183, 75) the crystals obtained by this process, or by any process said to yield crystalline B, contain Al, and some of them also C; Hampe gives the formula AlBis to the black crystals, and CAl, B, to the reddish yellow crystals, obtained by the foregoing method (c. Aleminium, nomines or; and

ALUMINIUM, BOROCARHIDE OF).

Properties. - Amorphous boron is a greenishbrown, opaque powder; tasteless; odonrless; non-conductor of electricity; very infusible, but melts when placed between the poles of a battery of 600 Bursen rells. Said to be slightly soluble in water; Reinitzer (Sitz. W. 82, 736) supposes that the body which dissolves is a hydride of B (v. Bonos, uypning or). Insoluble in alcohol or other. Heated in vacuo or in an martive gas, e.g. II, B becomes darker in colonr, heavier, and more compact. Heated in O, burns to B_2O_0 $(B_2^*O^*) = 317,200$: heated in air, B_2O_0 and BN are produced. Oxidised by heating with KNO, R.CO, KOH, cone. HNO, Aq, or aqua regia. Combines directly with many elements e.g. S. Cl. Br. N. The properties assigned by Wöhler u. Deville und others to crystalline B. are, according to Hampe's experiments, the properties of AlB, AlB, and CAlB, (v. ALUMINIUM, ROBIDE OF, AND BOROCARBIDE OF). The atom of B is trivalent in gaseous molecules (data BCls, BBrs, BFs). The atomic weight of B has been determined (i) by finding the V.D. of BCl, and BBr, and estimating the Cl and Br respectively in these compounds (Deville a. Dunns, A. Ch. [3]55, 180); (ii) by dehydrating boray (Berzelius, P. 2, 129; 8, 19); (iii) by converting dehydrated borax into Na SO, by action of HFAq and H₂SO₄Aq (Berzelius, P. 2, 128; also Artvolon, P. 2, 127); (iv) by determining the S. H. of Laron (r. infra). Boron is a uon-metallic element in its chemical reactions; its oxide, B_iO_o in an anhydride; boric acid, H.B.O. corresponds in composition to nitrons neid, but thermal data show that boric acid is dibasic; the acids H.BO, and H.B.O. (and several salts derived from the latter) are also known (v. Ponen, extractes or). B.O. seems to form compounds with P.O. SO. and WO. In some respects It shows analogies with C and Si :- physical properties; existence of acid containing F (HBF.); direct combination with N; existence of many borotungstates; probable existence of a sovered. When all action has coased the gaseous hydride, &c. In many points B resem-

position of compounds (B₂O₂, H₂B₂O₃, BCl₂, BCCl₃, &c.); B₂S₂, not a salt-forming sulphide in reactions with sulphides of very positive metals; existence of a B analogue of tartar emetio, &c. Boron is the first member of Group III (periodic law); the succeeding members of this group are all decidedly metallic; but Al2O, 3H,O dissolves in KOHAq, and forms aluminates (q, v,); Al.S₁ is also somewhat analogous to B_2S_1 : the compositions of several B compounds are similar to those of the corresponding compounds of members of Group III., e.g. M.O., MX, [X=Cl., Br., AlCl., and BCl., combine directly with PDCl., The differences between the chemical functions of B and the other elements of the group to which B belongs seem to be wider than is usual between the first and the following elements of the same group. The boron group comprises the following:

These elements are all metallic with the exeeption of B; in the reactions of B_iO_i towards certain acids (p. 527) It shows that it may act as a feely metal-like element. Borie acid is an extremely weak acid; its affinity is very small. The borates are very mustable salts, easily decomposed, even by water, to boric acid and basic oxides. The last member of the group, TI, shows distinct analogies with the 11th series member of the next group, viz. Pb; B shows distinct analogies with the 2nd series member of the next group, viz. C. B occurs in Series 2, all the succeeding members of this series -C, N, O, F ... are very negative and non-metallic; the general character of the series to which it belongs is stamped upon B, and the group-character is but feebly marked. It must, however, be remembered that very few compounds of B, except the horates and their derivatives, have been fully atudied.

Specific heat. The S.H. of B as determined by Kopp, Reguault, and Mixter a. Dana (A. 126, 362; Suppl. 3, 1, 289; J. 1861. 29; A. 169, 388) varied from 225 to 262 for the temperature-interval 30°-70°. In 1873-4 Weber car-fully determined the S.H. of crystallised boron (v. P. M. [4] 49, 161, 276); the following table summarises his results:

S.H. of crystallised boron. - 40° > H. 8.H. × At w. 1915 2.11

+77 ·2737 8.01 177 8.72 .3663 4.03

The S.H. increases as temperature rises, but the rate of increase per 1° is nough smaller at high than at low temperatures. The variations in the rate of increase are almost identical with those observed in the case of carbon (q. v.); assuming that this identity remains at tem. peratures above 233°, the value which the S.H. of crystallised boron will attain at about 1000° is approximately 5. Weller did not analyse the crystals of boron used; they were

ing to Hampe's investigation crystals thus prepared are a definite compound of B and Al (ante, p. 524).

. Reactions .- 1. Heated in air, B.O. and BN are formed. - 2. Heated in oxygen, B burns to BrO2.-3. Heated to redness in N. BN is formed, 4. B combines directly with many elements, e.g. Cl. Br, S, and some metals (v. Bonnes).— 59 Water is not decomposed by B at 110 but at a red heat. Steam reacts with B to form horie seid and H. 6. B is oxidiated by heating with nitric acid, come, valpharic and, or aqua regia; or by the action of molten nitre, or various oxides of heavy metals. 7. B is also oxidised by heating with petash (II is evolved). or with alkaline conbonates (C is separated, Berzelius, P. 8, 19), or with phosphoric acid (P is separated, Wohler n. Deville, A-Ch. [3] 52, 63) .- 8. B burns when heated in nitric oxids forming BaO, and BN: NoO is without action on B.-9. At a red heat B decomposes subplimetted hydrogen, hydrogen chloride, and ammonia, farming respectively B.S. and H. BCl, and R. BN and H. 10. Many metallic chlorides and sulphides, e.g. PbCL, AgC9, PbS, are reduced to no tal when heated with B. 11. From aqueous solutions of gold change B pp. Au. - 12. Aqueous solutions of caustic alkalis do not react with B. 13. So called crystalline boron reacts similarly to amorphous boron, but the reactions occur only at high temperatures; it is oxidised with much difficulty.

References. Gay Lussue a. Thenard, G. A. 30, 363; Davy, G. A. 35, 440; Berzelins, P. 2, 113; Wöhler a. Deville, A. 101, 113; 103, 347;

Boron, Bromido of, Blir., Mol. w. 250-22, (90-5) (Wöhler a. Deville, A. Ch. [3] 52, 89). S.G. 269 (W. a. D. Lc.). V. D. 127,

Formation. 1. By action of Br on B.O. and C at red leat (Poggiale, C. R. 22, 127). 2. By heating I O, with PBr, (Gustavson, B. 2,

Preparation. ... Amorphous B is loosely packed into a glass tube, the tube is gently watered and H is passed through it until every trace of moisture is removed; the H stream is stopped, the corks of the tube are removed for a moment or two, and then the B is gently heated in a stream of dry Br vapour, and the liquid BBr, is led into a dry flask surrounded by ice cold water. The BBr, is freed from Br by digestion with Hg, and distillation.

Properties and Reactions. -- Colourless, strongly luming, liquid. Reacts with H₂O to form HBrAq and H,BO,Aq. Forms a compound with dry NH, (r. also Nickles, C. R. 60, 800;

tiautier, C. R. 63, 920).

Boron, Chloride of. BCl,. Mol. w. 117-08. (15°23) at 760 mm. (10 gna. d. Acad. 26, 658), S.G. 11 1:35 (Wöhler a. Deville, A. Ch. [3] 52. 63). V. D. 58 2. H. F. (B, CF, = 101,000 (Troost n. Hautefenille, C. R. 70, 185).

Formation.—1. By direct combination of B and Cl (Berzelius, P. 2, 147).—2. By the reaction of Cl with B₂O₂ and C at a red heat (Dumas, A. Ch. [2] 31, 436; 33, 376).-3 By the reaction of B with dry HCL-4. By reaction between HgCl, PbCl, or AgCl, and amorphous B.-6. By heating B.O. with PCI, to 150° for prepared by heating borio acid with Al. Accord- some days (Gustavson, B. 3, 426; 4, 975).

Preparation.—Amorphous B is heated in H until quite dry, then in dry Cl (details v. Bonon, BROWIDE OF); the exit-end of the tobe is connected with a Y tube, the upper part of which is surrounded with a mixture of snow and salt, and the lower limb passes into a dry tube also surrounded by snow and sait.

Properties. Colourhas, highly refractive, liquid. Very expansible by heat. Funce in

air with decomposition.

Reactions. - 1. With water forms ff, BO, Aq and ffClAq; (BCP, Aq) 79,200 (Troost a. Hautefeuille, A. Ch. [5] 9, 70). - 2. Not decomposed by heating with zinc-dust to 200°; or with sodium below 150°, at 150° B is separated. 3. Heated for some time with superficially oxidised sodium-amalgam at 150 'B is separated (Gustavson, B. 3, 426). 4. Heated with P.O., the compound BCl., POCl, is formed (v. infra) (Gustavson, B. 3, 975). . 5. Heated with sulphur trioride, B.O. and SO Cl. are formed (Gustavson, I.c.) .- 6. With alcohol forms B(OEt), and HCl. -7. With NO, react; to produce BCl, NOCI, B.O., and O (Genther, J. pr. [2] 8, 853).

Combinations. 1. BCl, vapour passed into POCl, forms crystals of BCl, POCl, (Gustavson, B. 4, 975). This compound melts at 73' in a closed tube; by sublimation it separates into BCl, aml POCh; it is decomposed by water, or moist air, into H₁PO₄Aq, H₂BO₃Aq, and HClAq. The same compound in produced by the reaction of BCl₂ with P₂O₄, and of B₂O₄ with POCl₃ (G. Le.). --2. With animonia gas gives 2BCl, 3NH, with production of much heat (Berzelius, P. 2, 117). 3. The compounds BCl, CNCI, aml BCl, CN11, are known (Martins, A. 109, 80; Gantier, C. R. 63, 920).

Boron, Flueride of. BF., Mol. w. 68-27.

V. D. 33.7. S. (0°) 1013.

Formation. -1. By reaction, at white heat, of an intimate mixture of 1 part B.O. with 2 parts Col., free from silica; Gay Lussac a. Thenard (A. Ch. 69, 204) prepared BF, by this method, in 1810, using vessels of thirt, and collecting the gas over Hg. -2. By heating 1 part B₁O₂ with 2 parts CuF₂ and 12 parts cone. H₂SO₂A₁ in glass vessels (J. Davy, T. 1812, 365); or 1 part B₂O₃, f part CuF₂, and 20 parts H₂SO₂ (Ferrari, J. Ph. 19, 48). Prepared thus, tho gas always contains SiF, (Berzelius, P. 2, 116).

Preparation. - 100 parts KBF, are mixed with 15 20 parts fused and linely powdered B.O., and the mixture is hented with cone. ILSO, the gus is collected over fIg (Schiff, A. Suppl.

5, 172).

Properties .- Colourless gas, with suffocuting odonr, condensed to a limpid liquid at -110° and strong pressure (Faraday, 4, 56, 152). Acts on organio untter like conc. If SO;; incombustible; not decomposed by electric sparks; does not act on ginss; is very stable, not decomposed by Fc at red heat.

Benetions, t. with water forms horofluor-hydric acid HBF Aq (v. infra), or fluoboric

acid HBO, 311F (r. p. 630), according as the BV, is passed into water until the reaction is acid, or until the water is saturated (c. onfor). [BF*, Aq] = 24,510 (Hammerl, C. R. 90, 312). -2. Conc. sulphuric acid absorbs BF, (about 50

earth, metals react at red heat, forming boro-fluorides and B (Berzelius, P. 2, 188).—4. With alcohol boric soid and ether are formed.

Combinations. - With ammonia to form BF, NH, a solid body not decomposed by sublimation; also BF, 2NH, and BF, 3NH, liquids, decomposed by heat, by exposure to air, or by dry CO2, giving N.I, and BF3.NfI, (J. Davy, T. 1812, 368). According to Kuhlmann (A. 39, 320) BF, also combines with the oxides of N.

BOROFLUORIEDEMO ACID, AND BOHOFLUORIDES. IIBF, MBF, (Hydroftuoboric acid.) When BF, is led into water until the liquid shows a strongly acid reaction, and the liquid is cooled, of the B of the BF, separates as ILB,O, and the rest remains in solution in combination with H and F. By neutralising the liquid with KOHAq, and evaporating, a salt having the composition KBF, is obtained. If the acid liquid is evaporated HF is evolved, and ILB O. 611F remains in solution (v. PLUOBORIO acto nuder Boson, oxyactos or). According to Landolph (C. R. 86, 603) the seid HBF, may be obtained us a colourless liquid, boiling at 1302 with partial decomposition, by the reaction between BF, and unethol (C.H.,OCII, C.H.); the acid renets with a little water to produce HFAq and IIBO,Aq. A solution of IIBF, is also obtained by dissolving crystallised borie acid in dilute, cooled, HFAq.

Вовоги токтыем. These salts have been chiefly studied by Berzelius (P. 2, 113). They are obtained by the reactions between (1) metallic oxides or carbonates and HBF, Aq. (2) BF, or HBO23HFAq and metallic fluorides, (3) HFAq and metallic thorides mixed with HBO.; in the last case bull the notal of the fluoride usually forms an oxide. Most borofluorides are crystalling, soluble in water, decomposed by heat to Bl', and metallic fluoride; heated with cong. H.SO₄, BF₃, HBF₄Aq, and notallic sulphile, are produced; fused with alkali carbonates they form a mixture of alkali metal fluoride and alkali borate, this reaction affords the basis of a method for analysing the borofluerides (r. Marignac, Fr. t. 105). Many berofluorides are partially decomposed by water lorming so-called basic salts, e.g. Ba(BF), $2H_2O_2$ Ca(BF), fb(BF), some e.g. the Ba or Ph sait are partially decomposed by alcohol; the aqueous solutions of several -c.g. NH, BF, Ca(BF_i) - redden blue lituus.

Totassium borofluoride Kill Obtained as a gelatinous pp. by adding HBF Aq to a soluble K salt. Prepared by Stolba (C. C. 1872, 395) K satt. Trepared by Stobia (C. C. 1872, 305) by henting to boiling I part crystallised boric acid, 2, parts powdered CaR, and 5, parts cono. ILSO, cooling, filtering, ppg. KBP, by addition of a soluble K salt, crystallising from tot water—S. (cold) 70.—The salt forms white lustrous six-sided tables; may be crystallised from algoliol or alkali carbonate solutions; notice when beated and at a bight temperature. melts when heated, and at a high temperature

decomposes to BF₂ and KF.

The other borofluorides are Al₂(BF₄),
Nf1,BF₀ Ba(BF₄),
Pb(BF₄),
LiBF₀ Mg(BF₄),
NaBF₀ Y₂(BF₄), Zu(BF,),

Boron hydride. No hydride of B has been vols.), on adding water boric acid is ppd. obtained free from H; but the experiments of (J. Davy, T. 1812, 365).—8. Alkalı, and alkalı: Jones (C. J. 35, 41), and of Jones a. Taylor (C. J. Me, \$13), leave little doubt that a gaseous 5. By heating the compound of BCl, with NH, Ht hydride exists and show that its composition is to 200° (Gustavson, S. [3] 6, 831).

probably BH.

Preparation .- An intimate mixture of 1 part recently heated B.O. and 2 parts Mg dust is placed in a Hessian or iron crucible, the lid is firmly wired down, and the crucible is heated in an ordinary fire; a violent reaction occurs, the oracible is at once removed from the fire; the fused mass—a mixture of B, Mg, Mg, Bg, Mg, Ng, and MgO—is placed in a small flask along with a little H O, and conc. HClAq is allowed to drop into the flask from a stoppsred lunnel tube; the gas is collected over water, or is dried by CaCl, and collected over Hg. Mg.B. may also be prepared by direct combination of B and Mg, or by heating Mg in BCl, vapour (J. a. T., C. J. 39, 214). The gas consists of II mixed with a very small quantity of B hydride.

Analysis. A known volume of pure II was burnt, by hot CuO, to II,O, and the H,O was weighed; an equal volume of the gas prepared as above was lornt in the same apparatus, and the H.O was weighed; the excess of H.O in the second experiment over that in the first gave a measure of the II combined with B as B hydride. The results showed the composition of the hydride to be BH, where x>2 and is ap-

proximately ... 3.

Properties and Reactions .- Colourless, very disagreeable odour, sparingly soluble in water, solution seems to be unchanged on keeping. Gas burns with bright green thame producing ILO and B.O. Decomposed by passing through a hot tube to B and H. Reacts with AgNO, Aq producing small quantity of a black pp. containing B and Ag, and decomposed by H O giving B hydride. Beacts with K.Mn.O.Aq giving MnO, and H.BO.Aq. Combines with NH, (conc. N11, Aq) to form a crystalline compound, of unknown composition, decomposed by acids.

References .- Older attempts to prepare bydride of Loron: Wöhler a. Deville (A. Ch. [3] 52, 88); Genther (J. 1865, 125); Gustavson (Z. 1870, 521) (r. also Reinitzer, Sitz. W. 82, 736). Compounds of B with paraffin radicles are known, eg. BEt, (r. Bohon, Organic Deriva-TIVES OF).

Boron, Hydroxides ol, v. Bonon, expactes OF.

Boron, Iodids of, Not known. Wöhler a. Dewille (A. Ch. 13' 52, 90), by the action of I on B at a high temperature, obtained a body which thew regarded as an oxynolide. Agl does not react with B even at the melting point of Ag.

Boron, Nitrido of. BN. Mol. w. unknown, as compound has not been gasified. Obtained in 1842 by Balmain by melting B.O. with KCN (P.M. (3) 21, 170; 22, 467; 23, 71; 21, 191) Composition determined by Wehler in 1850 (A. 74, 70).

Formation .- 1. By heating B in N, or in NH1 -2. By heating to whiteness a mixture of 4 parts B.O. and 1 part charcoal powder in N. 3. By heating borax (Wöhler, I.c.), or borne acid (H. Rose, P. 80, 265), with NII,Cl, or K, I'eCy. (W. Lc.), or KCN, or Hg(CN), or rea (Darmstadt, A. 151, 255).-4. By heating 2BCl, 3NH, and passing the vapour, along with NH, through a hot tube (Martius, A. 109, 80).

Preparation. A mixture of 1 part dehy-drated borax and 2 parts NH,Cl-or 7 parts B.O. with 9 parts area (Darmstadt, Lc.)-is strougly heated in a covered Pt orneible, the finely powdered mass is boiled with much water containing a little HCl, washed with hot water, B O₄ is removed by eareful treatment with HFAq (Wöhler, Le., could not remove all B₄O₄ thus), and the BN is washed and dried.

Properties. White, light, amorphous, powder; insoluble in water; infusible; solt (like tale) to the touch; heated in the edge of a tlame exhibits greenish white phosphorescence; very stable and very slightly acted on by most rengeats, e.g. by heating in air, O. I. H, COp or CS, or with cone. HClAq or HNO, Aq or ROHAq.

Reactions. - I. At a very high temperature reacts with chlorine, to give BCI, (Durmstadt, A. 151, 255). - 2. Hented to redness in steam, or to 200° in a closed tube with water, NII, and H.BO, are formed .- 3. With molten potash, NII, and K borate are produced. 4. With molten potassium carbonate KCNO and KBO, are formed, if much BN is used KCN is also produced. 5. Oxides of Pb, Cu, or Hg are reduced by heating with BN, with formation of NO or N.O. (Wöhler, A. 74, 70).-6. Heated with conc. sulphurie acid, or with cone. hydrochloric acid to 200° in closed tubes, NH, and H.BO, are formed. - 7. With cono. hydrofluoric acid NILBF, is formed .- 8. Heated in an alcoholflame fed with oxygen, BN burns to B.O.,

Boron, Oxide of. B.O., (Boric anhydride.) Mol. w. niknown, as compound has not been gasified. [577] (Carnelley, C. J. 33, 278). S.O. 1.75 1.83 (Playfair a. Joule, C. S. Mem. 3, 57; r. also Ditte, A. Ch. (5) 13, 67). S.H. (16° 98°) [B] O³ | 317,200 (Troost a. Hautefenille, A. Ch.
 [5] 9, 70). 2374 (Reguault, J. Ch. [3] 1, 129). H. F

Preparation, - By heating B in O, or by strongly heating boric acid (q. v.).

Properties. - Semi-transparent, edourless, brittle, inodorons, glass-like, solid; volutilised at a very high temperature (Ebelmen, A. Ch. [3] 22, 211); volatilised in steam or alcohol-vapour; non-conductor of electricity (Lapschin a. Tichanowitsch, P. M. [4] 22, 308; Bowgoin,

C. R. 67, 798).

Reactions. -BO is a very stable compound; it is not decomposed by heating with powdered charcoal or with i' vapour. It is an unhydride, but appears to show a feebly basic character in its reactions with certain acids (infra 6-8). 1. Heated with potassium, so lium, ar aluminium, metallic oxide and B are formed - 2. Mixed with charcoal and heated in nitrogen, chlorine, browni w, or carbon displyle'e, BN, BCl, BBr, or B.S., is formed .- 3. Salts of most acids, e.g. ulphates, nitrates, carbonate, are decomposed by heating with BaO, to a high temperature, with production of borates and volatilisation of the acid (v. Tate, C. J. 12, 160) .- 4. Reacts with most metallic oxides at high temperatures to form borates.-5. With water forms boric acid (q. v.). - 6. B2O2 is said to react with fuming sulphuric acid to lorin a compound aBO, ySO, sHO; the values given to x, y, and sby different ohemists vary; thus Merz gives the tor-

mula 5B, O. 2SO, 2H, O(J. pr. 99, 181), and Schuliz-Sellack gives the formula B, O. 3SO, H, O(B. 4, 15). This compound is easily decomposed by heat to B,O, and BO, -7. B,O, is said to form a compound with phosphoric anhydride, B.O., P.O.; this body is produced by heating together H,BO, and cone. H,PO,Aq (Vogel, Z. 1870. 125), and removing excess of the latter by hot water; it is also formed, according to Gustavson (B. 3, 426; 4, 975), by heating B.O. with PCl, to 140° for 3.4 days, and also by heating B.O. with POCl, to 150 -170" for 8 to 10 hours, distilling off the POCL BCL formed (v. Bonon, chronibe or; Combinations, No. 1) and strongly hoating the residual solid. B O, P,O, is said to be insoluble in hot water, to be museted on by scids, dissolved by boiling KOHAq, and to be decomposed by heating with Na, giving Na phosphide, and probably phosphido of B.—8. A compound of B.O. with tangstic anhydride (B₂O_w9WO₁xH₁O) is described by Klein (BL [2] 86, 205), r. Tunastonomarks, under Tunasten. 9. B_iO, dissolves in hydrofluoric acid, forming B.O. 611F.11.O (... 11.B.O. 611F) (c. Fraionoute Acro, under Boios, oxyomes or, p. 530). - 10. When a solution of 1 part B.O. and 2 parts KILC, ILD, in 24 parts 11.0 is evaporated to dryness at 100°, and the residue is treated with alcohol, a white, amorphous solid remains, insoluble in alcohol but very soluble in water. This solid has the composition C.H.KBO .; its reactions are similar to those of tartar emetic; probably it is the K salt of an acid B.C, H,O,DH anulogous to the acid Sb.C₄H₁O_aOH obtained by Chirko a. Stallo (B. 13, 1787) (c. Meyrac, J. Ph. 3, 8; Soubeiran, J. Ph. 3, 399; 11, 560; 25, 741; 35, 241; Dullo ; S. 44, 333; Vogel, J. Ph. 3, 1; Robiquet, J. Ph. (3) 21, 197; Wackenroder, Ar. Ph. [2] 58, 4; Wittstein, R. P. [3] 6, 1, 177; Duve, J. 1869, 540; Biot, A. Ch. [3] 11, 82; v. also under Tsuriari s).

Boroa, Oxyacids of, and their Salts. definite hydrates of B.O. appear to be known; orthoboric acid B₁O₂3H₁O (=H₁H₂O₂), metaboric acid B₂O₂H₂O (=H₁H₂O₂), and pyroboric (or telraborie) acut 2B,O,41,O (11.B,O.). Another hydrate 2B,O, 3H,O was described by Berzelius (S. 23, 161) as obtained by heating 11,130, *oonsiderably over 100 % Most metallic borates may be regarded as derived from H.B.O.; some -e.g. borax from H B.O.; a few are perhaps derived from H.B.O., e.g. Ca.B.O.; and in addition to these several borates exist which at present must be regarded simply as compounds of B.O. with metallic oxides (v. infra). All the boric acids are ' weak ' acids, thrir salts are easily decomposed by reactions with other acids; the admity of boric acid is extremely #maB, Na,B,O, in solution is entirely decomposed by an equivalent of 11 SO, (Th. 1, 200). Borates of the less positive metals are usually formed only by Insing together B_iO_s and the metallic oxides. Many of these borates are decomposed by water; some of them are partially converted into carbonates when exposed to the action of moist air; borates even of the very positivo metals readily combine with boric sold to form soid salts, but these salts, although containing excess of borie acid, usually turn red litmus blue; even alkali borates are partially decomposed by water. Thomsen's thermal investigation of borie and proves that the asid obtained by dissolving B₂O₂ in water is dibasis; thus;—

m[mNaOHAq,B²O²Aq]
1 11,101
2 20,010
3 20,460
6 20,640

The normal Na borate is therefore Na,B,O, Whoa borio acid is added to a solution of this salt heat is produced; thus:

m Na²B²O 'Aq,mB²O 'Aq]
1 2,192
4 4,911

Acid saits are therefore probably formed, bot the reaction of the solution towards lithus is still alkaline (Th. 1, 206).

Onthonomic Acto. 11,180, (Boracic Acid; Boric Acid). [181°-186°] (Carnelloy, C. J. 33, 275).

Occurrence.—In the waters of many volcanio districts, e.g. in 'In scany; in many mineral springs; in combination with NA as borax—in the waters of certain lakes in S. America, Thibet, Ceylon, &c.; in several minerals, e.g. borneite (borate of Ma), boronative alcite (borate of Ca and Na).

Formation. 1. Be oxidising B with aquaregia, evaporating, disolving in water, and recrystallising. 2. By dissolving B_iO₄ in water.

Preparation,—3 parts crystallised borax are dissolved in 12 parts boiling water, and (after liftering if necessary) 1 part cone. 14,804 is added; boric acid separates on cooling. The crystals are gently heated, recrystallised from water, dried, fused in a Pt crucible (to remove all 14,804), again recrystallised from water, and dried by pressure between titler paper.

Projectics. White, semi-transparent hanima, triclinic (Miller, P. 23, 588), monochino (Eurgott, Sitz. W. 12, 26). S.G. J2 1-434 (Stella, J, pr. 90, 457). S. (19°) 3-9, (25°) 6-72, (77.5°) 7-9, (50°) 9-84, (65°) 16-34, (75°) 21-15, (87.5°) 26-37, (100°) 33-67 (Brandes a, Frinlaber, Ar. Ph. 7, 50; v. also bitte, C. R. 85, 1969). S.G. of Π_b BO_bAq saturated at 15° ≈ 1-0218 (Stolla, J, pr. 90, 457). Heat of solution, (II BO°, Aq] ≈ -5335 (Th. 3, 196). Soluble in alcohol and several oils (Rose, P. 80, 262); soluble in warm coac. H SO₄, HCl, or HNO₃Aq, but most of the boric acid separates on cooling. Aqueous solution turns blue littuus wine-red, and turmeric paper cherry red; alcoholic solution burns with green-adged flame.

Reactions.—I. Heated to 100° H.B.O. is produced (Schaffgotsch, P. 107, 427; Bloxam, C. J. 12, 177; Merx. J. pr. 99, 179); hoated to 110° for a long time, or to 160° in a current of dry air, H.B.O. is formed (Merx. Lc.; Ebelmen a. Bosquet, A. Ch. [3] 17, 63); heated to about 300° the wride B.O. renaius. Berzelius (S. 23, 161) said that H.B.O. (=2B.O.,3H.O) is formed by heating H.BO. to a temperature 'considerably above 100° '; Merz (Lc.) affirmed the production of 8B.O.,H.O at 270°—2. Boiling cone. H.BO.Aq dissolves a fow metallic sulphides and oxides (Tissier, C. R. 39, 192; 45, 411); decomposes alkali and alkaline earth arbonates (Popp. A. Suppl. 8, 10).—3. With alkalis and alkali-carbonates salts of the form

minute of mility. TH.H.O. are generally pro-duced; few metallic salts of the form M.BO. are cartainly known; hence H,BO,Aq reacts no H.B.O.Aq (salso Borates). Ethereal salts of e.g. Et,BO, Me,BO, are known, but none of them is directly formed from H.BO. (p. 530).

METABORIC ACID, H.B.O.; and Pynonoric or TETRABORIC ACID, 11 B.O. Formed by licating H,BO, (r. supra); glass like, amorphous solids. Some salts of If B O, are obtained by the reaction between H.B.O.Aq and alkalis; salts of H.B.O. are obtained indirectly (v. infra).

Bouards. No borate is quite insoluble in water; the alkali borates are very soluble. The less soluble borates are easily decomposed by water, the easily soluble salts are also decomposed, but less quickly; an alkali borate, for instance, in cone aqueons solution slightly reddens litmus, but when much water is added the litnus becomes blue. Solutions of alkali borates absorb CO, and HS; they decompose NH, salts when boiled with them; dilute solutions react with Hg and "Ag saits similarly to alkali solutions. A few borntes can be obtained as definite, fairly stable, saits by precipitation from solutions; KHB O, 2H O separates from a solution of K.CO, in excess of B.O.Aq to which much KOHAq has been added; MgBO, 411 O. is said to be formed by the reaction between borax solution and Mg2NO Aq. The more definite burates are generally obtained by melting together B.O, and basic oxides. As a class the borates very reactily undergo change; the composition of very many is therefore extremely doubtful. H. Hose, who invest tell many borases, did not attempt to wash his preparations, but pre-ed them between filter paper till dry, and there determined the quantity of the admixed foreign so ta. Most horates seem to belong to the two forms M.B.O. and M.B.O.; many may be represented as M.B.O., all B.O. and M.B.O., eH.B.O.; a few - e.g. 3M/O.B O. may be regarded as derivatives of HeliO₂. The best-marked borates are the suits of K and Na.

Potassium borates. -(1) Normal metaborate, K.B.O., Monoclinic crystals (a:b;c - 2744:1:2-676); by melting, at white heat, I part B.O. with 1:05 parts K CD, dissolving in water, evaporating to a syrup out of contact with air. and crystallising (Schuhes, Bestime ung der Krystalligestellen Av. (When, 1855), 31). (2) Acid melaborate, K.B.O. II, P.O. 411.0. Begular six-sided prisms; by saturating bailing K CO Aq with B.O.Aq. adding KOHAq to strongly aikaline reaction, evaporating, and crystallising (Laurent, A. Ch. 2 67, 215). Said to some time 8 crystallise with \$11,0 in rhombic prisms; and to lose H.O when heated in a chold vessel. -(3) Other acid salts. K.B.O. 211 B.O. 611 O. obtained like (2), but using less KO18Aq. 2K_B_O_4H_B_O_5H_O(or 411 O), by adding B.O.Aq to boiling K.CO.Aq un. I solution has a slightly said reaction (Rammelsberg, P. 95, 199; Reissig, A. 127, 33).

Sodium borates.—(1) Normal meta-borate, Na,B,O,4H,O. Large monoc nio like mass.

prisms; by fusing 1 part dry Na CO, with 1-17 parts B,O, for with 3-6 parts crystallised borax, dissolving in water, and crystallising (4. 76, 267) by evaporating aqueous borax M M

out or contact with air. Melts at 57%? as with 3H₂O separates on cooling. Mixed wit NaF, in solution, and crystallised, the sa-Na B.O. 6NaF. H.O separates (v. FLOOBORIO ACIE (Hahh, J. 1859, 128),—(2) Acid melaborate, Na. B.O. 3H, B.O. 7H, O. lustrous, hard, crystal fine crust, obtained by boiling solution of a equivs, borax with 1 capus, NII,Cl so long as NII, is evolved, and crystailising (Bolley, A (8, f22). Na.B.O. 411.B.O. 711.O. small crystals obtained by dissolving in water I camy, boras with 5 equivs 11,80, and crystalleing (Lanrent, C. R. 29, 5), (3) Ortholomic, Na BO, Said to be formed by fishing BO, with excess of to be formed by fusing BO, NaOll (Bioxam, C. J. 14, 113). (1) Term, or pyro, borates. (a) Borax. Na li O. 10H1O, ordinary or prismatic borax; Na li O. toll2O, octaholral borax. The former ocenrs native; it is obtained by purifying crude bounk, or by fusing I part dry Na CO, with 2:34 parts II BO., dissolving in warm water, and slowly crystallising from a solution of S.G. 1 14-1-15 (B.P. 101"). stopping when the temperature has fallen to 25' 30 . The saft with 511 O separates from aqueous solutions of ordinary borax of S.G. 1:246 at tempentures between 56° and 79°, or from supersaturated solutions of the rame suit protected from dust, or from aqueous solutions of any though evaporated at 10° to 12' (Gernez, C. R. 78, 68). Old navy borax crystallises in large, transparent, columbers, doubly relactive. monoclinic prisms (area: 10995;1:5629, # 573 25). S.G. 169 (Filliof, A. Ch. (3) 24, 415). 8. (0.) 2.8, (20.) 79, (10.) 179, (60.) 404, (80.) 762, (90 1 1197, (100) 2014 (Poppinte, A. Ch. [3] S. 953), S.G. of solution saturated at 45° π 1:0159 pontains 38:494 boraxi (Michel a. Kraft, A. Ch. (3) 44, 471). Insol. in abrolub. Heat desappears during colution; [NacB'O', t0HO, Aq1

25,860 (Thornam). S.H. (Br. 50.) 385 (Kopp. T. 155, 71). Refractive indices (23", Na light P fe = a 1-1163, for β = 1-4682, for γ : 1-4712 (holdrausch, W. 4, 1). The crystals effloresce in hir toronding to Sims only when they contain a circum carbonate); when heated they melt and give off 10H O, leaving burnt borax (Na.B.O.), which mells at a red licat to a glass like mass (citrified boras) of S.G. 236; exposed to moist air this take ap 1911.0. Solution of borax in water is alkaline to litmus; it di solves many or same compounds more readily than water, c.s. beneauce and and pullicacid; some compounds in obtable in water discover in horax * dution, e.g. dec le said, various gams, resins, and oils; As,On distolver casily; rilicia neid only very stibilly. Molten borax dissolves many metallic cycles and salts forming fusible double calls; hence its use as a max, and also in analysis. Crystals of cornolum were ob Al O, in molten borax, and erystals of ratile and tridymite by dispolving TiO, and SiO, respectively, in the same to'vent (cr. Rose, J. pr. 101, 228; 108, 208). On inched bonce ery tallises in bard, transparent, regular octahedra. S.G. 1-8. Unchanged in dry air, but in moist sir

solution at 100° and drying the residue at the same temperature for a long time.

(c) Na, B, O, .6H, O was found by Bechi (Am. S. [2] 17, 129; 19, 120) in an old lagoon; it has not been prepared artificially.

The remaining borates have been chiefly investigated by Berzelius (S. 23, 160; P. 2, 113; 9, 483; 33, 98; 31, 561); Arfvedson (Gm.-K. 6th ed.); Guaclin (v. Gm.); H. Rose (P. 9, 176; 19, 153; 86, 561; 87, 1, 470, 587; 88, 299, 482; 89, 473; 91, 452); Wöhler (P. 28, 525); Itammel:berg (P. 49, 415); Ebelmen (J. Ch. (3) 16, 129; 17, 51; 33, 31); Bouquet (A. Ch. [3] 17, 54); Bolley (A. 68, 122); Herapath (P. M. (8) 34, 575); Laurent (A. Ch. (2) 67, 215); Tissier (C. R. 39, 192; 45, 411); Bloxum (C. J. 12, 177; 11, 113). For an account of various supposed compounds of MO with BO, v. Ditte (A. Ch. (5) 30, 218). The following are the salts which have been chiefly examined; but the composition of many is not citled.

Aluminium. 2Al O₃ B O₆ SH O ; 8Al₂O₃ 2B₃O₄ 7H₄O (H. Ro, c). 3Al₄O₃ B₂O₄ (Ebelnere).

Ammonium. - (NII.) B.O. 11. B.O. 211. O (also 3H₂O₁ (Arfvedson), -- (N11_d), B₂O₂, 511_dB₂O₂, 4H₂O (Beehi, Am. S. (2) 17, 129; 19, 120).

(811,) B.O. 311 B.O. 311 O (Gmelin). (NII,), E,O,, 111, B,O,, 211, O (Rannaelsberg).

Bartum. - BaB O₄,10H O (Berzelius). $BaB_zO_zH_zO_z$ (Rose), - BaR O_cH₂B₂O_p4H₂O; BaB,O,2H,B,O,12H,O; 2BaB,O,H,B,O,11H,O (Laureut), Ba, (BO,), ; 2DaO.B.O, ; 5BaO.2B,O, (Bloxnin). 2BaO.3B₂O₂; BaB₄O₅ (Ditte, C. R. 77, 788).

Cadminm. CdB₂O₄; 2CdB₂O₄CdO₂H₂2H₂O (Rose).

Calcium .-

CaB₂O₄,211₂O; 2CaB₂O₄,H₂B₂O₄,411₂O (Rose).--CaB,O,H,B,O, (Tönnermann).-CaB,O,3H,B,O,6H,O (Laurent).-CuBlO₃: 3CuO.2B.O₄; 2CuQ.3I\(\phi\)O₄; 3CuO.3M(O.4B.O₃) (Ditte).

Colait. 2Coll.O. CoO.H. 3H.O (Rose). Copper. Composition very uncertain (v. Tümermann, Rose, Laurent, Polley; le Boux, C. H. 64, 126; Pasternack, A. 151, 227; Poussier, B. 6, 1138).

Didymium. DiBO, (Clive, Id. 12) 43, 36 t). Iron. - Fr(B,O,),311,O; found in a lagooncrater (Bechi, Am. S. [2] 17, 129; 19, 120). Basic salts of meertain composition are formed by the reactions between alkali borates and solutions of ferric salts; borntes of Fo and Naseem to be produced by ppg, iron alum with

alkufi locates (Rose). Lead. PhB₂O₂H₂O; 2PbB₂O₄H₃B₂O₄3H₂O;

PbB_O, II, B_O, 3H_O (Hernpath; r. also Rose).

Double salts; PbB_O, Pb(NO₃), II, O; PbB_O, PbC[, H)O (Herapath).

Magnesium.— MgB_OO, III, O; MgB_OO, 3H, B, O, 8H, O; MgB,O₃5U,B,O₄13H,O (Laurent; Rammelsberg), ---MgB,O₆8H₂O (Wöhler), ---Mg,dBO₃ (Ebelmen), Bouble salts: 6MgO,3X,O₂2B,O₃ (X = Cr or Fe) (Elehnen). Ditte describes various compounds of the form xMgO.yCaO.zB.O.

Nickel. = NiB.O. 211.O (Rose): also various vague lasie salts.

Rul idium ... Rb.B.O., 611,0 (Reissig, A. 127,

Silver.—Ag.B.O. H.O.; SAg.O.4B.O.; (Rose)... Strontium.—SrB.O.; SrB.O.; SSrO.2B.O.; 2SrO.3B.O. (Ditto).—SSrB.O.,2H.B.O.5H.O. (Rose).—SrB.O. H.B.O. 3H.O; SrB.O. 3H.B.O. 3H.O (Laurent).

Zinc. -- Very vague (v. Rose). Büscher (4. 151, 231) describes the double salt ZuO.4NH,.2B,O,.6H,O.

[? Samarium borate. - SmBO, (Clève, Bl. [2])

43, 170.1 FLUORORIC ACID, and FLUORORATES. B.O. sp. pears to react as a feebly basic oxide towards the auhydrides of a few acids, e.g. SO, and P.O. (c. Romas, exter or, Reactions, Nos. 6, 7). 11,130, dissolves in cone, HFAq; by concentration, and cooling over 11,80,, a thick syrup-like liquid is obtained (S.G. 1581) containing H.B.O. and HP in the ratio 11.3 O. 611P. This liquid is generally regarded as a definite acid, called fluoboric word. This liquid chars organic matter like Il SO. The rame liquid is obtained by saturating water with BF, and distilling (Gay-Lassne u. The nard, Recuerches physico-chimiques, 2, 38; Berzelius, P. 58, 503; 59, 644). The liquid is decomposed by waterinto HBF, Aq and H.B.O. (r. Bonoracountring acre, under Boros, recommon or). If this liquid is neutralised by NaOllAq or KOHAq, and the solution is evaporated, the salts M_B_O_c6MT.H_O (M = Na or K) are obtained (Berzelius). The same salls are also formed, when M = K by fusing KF with H.BO, and when M - Na by crystallising mixed additions of Na B.O. and NaF. His very doubtful whether the so-called fluoboric neid is a definite compound or not. According to Basarow (C. R. 78, 1698) the liquid prepared as described is decomposed by distillation; at 140? Bl, is given off, at 160° to 170 a thick, heavy, fuming liquid (S. G. 1-77) comes over, at 175"-185" a less funding liquid is produced, and as the temperature rises the distillate becomes lighter and fumes less in air. The heavy distillates are decomposed by water with separation of H_sBO_s. The salts obtained by Berzelius are separated by crystallisation into MF, which crystallises out first, and a mixture of MF with MaBaO, (Pasarow). tion of the so-called neid react; with AgNO, Aq to give Ag₂B₂O₄ and Ag₂O. Busarow regards fluohoric acid as a mixture of H.B.O. with HBF. and 11F. Landolph (B. 12, 1583) describes the bodies H.B.O., 3HF and H.B.O., 2HF; the first is obtained in small quantities by the reaction between BF, and C,H,, the second is one of the products of the action of BF, on hot anethol. Both bodies are funning, acid, liquids; they seem to be fairly stable; the first is unchanged

by distillation. Boucrondstates. Many compounds of the form xB,O_s, fWO_s, zMO (MO = metallicoxide) have leen oldained the acid B.O., 9WO, 2H, O. 18aq has been prepared. The principal borotungstates will be described under Tungsten as TUNGSTOBORATES.

Detection and Estimation of Boric Acid. Free boric acid is detected by its action on turmeric paper, or by the green colour which it imparts to the flame of burning alcohol; borates do not give these reactions, therefore they must be decomposed by If SO, Aq before applying the alcohol test, or by HClAq before applying the

semionic paper test. There is no very satis | 2B(OEt)₂+3ZuMe₂=2BMe₂+3Zu(OEt)₄(Frank-term mathed for satimating haris acid. The | Iand. O. J. 15, 373). Pungent gas. V. sl. sol. sectory method for estimating boric acid. The most insoluble salt is KBP, ; it is obtained from borio acid or borates by adding excess of ROHAq, then evaporating with excess of HFAq, dissolving out sulphate, nitrate, &c. of K, by KC, H₂O₂Aq, washing with alcohol, and drying at 100°. Marignac (Fr. 1, 400) evaporates the solution of the borate with excess of NIL ClAquard MgCl.Aq, with various precautions, and finally obtains a mixture of MgO and Mg horate in which he then estimates the amount of MgO and so gets the amount of boric acid (r. a)-o Ditte, C. R. 80, 400 a, 561). As borie acul interferes with estimation of some other bodies, e.g. phosphoric acid, it is sometimes necessary to remove it; this may be done outher by boiling with alcohol and H SO Aq (Et, BO, goes off), or with HF and H SO Aq (BF, goes off). Recenbladt (Fr. 26, 18) describes a method lased on the volatilisation of boric acid by distallation with methylic alcohol; the method gives good results (c. also G ech, C. N. 55, 7).

Boron, exychlorides of. Two are known, BOCl and BOCly; meither exists as a cas; both are decomposed by heat. BOCl it a white gelatinous soud, obtained by heating B.O. with BCl, in the ratio B₂O :2BCl, to 150 ; at a high tomperature it is descenposed to BCl, and BO, (Gustavson, Z. 1870, 521). BOCk, is described as a yellowish green liquid; it was obtained, along with BCl, by parting Cl aver a heated mixture of B O2 and C, it moving Cl from the gaseous products by Cu turnings, condensum, and removing BCl, by evaporation (Councier, J. pr. [2] 18, 3000. The conditions under which is formed are not definitely known; Councier obtains I the lost result when relatively little carbon was used, and a fairly rapid stream of Cl was pa sed through the tube. BOCh is decomposed by heat into BCl, BO, and Cl; and by water to H.BO, HCl, and Cl. chloride of B is formed by the action of ozone on BCl, or by passing electric sparks through a mixture of BCl, and O (Michaelis a. Becker, B. 14, 914).

Boron, sulphide of. R S., Med. w. unknown, as compound has not been pasified.

Formation. 1. By heating B in S vapour to white heat (Berzehau, P. 2, 145). -2. By gently heating B in dry H S (Wohler a. Deville, A. 105,

Preparation. Small pellels are made by mixing powdered B.O. with soot and oil and out of contact with air; these are heated to full reduces in a poscedain tabe, in a stream of thy CS;; the tube is connected with two small thacks surrounded by snow and sait. The B.S. collects on the surface of the con b mold CS₂; it is separated from CS, and direct in an atmosphere of H (Frenzy, A. Ch. [3, 38, 312). B.S. is a white solid (with a yellowish times, Fremy), consi-ting of groups of small crystals; it smells strongly, and its vapour acts on the eyes; it is rapidly decomposed by mediture to B,O, and H S; it may be melted in an atmosphere of H, and volatilised in a current of H S. M. M. P.

BORON, ORGANIC DERIVATIVES OF. Boron tri-methide C.H.B i. BMc., V.D. 191 of the solid product from other. (cale. I 93). From ethyl borale and ZnMe, thus:

land, O. J. 15, 373). Pungent gas. V. sl. sol. water, v. sol. alcohol and ether. Takes fire in air, hurning with a green flame. Not affected by cone. H.SO, or by HI. Combines with potash forming BMc, KOIL. Combines with ananonia forming BMc, NH, [56°, (110°). Combines at o with NaOH, CaO, and BaO.

Boron tri-ethide BEG, Mol. w. 98, (95°), S.G. 44: 600, V.D. 840 (calc. 3:10) From BCl, or B(OEt), and ZnEt. (Frankland, Tr. 1862, 167; Pr. 25, 165). Pung at oil. Takes the mair. Slowly decomposed by HCl, evolving C.H., Violently attacked by chlorine,

Combinations, -1. With ama min it forms a liquid 10 t,NH, 2. By careful exidation, first in air, then in oxygen, it forms in exide BUL O. (125"). Water decomposes this oxide, forming et hyl-borie nerd, Blit(OII),. Ethylborie acid is crystalline, and may be sublimed; its solutions are need, but it does not form salts. A compound BEt(OEt) B(OEt). te. 112 I appears to be formed by the action or Zulit (I mol.) on borie ither; it is decomposed by water into alcohol, BEt(OII), and B(OII), and by ZuEt, it is converted into DEC (OEt) (103), which absorbs oxygen, becoming BEQUEO . Dischyl-boric ether, BEL (OEt) is caponified by water, and the acid absorbs oxygen, begoining crystalline BEt(OEO(OH), which is converted by water into ethyl borin neid BEGOID.

Boron-phonyl-di-chlorido C H BCl., |ahout (175). Prepared by heating horon bri-chloride with mercury diphenyl at 2002 (Mighaelis a, Becker, B. 15, 180). Colourlesa faming this. By the action of water it gives phenyl lorie neid, and by the action of alcohol, phenyl-boric other.

Boron - je toly1 - dichloride Call (CHa). BCl. . Colombes crystals. Prepared by heating boron tri chlegide with mercury di-p-tolyl (Michaelis a. Becker, B. 15, 185),

BORO-TARTAR EMETIC v. Boron, exide DE; R orthons, No. 10; also under Tartrates. BOROTUNGSTATES. Compounds of the form xB₂O_xyWO_xxMO(M = metal), v. Tunosronorvies, under Tungsten.

BRASILEIN v. BRAZILEIN.

BRASS v. Coppen, Alloys of

BRASSIC ACID C HigOn [60°]. Prepared by just melting its isomeride, erucic acid, with blute HNO, and adding sodime nitrile. Lamine (from alcohol). Combines with bromine. Potash fusion gives arachic acid.

Salts. NaA'; Immine Gran alcohol). Ethyl ether Eth'; [30]; (dove 360°); clistenin (plates; obtained by otheritying the and or by the action of nitrous acid upon ethyl fracate.

Glycerin-di-brassic ether C.H (OH)A' .; (librarsidin) [65%; crystal, al. sol. ether. Formed from glycerin-th crucic ether by nitrous acid.

Glycerin-tri-brassic ether C.H.A.: Technisselin, '17 hafterheating [36°; colourless crystalline powder; v. soi, ether, nearly insol. absolute. Obtained by the action of nitroas and upon rape-seed oil, and crystallisation

Amide Calla CONH : [90]; colourless

needles; formed by the action of NH, gas upon the anhydride.

Antiide C_nH₄₁.CONHPh: [78°]. Anhydride (C₂H₄₁.CO)₂O: [29°]; glistening tables; v. sol. other and benzene, fasol. alcohol and water; formed by the action of PCl, upon brassic acid and subsequent addition of alcohol (Reimer a. Will, B. 19, 3320; cf. Wesky, J. pr. 58, 419; Haussknecht, A. 143, 40; Fitz, B. 1, 411; Goldschmiedt, Sitz. B. 74, 394).

BRASSYLIC ACID Call of Mol. w. 216. [109°]. Formed, together with its aldehydo and dioxybelienolic acid, by the action of fuming HNO, on behenolic acid (Haussknecht, A. 113, 45). Crystalline, v. sl. sol. cold water, v. sol. alcohol and ether.

Salts.-CaA" 3aq.-Ag2A".

BRASSYLIC ALDEHYDE C,H, O, chief product of the action of faming IINO, on behenolio acid (v. supra). Oil, lighter than water, volntile with steam. Sol. NaOHAq and reppd. by HCl. Oxidised by bromine to brassylie acid.

BRAZILEIN C16 II 12O, II2O.

Formation. - By the oxidation of brazilin by alr in presence of alkalis, or in ethereal solution by HNO₃ (Reim, B. 4, 334; E. Kopp, B. 6, 416; Liebermann a. Burg, B. 9, 1883; Buchka a. Erck, B. 18, 1142).

Preparation. - Extract of Brazil-wood is discolved in hot water and, after cooling, NII, in slight oxcess is added. The solution is exposed to air, when a pp. is formed which is crystallised from hot dilute acctic acid (Humanel a. A. G. Perkin, C. J. 41, 373).

Properties. - Minute crystals with grey lustre. Reddish-brown when powdered. Very slightly soluble in cold water, more so in hot water. The solution is yellowish-pink with greenish-orange fluorescence. Alkalino solutions are carmino

red, but slowly turn brown in air.

Reactions. -1. If hot glacial acetic acid be slowly added to a solution in cold cone. It. So, minute orange needles of iso-brazilein sulphate, On It, O.SO, II, are got. Its alkaline solutions are carmine red quickly turning brown in air. Alcohol turns iso brazilem sulphate scarlet, forming the basic salt $C_n H_n O_2 Q(C_n H_n O_8 O_1)$.

2. Conc. HCl at 100° forms $C_n H_n O_8 C_1 H_1$.

7. Conc. HCl at 100° forms $C_n H_n O_8 C_1 H_1$.

8. Montte rod priscus with violet lustre, called iso-brazilein

3. HBr at 100° forms, similarly, $\bullet G_{10}\Pi_{11}G_{1}Br$, Brazilein resembles humatem (q, v,) in these reactions.

BRAZILIN $C_{10}\Pi_{10}G_{2}$. Occurs in Brazil-wood (the wood of $C_{10}G_{2}^{*}$) in the military of $C_{10}G_{2}^{*}$) occurs in Brazil-wood (the wood of $C_{10}G_{2}^{*}$) in the military of $C_{10}G_{2}^{*}$) occurs in Brazil-wood (from $C_{10}G_{2}^{*}$) in the military of $C_{10}G_{2}^{*}$) and in $C_{10}G_{2}^{*}$ (Gladstone, P. M. [31] 35, 315). Br, and $H_{2}O_{2}^{*}$ (Gladstone, P. M. [31] 35, 315). Br, and $H_{2}O_{2}^{*}$ (Gladstone, P. M. [31] 35, 315). Br, and $H_{2}O_{2}^{*}$ (Ma.SO, Aq + H, O + Br₂ = Na.SO, Aq + H

deposited during the preparation and storage of commercial extract of Brazil-wood consist of brazilin and its lime compound. The crusts are washed with dilute (5 p.c.) HCl and then extracted with very diluto (12 p.c.) alcohol.

Properties .- Colourless' crystals (containing aq). Sol. water, alcohol, and other. Turns orange in air. Forms a carmine solution in aqueous NaOll in air; this solution is bleached by sine-dust, but re-oxidised to brazilein by air. Aqueous scintions are also turned red by NB or haryta when exposed to air,

Reactions, -1. Resorcin is among the products of its dry distillation. -2. KClO, and HCl give iso-tri-ohloro-glyceric acid (Benedikt, Salt.-PbA"aq: small colourless needles

Tri-acetyl-derivative C. II, (OAo), O, 1 [106°]; slender colourless needles.

Tetra-acetyl-derivative C, II (OAc),O: [151°]; glistening needles (Buchka a. Erck, B.

BREIDIN v. ARBOL-A-BREA. BRIMSTONE v. SULPHUR. BRITANNIA METAL v. Tin, Alloys or. BRITISH GUM v. DEXTRIN. BROMAL v. BROMO-ACRTIC ALDERVIDE.

BROMALIDE C.H.Br.O. i.e. CBr₃.CII CO.O CII.CBr₃.

Tri-bromo-ethyl. idene tri-bromo-lactate. [158°]. Formed by heating bromal hydrate with II SO₄; or by warming a mixture of bromal and tri-bromo-lactic acid (Wallach, A. 193,1; Wallach a. Reinecke, B. 10, 2128). Monoclinio crystals, insol. water, sol. other. Decomposed by alcohol.

BROMANIL is TETRA-BROMO-QUINONE (q. v.). BROMATES AND PERBROMATES. - Salts of bromie and perbromie acids; v. Bromine, OXYACIDS OF.

BROMHYDRIC ACID. HBr. (Hydrobromic acid. Hydrogen bromide.) Mol. w. 80-75. [-73° (-69) (Faraday, T. 1823, 189). V. D. 39·1. S. (-25° to 0° at 760 mm.) about 690; S. (-25° to 0°at 2 min.) about 345 (Roozchoom, R. T. C. 4, 102). II. F. [II,Br] = 8,440; [II,Br,Aq] =28,376 $[\Pi, \text{BrAq}] = 27.837 \quad (Th. 2, 29).$ $\Lambda - 1$ × At. wt. = 20.6 (Gladstone, T. 1870. 9).

Il and Br do not combine at ordinary temporatures even in direct sunlight.

Formation. I. By burning II charged with Br vapour. -2. By passing a mixture of H and Brover hot Pt; for details of method v. Harding, B. 14, 2085. 3. By the action of electric sparks on H and Br. -4. By the action of Br on H2O, more quickly in presence of exidisable bodies such as P. S. As, or lower oxides of these elements. - 5. By passing 11.0 and Br through chlorhydrin. Its aqueous solution is orange. By passing ILS into Br and H₂O (Naumaon, 3, HBr at 100° forms, similarly, $G_{l_0}L_{l_1}O_{l_1}Br$. By passing ILS into Br and H₂O (Naumaon, B. 9, 1577). –7. By passing III into Br (Haute-

(Champion a. Pellet, C. R. 70, 620).

Preparation.—1. When small quantities are required, Hir may be prepared by the action of P and Br on 11,0 (111,0 + P + 5Br - H,PO, + 5HBr). A glass tube is bent 3 times at about a right angle; a little Br is placed in one bend and a few pieces of P in the other; pieces of glass moistened with a very little water are placed in the limb of the tube above the P; a cork with delivery tube is fitted into the open end of the tube above the glass, and the other end of the

tube is closed by a cork. The Br is then very gently warmed; the reaction occurs when the Br vapour reaches the moist P, and the HBr passes off through the delivery tube .- 2. When larger quantities of HBr are required it is advisable to use amorphous P. In a flask fitted with a cork carrying an exit tube and a small stoppered separating funnel, is placed 1 part amorphous P mixed with some dry sand, the I is moistened and then covered with a layer of dry sand (Linnemann, A. 161, 198 note); the exit tube is connected with a U tube nearly filled with glass beads moistoned with cone. HBrAq and piece: of ordinary I' (any Br which may come over is converted into IIBr in this tube); this is followed by a drying tube containing CaCl, or P.O., and from this a delivery tube passes into a dry cylinder filled with dry Hg standing in a Hg trough. Ten parts of Br are placed in the separating funnel and allowed to drop slowly into the flask; HBr is evolved. Towards the close of the operation the flask is gently warmed. If it is desired to prepare an aqueous solution of HBr, the U tube is fitted with an exit tube passing into the tubulus of a retort placed vertically and arranged so that the beak dips a little way under the surface of water in a bottle; should the flow of Hisr slacken, the water rises into the lody of the retort but cannot flow back into the generating vessel. -3. By the action of con. H.PO.Aq on KBr; 1 part KBr, 1 part H₂PO₀ and 3 parts H₂O being used (Bertrand, C. R. 82, 96) .- 4. By decomposing the bromide of an alkaline earth metal by ILSO Aq; Bertrand (l.c.) employs two parts Calle, 2 parts cone. If St. and 1 part H.O. If an alkali bromide is used, the HBr contains much Br and some SO, -5. An aqueous colation of HBr may be obtained by rharly adding P in small pieces to Br mixed with a considerable quantity of water in a vessel -arrounded by ice, then addin; a little more Br and then a few pieces of P (repesting if a strong solution is required), and distilling from H.PO.Aq (v. Topsöc, B. 3, 400).

Properties. Hitr is a colourless gas, with pangent, neid, very irritating, odour; excites inflammation when applied to the skin; fames in moist air; dissalves very largely in water; (v. intra) is absorbed by, and melts, ice; at +73; (v. intra) is absorbed by, and melts, ice; at +73; fliqueties, and then crystalrises. Melsona (C. R. 77,781) obtained liquid Hitr by saturating wood charcoal with the gas (15,500 grain-units of heat are produced for every 81 grains Hitrabsorbed, Favre, A. Ch. [5] 1, 200), placing the charcoal in one end of a closed glars tube bent to an obtase angle, the other end of which was well cooled, and heating the charcoal in a water bath. An aqueous solution of Hitr form: a colourless, strongly acid liquid; the affinity is a very little less than that of HCMq r. Arrivary.

The cone solution fumes in, but is not decomposed by exposure to, a.r. S.G. of solution saturated at 0°2-1·78; 1 e.e. contains 1·46 grams HBr (=82·02 p.e. HBr by we highly which almost agrees with the composition calculated from the formula HBr.H.O (Bineau, A. 44, 237). Boozeboom (R. T. C. 5, 363) has obtained the hydrate HBr.H.O as a solid at low temperatures and under a pressure of 3 atmos. If cone, HBrAq is distilled at 760 mm, pressure TBr is evolved. if HBrAg containing less

than 47 p.e. HBr is distilled at 760 mm. H₂O is evolved, in each case the B. P. becomes constant at 126° and the solution contains 47:38 - 47:80 p.c. HBr; the composition of this liquid is almost exactly that expressed by the formula HBr.511 O (V. D. = 14-1 agroeing with V. D. calculated for HBr + 511 (1); but it is not probable that the liquid consists of a true hydrate, as the composition varies with the pre sure; thus if the pressure is 1.952 mm, the constant B. P. is 153 and the liquid contains 1963 p.c. HBr (c. Roscoe, A. 146, 203). If dry nir is passed through HBrAq at a constant temperature, either HBr or H O is removed, and the composition becomes constant; at 16° the liquid finally contains 51 65, and at 100 49 35, pec. HBr (Roscoe, Lc.). The S.G. and p.c. composition of HBrAq are given in the following tables (Topsöc, B. 3, 401; Wright, C. N. 23,

212).					
Temp.	80,	P.e. HBr.	Tean	. S.G.	P.o. HDr.
11'	1.055	7:67	137	1.302	33.84
1.4	1.075	10:19	13	1.335	36:67
11	1:089	11:94	13	1 349	37.86
14	1.007	12:96	13	1:368	39:13
11	1.118	15:37	13	1:419	43/12
11	1:131	16.93	13	1:431	43.99
11	1:164	20:65	13	1:438	41 03
13	1.200	24:35	14	1:451	45:45
13	1.232	27:62	13	1.160	46:09
13	1.253	29.68	14	1.485	47.87
			14	1:490	48-17
	8.0.	Temp. 182	(Wrigh	D P.c. It	Br.
	1.030			10:4	
	F490			23.5	
	1:218			30.0)
	1:385			40.8	1
	1.475			48-5	
	1.515			40.0	

Reactions. 1. Not decomposed by heat mone, even at 706 (Hantefemille, C. R. 61, 705). 2. Decomposed by heating with many natale. eg, K.Na, Sa amrigam, Sn, Ac, with formation of metable bror ide and H. 3. Chlorine forms HCl and Br. 4. Cone, naric or sulphuric acid forms Br. H.O. and NO₂ or SO₂ 5. Lead or eilver oxide forms metallic brounde and H₂O at ordinary temperatures; picet of the other metallic oxides decompose Hitr in a similar way on warming, 6, Metallic perocides, and acids containing metals (e.g. HSbO.), form metallic bromides and Br. - 7. Agreeds solution of HBr is deconneced by nost metals with learnation of portallic brounds and II: nest notallic oxides desolve in HB(Aq forming brounds. The heat of neutralisation of HBrAq ly MOHAq or MiDHi, Aq, when 'il an alkali or alkaline earth metal, is the same us the heat of pertralisation of HCl? q. viz. 13,750; but the quantity of heat produced by the action of HBrAq on the hydrated oxides of Pt and Au, and on HgO, is much greater toan the quantity of bent produced by the action of HGIAq en the same compounds; the differences are

for AuO₂H₂ 13,810 "PtO₂H₂ 11,890 "PtO₄H₄ 16,300 "HgO 12,296.

If cone, HBrAq is distilled at 760 mm, pressure—The action of IllirAq on these hydrated oxides HBr is evolved, if HBrAq containing less is very different from the action of the same

acia on the hydrated oxides of K, Ca, Mg, &c.; in the former cases there is little doubt that acids of the form H.HgBr., H.PtBr., H.PtBr. and HAnBr, are formed in the solutions (v. Thomsen, Th. 3, 538). Many double bromides of Au, Hg, aud Pt - e.g. PtBr, 2KBr-are rather to be regarded as alkuli salts of these acids than as double salts (v. Th. 3, 417; also Gold, Mencury, Palladium, Platinum).—8. HBrAq is decomposed by bromic acid solution;

 $HBrO_0Aq + 5HBrAq = 3H_2OAq + 6BrAq$. 9. Conc. sulphuric acid hented with HBrAq forms H₂O, SO₂, and Br; dilute H₂SO₄Aq does not decompose HBrAq at ordinary temperatures. 10. Chlorine sets free Br from HBrAq. - H. HBrAq is partly decomposed by potassium permanganate solution in the cold, quickly and completely on heating. -12. By electrolysis of HBrAq, IIBrO, Aq is produced (Riebe, C. R. 46, 848) .- 13. Bromine dissolves in HBrAq forming a dark-coloured liquid.

Combinations, -1. With ammonia and phosphine; Ogier (C. R. 89, 705) gives the thermal data, [NII3,IIDr] = 45,600; [PII3,IIBr] = 23,000; neing gaseous constituents and forming solid MH Br. 2. With water probably to form the hydrate HBr.H.O (v. Properties). The heats of solution and dilution of HBr have been measured by Thomson (Th. 3, 13 and 72); the results indicate the existence in the solution of the hydrate HBr.H.O, but do not suggest the formation of uny other definite hydrate on dilution. The heat of dilution appears to be a continuous hyperbolic function of the quantity of water added, provided the composition of the acid to start with is IIBr.H.O; the results cannot, however, be expressed by quite so simple a formula, involving a single constant, as is applicable in the case of chlorhydric acid (q. v.), It is quite possible that the reactions of LiBrAq are the reactions of the acid HBr.H.O (? .. H.Br.Oil), and that HBr itself is not an neid (v. Presidential Address to Section B. by P. of. Armstrong, B. A. Meeting, 1885). Berthelot (Bl. [2] 19, 385; C. R. 61, 114; 66, 742) thinks that HBrAq contains a number of hydrates, come partially dissociated, and also the compound IIBr (v. Chromiyonac acro). Roozeboum (R. T. C. 4, 108, 331; 5, 351, 363; also Van't Hoff, ibid. 4, 414) has determined the relations between vapour-pressure and temperature of solutions of hydrated HBr. . M. M. P. M. BROMIC ACID HBrO, v. BROMINE, OXX-

BROMIDES. Binary compounds of Br with more positive elements i.e. with any element except F, Cl, or O. Br forms binary compounds with most of the elements. The greater number may be produced by direct combination. The formation of metallic bromides is usually accompanied with production of much heat; thus, [K', Br'] = 190,620; [Ca, Br'] = 140,850; [Zu, Br'] = 75,930; [Hg, Br'] = 50,550; [Au, Br3] 8,850. (Liquid Br was used.) Some metallic bromidee are formed by the action of Br on the oxides; e.g. AgBr by Br on Ag.O. Alkalis and alkalino earths in aqueous solutions are decomposed by Br, giving bromides and

bromates, certain metallic sults, in aqueous

are decomposed wholly or in part by Br, giving metallic bromides and free I. Many metallic ohlorides are partly decomposed when heated in olosed tubes to about 300° with equivalent quantitiee of Br; after a time equilibrium is established in the eystem consisting of chloride. bromide, Cl, and Br; thie equilibrium is not overthrown by increasing the mass of Br, the temperature, or the time (Potilitzin, B. 14, 1044; 15, 918; 18, 3051). Metallic bromides are usually formed by the action of HBrAq on the oxides (comp. BROMHYDRIO ACID; Reactions, No. 7). Most metallic bromides are white; they are generally isomorphous with the corresponding chlorides; most of them are not decomposed by heat alone, but some, e.g. those of Au and Pt. give up all their Br when heated. Some metallic bromides are decomposed by II,O, e.g. those of Bi and Sb; others are decomposed when their aqueous solutions are evaporated, e.g. AlBr.: most are decomposed by heating in air in presence of steam. Metallic broundes are decomposed by certain peroxides and oxidising agents, e.g. MnO2, K2Mn2O8Aq, K2Cr2OAq, HNO3Aq, with separation of Br; conc. Il SO Aq sets free little HBr, but decomposes most of the bromides to sulphate and Br with simultaneous formation of SO2; HCl and HClAq form HBr and metallic chloride. When a metallic bromide is heated with solid K,Cr,O, and conc. II,SO, free Br is obtained (distinction from chlorides). Aqueous solutions of alkali bromides dissolve large quantities of Br, probably with formation of perbroinides in solution; Berthelot (C. R. 91, 195 and 706) gives the numbers (using gaseous Br) [KBrAq, Br"] 11,500. Aqueous solutions of alkali, alkaline earth, and magnesian bromides partly decompose AgCl when the salts are shaken together for a few minutes at the ordinary temperature; the percentage of AgBr formed varies from 95 when LiBrAq is used, to 848 when CdBr, Aq is employed (Potilitzin, B. 18, 1522). The binary compounds of Br with the non-metals are fairly stable bodies; they are usually produced by direct combination. Bromides of B, P, C, and Si are stable as gases. Br forms definite, stable, compounds only with the more metallic and positive members of the oxygen group of elements; broundes of To are gusitable, Sc.Br. is fairly stable, but is decom-posed by heat, S.Br. is a feebly-marked body, and no oxide of Br is known. Bromide of iodino is a fairly well murked compound, [1, Br] = 2,500 (Berthelot, C. R. 90, 811; using liquid Br and solid 1). Bromine chloride is very easily decomposed, and no compound of Br and F is definitely known. If N bromide exists it is extremely unstable (v. also Halogen Elemente, minary compounds of; for the individual bromides v. the articles on the various elemente.) M. M. P. M.

BROMIDES, ORGANIC v. BROMINE, ORGANIO

BROMINE. Br. At. w. 79.75. Mol. w. 159:50. [-24:52] (Baumhauer, B. 4, 927), [-7-22] (Philipp, B. 12, 1421; according to Philipp, the lower M.P. was due to presence of Cl). [-7.05°] (Ramsay a. Young, C. J. 49, 453); (63°) (Pierre, A. Ch. [3] 20, 5); (59.27°) (Thorpe, solutions, form bromides and peroxides; c.g., C. J. 37, 172); (58-72) (Ramsay a Yonng, L.). salts of Mn, Ni, Co, and Pb. Metallic iodides; S.G. 2 3-1872 (Pierre, L.); 2 3-18828 (Thorpa,

Lc.), S.G. at B.P. 2-9822 (Thorpe, Lc.). V.D. | 80 (v. p. 536, Properties). S.H. (solid -78° to -20°) 08432 (Regnault, A. Ch. [3] 26, 286). S.H. (liquid 13° to 45°) '1071 (Andrews, C. J. 1, S.H.p. (equal mass of H O - 1) (835-2281) '05552 (Regnault, Acad. 26, 1): S.H.v. (equal mass of HO 1) 0429; (equal volume of air =1) 1:395 (Clausius, Mechan, Warmetheorie, 1, S.H.p. (20 | 388°) 1 293 (Strecker, 62 [1876]. W. 13, 20; experimentally determined). Expansion (0° to B.P.) V

(Thorpe, hc_0), 8, (5) 3.98, (10) 3.327, (15") 3.226, (20) 3.238, (25) 3.167, (30) 3.126, (Pancer, C, J, 15, 477). Heat of fusion . 16,185, Reguardt (A. Ch. [3] 26, 268). $\frac{1}{d}$ > At. wt. $\approx 10^{\circ}23$ (Gladstone, T. 1870, 9).

Emission-spectrum; marked lines are 3980, 6356, and lines beginning with 6999 (Salet, A. Ch. 4 24, 26). Absorption-spectrum characterised by many bands between 68015 in the red and 52414 in the green (Boscoe a. Thorpe, T. 1877, 207).

Bromme was discovered by Balard in 1826 (B. J. 7, 102); it was previously obtained by Liebig, but supposed by him to be feding chloride (v. Hofman's Life Work of Lieber); and by Joss, but regarded by him as selection (J. pr. 1, 1296.

Occurrence. Never free; widely distributed, but not in large quantities, chi-fly in combina-tion with K, Na, and Mg. In rea water (for quantities, r. Becchand, B. 18, 2888), sea-wood, saline springs, and many marine plants and unimals (Kundt a. Wöhler, P. 10, 509; Strohmeyer, S. 49, 249; Hembshidt, B. J. 7, 110). According to Marchand (C. E. 31, 495) all waters, including rain and snow, contain traces of brombles. In various minerals, chiefly as AgBr in Mexico and Chili (Bertmer, A. Ch. 77, 417; 79, 164); in minute quantities in Selesian zinc ones (Hollander, R. J. 8, 82); in Clair salt-

petre (Grineberg, J. pr. 60, 172); we, we, Preparation. The starting point is the concentrated liquor of certain value springs, the residual liquor obtained in working the salt deposits at Stassburt, or the solution of the ach of sea plants. The liquel is freed from the less soluble salts, chiefly chlorites and sulphates, by concentration and cry-tallisation, mixed with MaO, and HCiAq, and distilled. The quantities of the reacting materials are arranged so that there is always an excess of brounds, in order to prevent formation of bromine chloride (c. Mohr, A. 22, 66). In some cases the con-centrated figure is 1 at 4 with H.SO,Aq, sul-phaces are then removed by crystallisation, and the mother liquor is districtly with MoO₂ and H₂SO₆Aq (**, Herrmann, P. 13, 175; Pt, 613). The Br is condensed in water and converted into Nalle and NaBrO, by treatment with NaOHAu, the liquid is evaporated to dryne the residue heated to decompose NalirOa and the NuBr is decomposed by pure MnO, and H.SO,Aq. Iodine may be removed from the : original fiquor, before the composing by MnO, and acid, by the action of Cl, or by ppg. as Cul, (Bussy, B. J. 18, 117; Balard, B. J. 7, 102). Chloring may be removed by decomposing the B. J. 8, 83; Solly, A. 20, 121). Dissolves

distillate from the first action of MnO, and said by BaOAq, evaporating to drynoss and heating, dissolving out BaBr, in alcohol, filtering from insoluble BaCl,, evaporating to dryness, and decomposing by MnO_c and H₂SO₄A₁ (Piria, B, J, 19, 277). Adrian (J. 1870, 248) removes Cl by washing the distillate with water and small successive quantities of ether; he then digests with starch paste to remove I, and again distils. Stas (Fr. 25, 213) frees from 1 and Ci by dissolving in KBrAq, adding ZaO and distilling. Cyanogen is occasionally found in samples of Br; it may be detected by conversion into K.Fe(CN), by digesting with iron tilings, and rapidly filtering (Phipson, C. N. 28, 51). Biomoform is another impurity; it is detected by its odour, after addition of KlAq sufficient to convert all the Br into KBr, and removal of the I by the action of Na S.O.Aq (Reymann, B. 8, 790). Bromine is prepared from laboratory residues containing Br compounds by making alkaline with KOHAq, and distillation with solid K Cr.O, and excess of H SO Aq (2 parts strong acid to I part water by weight) added gradually through a famuel tube (Bolas a. Graves, C. J. (2 9, 781). To prepare pure Br for atomic weight determination, Stas removed 1 from commercial KBr by dissolving in water, adding BaAq to f of the liquid till the I which at first separated redissolved forming a clear orangevellow coloured liquid, adding the other ? of the liquid and shaking repeatedly with pure CS. The liquid was then wanned to remove all CS,; the KBr was oxidised to KBrO, by the action of CI in presence of pare KOHAq (for details v. Stas, Nouv. R. 159; or pp. 159-160 of Aconstein's German translation Univenchungen aber die Genetie der chem. Proportionen, &c.); the KBrO, was purified by repeated crystallination, and a part of it was converted back to KEr le herting in a porcebin vessel in small nucce ive quantities. By decomposing a mixture of KBr and KBrO, (in the ratioaKBr, KBrO.) with pure H SO, Aq, Br was obtained. A portrop of this I'r was then digested with milk of lime and $\mathrm{NH}_4\mathrm{Aq}$, whereby $\mathrm{CaBr_2Aq}$ was obtained ; this liquid was saturated with another portion of the purified Br; water was added to pp. Br; the ppd. Br was separated, and discated several times with joire dry CaBr, (prepared by the action of part of the joirfied Br on CaO); the Br was Jon shaken in contact with pure P.O. then allowed to remain in contact for 12 hours with pure BaO which had been strongly heated. and finally poured off and distilled. All operations were conducted in apparatus made wholly of class.

Properties. --- At ordinary temperatures a dark brown red volatile liquid with most irritating odour (Βρώμος - & tench); in thick layers almost black; vapour is yellowish red, and becomes less transparent us temperature is increased (c. Audrews, B. A. 1871. (Sec.) 66); solidities to a grey-brown crystalline mass with semi-metallic lustre. Very poisonous. Vapour acts on inucous membrane and causes violent irritation. Non-conductor of electricity; but an aqueous solution of Br is a better conductor than pure water (such a solution contains some HBr) (Balaid, A. Ch. [2] 32, 871; De la Rive,

change; solidile also in conc. aqueous solutions

Clici, solution is accompanied by chemical

heat $(Br^2,Aq) = 1080$ (Th. 2, 25); solution is pale orange-yellow. Dissolves more readily in alcohol, and in all proportions in ether, CS, and

of KBr and many other metallic bromides, fre-

quently with formation of perbromides; also in

cone. HClAq and HBrAq, and in liquid SO, (Sestini, Z. 1868, 718). Br is absorbed by wood

charcoal with considerable rise of temperature (Molsens, C. R. 77, 781). In presence of H₂O₃ acts as a bleacher und disinfectant.

At very high temperatures (approximately 1570°) the S.G. is less than the calculated; Mcyer a. Züblin (B. 13, 405) obtained values lying between those calculated from the formula Br.

and #Br2; when the Br was obtained by decou-

posing PtBr, at high temperatures the S.G. at

1570° nearly agreed with that calculated for 3 Br. (3.66). The S.G. of Br vapour diluted with 10 volumes air, at 50° under the B.P. of Br,

was determined by Langer a. Mayor to be 5-52 (B. 15, 2769). Jahn's determinations (B. 15,

1238) show that bromine does not attain the S.G. calculated for Br2 until it is heated about 160° above its B.P.; the deviations are small;

the S.G. at any temperature up to about 2200

may be approximately found by the formula S.G. = a + bt, where a = 5.8691 and b = -0.0163.

(For S.G. of Cl and I gases v. these elements: v. also Halogen Elements.) Reactions, -1. Br dissolves in water with production of heat [Br2, An] = 1080 (Th. [2] 25); the water is slowly decomposed, more quickly

The atomic weight of Br has been determined (1) by analyses, and determinations of V. D., of many gaseous computends, e.g. Brll, Br,Cd, Br,B, Br.Sn. &c.; (2) by determination of S. H. of solid Br: (3) by comparison of bromides &c. in direct snulight, with formation of HBr and O. with isomorphous chlorides and iodides &c.: Bromine water, therefore, acts as an oxidiser; e.g. HNO Aq is oxidised to HNO Aq (Schönhein, (4) by syntheses of AgDr by Marignan (B. J. 24, 193); by syntheses of AgBr by Stas (Nouv.

J. pr. 37, 141), Mu(C.H.O.) Aq to MnO. (Kämmerer, B. 4, 218); sagar, mannite, beuzene, &c., &c., to various oxidised derivatives (Blomstrand, A. 123, 248). If NO is led into Br

R. 158, 171); by reduction of AglirO, by Stas (Nouv. R. 199); by conversion of KBr to AgBr by Stas (l.c. 303); by conversion of AgBr to AgCl by Duners (rl. Ch. [3] 55, 162). In gaseous cooled below 0' the gas is absorbed, and on adding water HBr and higher oxides of N are formed (Landolt, A. 116, 177) .-- 2. Steam mixed with Br and passed through a red-hot tabe

molecules containing Br the atom of Br is monovalent. Br acts as a strongly negative non-metallic element; it combines directly with most metals forming salts; its compounds with yields IIBr and O. - 3. Hydrogen peroxide non-metals one of which is II are arids. The evolves O, and IIBr is formed (Schönbein, A. inbatitation of H by Br in earbon agids accurs to 108, 169).-4. Anneous solutions of notash or soda are decomposed by Br forming KBrAq (or increase the relative allimity of the neids

NaBrAq) and KBrO,Aq (or NaBrO,Aq): CaO and

(r. Appinity, p. 83); generally speaking, the introduction of Br in place of II in earlien compounds is accompanied by the production of DaO lorm bromides and O. ...5. Aqueous ammonia yields NH BrAq and N. - 6. Urea is decomposed

by BrAq with evolution of N.- 7. Hydriodic acid and iodides in solutions are decomposed by

1859. 67) .- 10. With carbon disulphide, crystal-

line C.S.Br. is formed (Hell a. Urech, B. 15, 987).

11. Carbon compounds are usually acted on by

Br; sometimes II is withdrawn, sometimes this

is necompanied by substitution of Br, and sometimes more complete decomposition results .-

12. Br partly decomposes metallic chlorides when

heated in equivalent quantities to 270 300°; If

the mass of Breis increased, the amount of

decomposition increases up to a limit which is

not passed by increasing the mass of Br. the

temperature or the time of action (Potilitzin, B.

14, 1014; 15, 918; 16, 3051). According to Hampidge (B. 17, 1838) AgCl is partly decom-

posed by contact with water and an equivalent

Br with formation of HBrAq, or MBrAq, and

i.- 8. Sulphuretted hydrogen yields IIBr, and S which partly combines to form S.Br.; this decomposition proceeds either in presence or absence of water (Nanmann, B. 9, 1574). -9. In contact with excess of silver nitrate solution, AgBr and HBrOAq are produced (Spiller, J.

grant-units less than that of the chloride, and about n 26,000 gram units more than that of the iodide, of the same metal, where n is a whole munher, usually 1, 2, 3, or 4; metallic

Cl and 1; the heat of formation, in solution, of a metallic bromide is usually about n 11,000

more or less acolic properties. In its chemical relations Br stands between

bromides are wholly or partly decomposed by Cl, and metallic iodides by Br; metallic chlorides

are partially decomposed by Br (v. Reactions,

No. 12). The relative utilinities of the acids

HCl, HBr, and HI in aqueous solution are,

however, nearly the same (v. Arrinity, p. 77). In its compounds, Br appears to be positive to Cl, F, and O. No oxide of Br is at present known; the oxyncids of Br exist only in pre-

sence of water; they are much less stable than the axyacids of iodine; one of the oxyacids of Cl (HClO.) has been obtained in separate and definite forms, although it is an extremely unstable compound; oxides of Cl are known as gases, and an oxide of 1 (I.O.) exists as a solid body. No oxide or oxyacid of F is known (r.

BROMIDES, HALCOEN ELEMENTS, and HALCOEN ELEMENTS, BINARY COMPOUNDS OF). The S.G. of Br gas at 99 was found by Mitscherlich to be 5:54 (air = 1) and by Meyer

quantity of Br; thus 5.2 p. c. Cl was removed from AgCl after 24 hours' action at 11°, and 14:53 p. c. alter 12 hours' action at 44°.

Combinations .- 1. With water at 4° forming a. Züblin to be 5:38 at 100°; the S.G., calen- red octahedral crystals of Br.10H.O which are

decomposed to Br and H₂O at 15° (Lowig, P. 14, 114; 16, 875). For dissociation-pressures or Rooseboom (R. T. O. 4, 65).—2. Combines directly with most of the elements, especially the metals, often with production of much heat and sometimes light (v. Browness). Does not combine directly with C or O. According to Merz a. Weith (B. 6, 1518) dry Br and Na da not combine even at 200°. Combines with liquid Cl at = 90° (Domny n. Mareska, C. R. 20, 817). No oxide of Br is at present known.

Detection.—The physical properties of Breaable this body to be easily detected when monombined. Broundes are decomposed by ClAq, giving Br and chloride of the metal. Solution of N.O. in case, H.SO.Aq does not decompose broundes, but does decompose iodides with production of 1; on this reaction is based a method for detecting broundes are decomposed by leating with KCr.O. and come. R.SO.Aq, with formation of Br; chlorides yield GO.Cl.

Estimation. Free Br may be estimated volumetrically by measuring the 1 set free by from RLAg, or by finding the mass of AgO, which it oxidises to AgO, in an alkaline relation; Br may also be combined with H to form HBrAg, by freatment with H S, or SO, in appeaus solutions; the HBr may then be estimated by pp., with AgNO, Ag. Br in broundes is usually estimated in AgNO, Ag. Br in broundes complished by addition of AgNO, Aq.; in soluble brounders may be fused with Na, LO, dissolved, and acid data. I with HNO, Aq. In presence of chlorides, or includes, Br in broundes must be determined by indirect methods (a. Manuals of Analysis).

Bromine, Chleride of. BcCl. Mol. w. nnknown. Calorine is absorbed by Br with formation of a ref yellow, mobile, very volatile, liquid, which gives off a dark yellow, very irritating, vapour with strong blenching properties; many metals burn in this vapour to chloride and bromide (Balard, A. Ch. [2] 82, 371). If the Br is could to 0' the liquid finally contains Br and Cl in the proportion BrCl; at ordinary temperatures less Cl than is required by the composition BrCl is absorbed (Bornesmann, J. 189, 183). At temperatures above + 10° the compound BrUl decomposes with evolution of Cl. By adding a little H_iO to BrCl, and cooling to 0°, yellow-brown crystals of BrCl.10H O separate (Burnemann, Lc.); these melt at 7°, and are decomposed by SH, to S. NH Br, and N chloride (Löwie, Dr. Brom und seine chemischen Perhaltaisse) Heidelberg. 1829, p. 64). An aqueous solution of BrCi, obtained by dissolving Br in ClAq, is decom-posed by alkalis giving alkali bromate and chloride; in sunlight HBrO Am and HClAq are formed; reducing agents, e.g. SO,Aq. Zn powder, Fe fings, P, NO, &c., withdraw Cl, and set Br free (Schönbein, J. pr. 88, 483).

Bromine, Cyanide of: better called Cyanogen Bromide. Obtained by action of Br on Hg(CN), KCN or HCN: P. CYANG ACRES, vol. ii. p. 313.

KCN, or HCN; r. CYANG ACHS, vol. ii. p. 313.

Bromine, Hydrate of Br.10H,O. cidained by cooling a saturated solution of Br in H₂O; Beautys: Combinations, No. I.

v. BROMINE; Combinations, No. 1.
Bromine, Iodide of; better called Bromide of Iedine; v. Iodine.

Bremine; Oxyacids of. No oxide of Br is known. The soids HBrO and HBrO, exist in aqueous solutions only; both solutions are decomposed on heating, HBrOAq even at 30°. Perhronic acid, HBrO,, was said by Kämmerer to be produced by the action of Dr gas on HCa), Aq (J. pr. 90, 190); but later experiments have shown that this acid has not yet been obtained fr. Pattison Muir, C. J. 30, 469; MacIvor, C. N. 33, 35; Wolfram, A. 198, 95). BrAq is not exidised by such rescents an K.Mn.O.Aq, K.Cr.O.Aq, HNO, Aq, or HClOAq; but the action of HClO Aq er HClO Aq produces HBrO, Aq. Dilute solutions of HBrO and HBrO, are also formed by electrolysing HBrAq. MileAq, or BrAq; also by the action of metallic oxides on BrAq. No salts of HBrO are known except in aqueous solutions; salts of HBrO, exist us solids, they are all easily decomposed by heat with evolution of O, and frequently also of Br. The addition of O to KBrAq would be attended by disappearance of much heat; Thomsen gives these numbers (KBrAq, O) also [Br'(O,Aq)] = 16,200. The following data show that the heat of formation of the oxyacids of Br is less than that of HBr, and is also less the more O the acid contains: - $\begin{array}{lll} \{H, Br_i Aq\} & 28,380 \;; & \{H, Br_i O_i Aq\} - 26,080 \;; \\ \{H_i Br_i O_i Aq - 12,120 \;(Th. \; 2, 400), & These \; \text{data} \end{array}$ are analogous with those for the corre ponding compounds of Cl, but deffer from the data for the similar compounds of I (v. Hander Elemisis).

1. Hypermomors: Acid, and Hypermomyres, HBrOAq and MBrOAq. Gay Lussue obtained a gas by the action of Br on dry HyO and supposed it to be an oxide of Br; Pelcoze, and mure recently Dancer (C. J. 15, 477), proved that only O is thus produced. An aqueous solution of iBrO i; altained by the action of BrAq on the oxide, or nitrate, of Hg, or Ag; Hg O and PbO do not oxidise BrAq (Spiller, C. N. 5, 249).

Formation. By the action of BrAq on HgO repeating several times HgOBr, HBrDAq, and HgBoAq, are formed; by di-filling in vacuo, a liquid containing 62 pre, Br as IIBrO is obtained, but it cannot be quite freed from HgBrO (Bidard, A. Ch. 32, 337).

Preparation.—Pure BrAq is shaken with excess of ApNO_Aq until the colour and adout to Hr have gone; the straw coloured liquid is then at once distilled in racuo; at 50 mm, pressure it holds at 40?. The distillate gets richer in Hino until 736 p.c. Br is present as Hiro (then it gets peerer in the acid); about 46; p.c. of the Br used is changed to Hiro.

Properties and Reactions. Solution with 6c2 p.c. Br as IIIsO decomposes at 30° giving Br and IIIsO Acq. solution with 736 Br as IIIs O decomposes, into o one products, at 60°. IIIstO q is a strongly bearding liquid; it is decomposed by AgO (and slowly by contact with AgNO, Aq) with formation of O and AgBr, by II,O, Aq with evolution of O; and by IIClAq, IIBAq, and IIIAq, with formation of Br (Schonbein, J. pr. 88, 475).

No hypotromites have been obtained except

No hypobromites have been obtained except in dilute aqueous solutions. By the action of alkalis, alkali carbonates, or phosphates (Fritzsche, A. 40, 251), on BrAq, yellow, strongly bleaching, liquids are produced; these liquids are very unstante, decomposing in air, quickly at 30°. They decompose ures with evolution of N;—CON,II;+3HBrOAq=CO₂+N;+3HBrAq+2II;OAq (E. Kuop, C. C. 1870, 132). Balard (A. Ch. 32, 337; J. pr. 4. 165) described bodies re-embling bleaching powder, obtained by the action of BrAq on CaO.H. and BaO.H.; by the addition of water and filtration, beaching solutions were produced, very unstable, and easily decomposed, even by Co.,

H. BROMIC ACID AND BROMATES, HBrO.Aq and MBrO_s. Broinic acid, HBrO_s, is known only in aqueous solution.

Formation. 1. By electrolysis of HErAq (Ricke, C. R. 46, 348). 2. By action of heat

on hypobromites in solution.

Preparation. BaBrOs is prepared by adding Br little by little to cone. BaO,H,Aq imil the liquid is slightly red, when BaBrO, pps. aml BaBr, remains in solution. The BaBrO, is crystallised from hot water, dried, and ground to fine powder; 100 parts are digested for some time in the cold or at a very gentle heat, with 24 parts cone. H.SO, mixed with 240 parts H₂O; excess of H₂SO; is removed from the liquid by gradual addition of BaO₂H₂Aq; the acid liquid is evaporated by vacuo (Rannaelsberg, A. 40, 117). Kämmerer passes CLO into Br under H.O until the colour of the Br has disappeared; 5Cl,O + Br,Aq + H,O ≈ 2HBrO₃Aq + 10ClAq (J, pr. 85, 452),

Properties and Reactions, - HBrO Aq may be concentrated in vacue until the liquid contains 50 59 p.c. HBrO,; the composition then nearly agrees with the formula liBrO, 711,0. When concentrated by heating in an open vessel decomposition begins when the liquid contains 4:27 p.e. HBrO, with production of Br, O, and H.O. HBrOaAq is a colourbess, acid liquid, with a bromine-like smell. Oxidisalde bodies decompose HBrO₃Aq with separation of Br; c.g. $58O_2 + 2HBrO_3$ Aq + 4HLO = 5H SO₂Aq + Br Aq; $5H_2S + 2HBrO_5Aq = 6H_2OAq + 6S + Br_2Aq$. To loff₂σ (24) 101 (5) at 7 (14) and 7 (14) at 7 (14) H2O and Br; HClAq and H1Aq form H2O and BrCl or 1Br. The heat of neutralisation of HBrO Aq is the same as the mean value for the stronger monobasic acids; [HBrO A], NaOHAq] - I3,780 (Th. 1,240). Bromie acid is monobasic , and forms one series of salls, the bromates, hydroxides, or earbonates, of the metals; the a kuli and alkaline earth salts are also formed. niways with brombles, by the action of Br on aqueous solutions of the nikuli or nikaline earth hydroxides. (For special methods r, individual salts; also Kämmerer, J. pr. 85, 452.) The bromales crystallise well; they are all soluble in water, the least soluble are AgBrO, and They are decomposed by heat; sometimes O is evolved and metallic bromide remains, e.g. bromates of alkedis, Hg, and Ag; sometimes Br and O are evolved, and oxide remains, e.g. bromates of Mg, Al, Zn; or a mixture of oxide and bromide remains, e.g. bromates of (Kremers, P. 99, 443). S. (6°) 3·1, (20°) 6·9, Pb. Cu, &c. Dilute HNO,Aq, H,SO,Aq, or (40°) 13 2, (100°) 50 (Kremers, P. 97, 1). Insol.

H,PO,Aq, decomposes bromates giving HBrO,Aq most of which decomposes to Br, O, and H.O. Solutions of bromates react similarly to solution of HBrO, towards SO, H2S, HClAq, HBrAq, and HIAq. The bromates have been chiefly investigated by Rammelsberg (A. 40, 147; P. 90, 16); Löwig (B. J. 12, 120); and Marignac (C. R. 45, 650; J. 1857, 127). The following are the better-studied salts.

Barium bromate Ba(BrO.) ... H.O. matic, monoclinic, crystals; isomorphous with Bu(ClO₃)₂ (Marignac a. Rannuelsherg); S. (100°) 4·2; (15°-18°) 77. Prepared by decomposing

KBrO, Aq by Ba(C,H,O,) Aq. Cadmium bromate Cd(BrO,), H,O. lumnar trimetric crystals; prepared by CdSO₄Aq + BaBrO₄Aq. S. (15°-18°) 125. On heating, leaves CdO and CdBr, (Rammelsberg),

Calcium bromate Ca(BrOs), H.O. Monoclinic plates (Marignac); prepared by CaO..H.Aq + HBrO.Aq. S. (15°-18°) 99, CaO.H.Aq + 11BrO.Aq. S. (15°-18°) 99. Heated to 180° loses H.O, at higher temperature gives O and CaCl (Rammelsberg).

Cobult bromate Co(BrO3), 611,0. Trans. parent, hyacinth-coloured, monometric octahedra; prepared by HBrO,Aq + CoCO, Bu(BrO₃), Aq + CoSO₃Aq. S. $(15^{\circ}-18^{\circ})^{\circ}$ Heated, gives CoO (Rammelsberg).

Copper bromate Cn(BrO3),5H,O. green crystals, efflorescing over H.SO, in vacuo; very soluble; lose 11.0 completely, and a little Br also at 200°. Prepared similarly to Co(BrO₃)₂,

Lead bromate Pb(BrO3),11 0. Instrous, monoclinic prisms; isomorpheus with Sr(BrO₃) ₂H₂O (Rammelsberg). S. (15° 18°) 1·33. Prepared by HBrO₂Aq + IBCO₃, or cone. Pb(C₄H₃O₄),Aq + KBrO₂Aq. Heated over 180° gives Br, PbO, and PbBr, at higher temperatures gives Pb₂O₄, Br, and PbBr₂.

•Magnesium bromate Mg(BrO.) .. 6H.O. Large, efflorescent, monometric octabedra; S. (15°-18') 71.5; loses all H₂O above 200°. Prepared by MgO + HBrO3Aq, or

MgSiF₆Aq + KBrO₃Aq.

Mercury bromates Hg2(BrO3)2, and Hg(BrO_s), 2H_sO. Merenrous bromate is prepared by Hg (NO₃),Aq + KBrO₃Aq or HBrO Aq + Hg O; mercuric bromate by HBrO_sAq + freshly ppd, HgO. Both form small white crystals; the increurons salt forms yellow basic Hg,(BrO,), Hg,O by the action of 11,0; when heated it decomposes with dctonation (Rammelsberg). The mercuric salt decomposes at 130°-140°, with slight explosion, to HgO, HgBr, and HgBr, S. (15°-18°) 17, (100°) 1·6.

Nickel bromate Ni (EvO3), 6H2O. Prepared *as, is isomorphous with, and generally resembles, the Co salt (q. v.). S. (15°-18°) 28. (Rammelsberg; v. also Marbach, P. 91, 412).

Polissium bromale KBrO. Prepared by adding Br, or BrCl, to conc. KOHAq; or by adding Br to K.CO.Aq which has been previously saturated with Cl. Rhombohedra (Rammelsberg; Marignac, J. 1859. 139; e. also for crystalline forms, Löwig, B. J. 12, 120 ; Fritzsche, A. 40, 251; Marbach, P. 94, 412). S.G. 17 10 3 271

in alcohol. B. P. of saturated KBrO, Aq = 104°. KBrO, when heated, melts at 350°, then begins to decompose, at first slowly, then rapidly and explosively, with evolution of O and a little Br. According to Fritzsche (A. 40, 251), crystals of KBrO, prepared from exactly neutral solu-tions, or from solutions containing a little acetic acid, decrepitate before decomposing, and lose 13 p.c. of their mass phiefly water); if the resulting powder is placed in warm water, O is evolved at the surfaces of the undissolved particles, but most of the O is negling bearbed by the liquid; on evaporation, pure KlirO, crystallises out. Fritzsche supposes that KBiO, is partly decomposed by heat to hypobromite and perbromate, that the latter acts on water, evolving O and forming KBrO, Aq, and that the O is absorbed by the KBrOAq with formation of KBrO, Aq. KBrO, is decomposed by cone. H2SO, Aq, with evolution of O and Br (Liewig); by HNO, Aq, with production of KNO, Aq, Br, and O (Penny, A, 37, 206). KBrO, Aq decomposes H.S. separating S, and forming HBrAq, and 11.SO,Aq. Heated with combustible bodies, KBrO, evolves O rapidly and explosively.

Silver bromate Ag BrOs Dimetrie prisms (Marignae). Propored by ArNO, Aq e HBrO Aq. or KBiO, Aq. and crystallising from hot water. Stable in air free from organic matter. Decom-

poses on heating.

Sed am browate NaBrO₈. Prepared as KBrO₈. At 4° crystallics with xH O forming efflere wint needles (Lowig); above 4^2 the anhydrous salt separates, isomorphius with KBrO, according to Lowie (B. J. 12, 120). $S_iG_i = 3.339, \quad S_i(0^\circ) 28, (20^\circ) 38\%, (60^\circ) 62\%,$ (160°) 59 (Keeniers, P. 97, 1). B. P. of suturated NaBrO Aq = 109". Decomposes when heated as KBrO, (q. v.).

Street from bromate Sr(BrO.)...11.0. clinic priems. Is amorphous with the Us falt (Barminch Ferg), S. (15° 18) 33. (160 8 11,0) at 120°. Prepared by SrCO, 4 HBrO, Aq. Zine brownits Zn(BrO,) 5H,O.

metric octahedra, i outorphous with the Mig salt; prepared as Co(BrO₂)...6H₂O. Molts at 100%; loses 6H₂O at 200 with partial decomposition to ZhO, Br, and O. S. (15" 18) 100. Soluble in NII, Vq (Rananelsberg).

Besides the foregoing, bromates of Ce, In, and Dr of the form Millro,) .611 O have been prepared (Bammelsberg, Matiguae, Hermann J. pr. 82, 385). Promates of Al, Cr. Fe, mel V; of Pd, and Pt; of Bi; and of Sn. 10 m to They have, however, either not been ; exist. obtained in definite fram, or have been very slightly examined.

whilly examined.

Bromine. Sulphiet of: better called Sul
Bromine (e. Schenne). M. M. P. M. .. phur Bromide (v. Scarners). BROMINE, ACTION ON ORGANIC BODIES

B. BROMO COMPOUNDS.

BROMO .. Use of this prefix applied to inorganic compounds: for brome compounds and brome-salts v. the element the brome compound of which is sought for, or the saits to the names of which bromo is prefixed. Thus bromochloride of carbon will be found under Carson; bromo-chromate of potassium under

BROMO-ACENAPHTHENE v. ACENAPHTHENE.

BROMO-ACENAPHTHYLENE . ACENAPH-THYLENE

BROMO-ACETAL e. BROMO-ACETIC ALDERIYDE. .BROMO-ACETAMIDE e. BROMO-ACETIC ACID. v-Bromo-acetamide v. Aceto-bromo-amide, p. 5. BROMO ACETAMIDO- e. Bromo amido-, BROMO-ACETANILIDE & BROMO-ANHINE. BROMO-ACETIC ACID CH,BrO.

OH Br.CO H. (51°), (208°).

Formation .- I. By bromination of acetic acid (Perkin n. Duppa, A. 108, 106; Hell a. Mohlhauser, B. 11, 211; 12, 7, 5). 2. By heating ethyl acctate with because at 150° (Crafts, A. 129, 50). - 3. Prom plycollic neid and HBr (Kekulé, A. 130, 11), ... t. By atmostdarie oxidation of an alcoholic rolation of brome ethylene (Glöckner, A. Suppl. 7, 107). 5. By the action of bromine on dry glycerin (Barth, A. 124, 341). 6. From rhloro acetic ucid and HBr (Demale, R. 9, 560). 7. From ethylene bromide and furning HNO, (Kachler, M. 2, 259).

Preparation. - Br. placial acetic acid, and some CS are boiled with inverted condenser; the yield is nearly theoretical (Michael, Am. 5, 202). Properties. Deliquement rhombohedra; v.

sol, water. Blisters the skin,

Reactions. 1. Heated with zinc it yields Zn(OAc), and ZnBr₀, 2, NH, forms glycocoll, 3. Sitter beneate forms glycollide, benzoie neid and Arlie. 4. Silver powder at 130° forms streetine arid. 5. Ferryl sulphale (C₃H₂)S forms benryl bromide and S(CH₂CO₃H), (Letts, Tr. E. 28, 642). Allyl sulphide acts similarly. 6. Bromo acche neid and its ethyl salt unite directly with Me S and its hom logues, forming hydrodrounds of 'thetines' (Crom Brown a. Lette, B. 7, 695).

Salts. The NH, K, Ba, and Ca salls are v. sol. water. PbA': lamine, sl. sol, cold water. ApA': erystalline; explodes at 90",-Witch Ap 1 (Clarke a. Owens, D. 11, 35).
Methyl ether Med. (141) (P. a. D.).

Ethol ether EtA'. (170'). Converted by Na interaction ther ther (v. p. 2), Chlerotte describer CH₂ClCH₂A', (214).

S.G. 92 1 65. From chloro cthyl chloro acctate and Br (Henry, C. R. 97, 1308). Decomposed by heating with water into glycol chlorhydrin and brome acetic acid.

Bramo-ethyl ether

CH ,CHBr,O.CO.CH Br. (135°) at 370 mm. S.G. 3º 1962. From CH, CHCLOAc (c. p. 105) and Br at 1002 (Kersel, 7) 10, 1999; 11, 1916). Oil. Boiling alcoholic KOH forms branco acctic ether. EtBr, acetal, and crotonic ablebyde.

Di-bromo cthyl ether

C.H.Br. O.CO.CH.Br. A non volatile oil, ohturned by heating the preceding (I mot) with Br (1 mol.) at 120°.

Tri bromo ethy : ther

C.H Br., O.CO.CH Br. An oil found by heating bromo (thyl bromo ac tag (I mol.) with Br (2 mols.) at 160 .

Tetra bromo ethnil ether

C.HBr. O.CO.CH Br. (177'). From the preceding (I mol.) and Br (I mol.) at 170°. Decomposed by alcohol into little and bromo- and di-bromoacetic ethers.

Penta-bromo-ethyl ether C.Br. O.CO.CH Pr. (c. 197°). Formed by brominating the preceding.

secomy tether O.H.A. (2019). Chloride v. BRONO-ACRTYL CHLORIDE.

Bromide v. Bromo-Acetyl Bromide. Anhydride (CH2Br.CO)2O. (245°). tained by distilling Ac.O.CO.CH, Br which is

formed by the action of AcONa on Br.CO.CH, Br (Naumann, A. 129, 273; Gal, C. R. 71, 273).

Amida Cll, Br. CO. NII, [165°]. From bramo acetic other and 20 p.c. NII, at 0° (Kessel,

B. 11, 2116). V. sol. water, sl. sol. alcohol,

insol. other.

Nitrile CH, Br.CN. (1499). S.G. 12 1.771. V.D. 4 06. Brone-acctonitrile is formed by the

action of bromme-water on iodo-acetonitrilo (Henry, C. R. 103, 413). The dibromido of

acetonitrile (p. 35) may be looked upon as a

imide (CH,Br.CO),N11 [98] gradually separato (Engler, A. 133, 137; 142, 69).

Di-bromo-acetic acid Clltir, CO H. [45% 50%]. (283°). Formed, together with C11,Br.CO,11, when a mixture of bromino and acctic acid is exposed to sunlight (Perkin a. Duppa, C. J. 11, 22; Schäffer, B. 4, 368). Formed also by the

action of Br on acctic other at 130 ' (Carins, B. 8, 836), and as a by product in the preparation of tri-bromo acetic aldehyde by passing bromine-vapour into alcohol. Crystalline mass, v. sol.

alcohol and ether; its vapour is very purgent. Sutts. Excepting Ag and mercurous salts, the dibrome-acctates dissolve readily in water. NH,A'. KA' aq. BaA', Jaq. BaA', Saq (Benedikt, A. 189, 160).—PhA'₂: stellate groups of nsedles,—AgA': needles; decomposed by boil-

ing water into AgBr, glyoxylic acid, and dibromoacctic neid; decomposed by ether at 100 into Agllr and an oil, C. H.Br.O., whence water forms di-bromo-acetic and glyoxylio acids (Perkin, C. J. 82, 91).

Ethyl other CHBr., CO.Et. (193°). Formed by heating the acid with alcohol or by adding bround hydrate (4 pts.) to an alcoholic solution

of KCy (1 pt.) (Remi, J. R. 7, 263). Amide Cillir, CO.NH. [156]. Formation.

-1. From di bronno acetic ether and NII, (Schäffer, B. 4, 369; Kesset, B. 11, 2116). 2. From penta bromo aceto acetic amide CBr, CO,CBr, CO,NH, and boiling water (Stokes

a. V. Pechmann, Am. 8, 375). 3. From pentabromo acctone and N11, (Ctocz, J. 122, 121). 4. From asparagine and teromine (Guareschi, B. 9, 1435). Properties, ... Needles; may be sublimed; m. sol. cold, v. sol. hot, water, alcohol, and ether,

Nitrile CHBr., CN. [112]. Formed, together with bromoform and CO, by the action of Br on aqueous evano neetic acid (Hoff, B. 7, 1571). Trl-bromo-acetic acid Clir, CO.H. (8.); (135") (Gal, C. R. 77, 786). (250").

Formation. 1. By the action of water on tri-bromo-acetyl bromide. - 2. By heating aqueous analonic acid with bromino (Petrieff, B, 8, 730). 8. By heating tri-bromo-acetic aldehyde with

HNÖ, (Schiiffer, B. 4, 370). Properties. Mouoelinic tables, v. sol. water;

its vapour is pungent. Decomposed by boiling, giving off Br and HBr. The salts, excepting the silver and mercurous salts, are v. sol. water and alcohol, but decomposed by heat, both when dry and when in solution, into bromoform and a metallic carbonate.

Balts. Maa'2]aq: lustrons lamins. BaA', Saq: tables. PbA's stellate groups of needles. - AgA': very unstable lamine.

Ethyl ether Eth'. (225°). Amide CBr., CO.NH, [12]

[121°]. Formed. together with the preceding by the action of bromins on asparagine suspended in water

(Guareschi, G. 6, 375). Formed also by treat. ing hexa-bromo-acetone with ammonia (Weidel a. Grüber, B. 10, 1148). Monoclinic lamine; may be sublimed; sl. sol. alcohol, other, and cold water. Split up by boiling alkalis into bromo-form, CO., and NH.,

BROMO-ACETIC ALDEHYDES. Bromo-acetic ortho-aldshyde.

Ethyl ether CH_Br.CH(OEt) ... hydrobromide of bromo-acetonitrile; when water, acctal. (1712). From acctal and bromine is added to its alcoholic solution needles of the (Pinner, B. 5, 119) or by treating vinyl ethyl oxide with Br and decomposing the product CIL Br.CHBr.OEt with NaOEt (Wislicenus, A. 192, 112). Alcoholic KOII at 170° converts it into

Cit,Otl.Cit(OEt),, while NaOEt forms at 100° CII,(OEt).CII(OEt)... Di-bromo-acetic aldonyde CHBr2.CHO. (142°). Formed by dropping Br (2 mols.) into a solution of paraldehyde (1 mol.) in acetic other (Hagemann, B. 3, 758; Pinner, A. 179, 67). Liquid, v. sol. water and alcohol. Blisters the skin. Stowly changes to an isomeric modification, insol. water. Combines with water, forming

the solid hydrate CliBr, CH(OII), Tri-bromo-acetic aldehyde CBr. CHO. Bromal. Mol. w. 281. (1747). S.G. 3:34. Formation .- 1. By the action of Hr on alco-

hol (towig, A. 3, 280; Schiffer, B. 4, 366). 2. Together with the preceding, by this bromina-

tion of paraldehyde. Properties. Pungent liquid; decomposed by aqueous alkalis into bromoform and formic acid. Reactions. -- 1. Funning HNO formstri-bromo. acetic acid.—2. Alcoholic KCy forms in bromo-acetic ether and 11Cy (Remi, B. 8, 695).— 3. Conc. 11.SO, forms bromalide C.H.Br.O.

tri - bromo - ethylideno tri - bromo - lactate ,0.CO CBr_rCH([1580]. This body is ∖o.ċп.cвr, also formed by the action of tri-bromo-lactio acid on bromal (Wallach, A. 193, 52). It is insol. water, and decomposed by alcohol.— 4. Tri-chloro-lactic acid forms the corresponding

,0.CO [150°].-5. Lactic acid 0.CH.CCL

forms CBr.CH([97°] (Klimenko, o.cu.ch, B. 9, 968). Combinations .- 1. With water: Tri-bromo-

acetic orthoaldehyde or Bromal hydrate CBr, CH(OH), [54°]. Crystallises on evaporating an aqueous solution of bronal. Decomposed by distillation into H.O and bromal.

2. With also hol: CBr_s.CH(OII)(OEt). Bromal alcoholate [44°]. Thick needles; sl. sol. water; resolved by distillation into its components. - 3 With sodium biaulphito CBr, CH(OH).SO, Na: laminm .-- 4. With ammonia CBr. CH(OH).NH. crystals, insol, water; decomposed at 350 (Schliff a. Tassinari, B. 10, 1786) .- 5. With acetamide:

CBr. CH(OH).NHAc. [160] (S.a. T.) .- 6. With

BROWIDE.

BROMO-ACETIC CHLORIDE r. BROMO-ACETYL GRIORIDE BROMO-ACETIC OXIDE v. Anhydride of

BROMO-ACETIC ACID. BROMO-ACETO-ACETIC ACID ather C.H.Bro, i.e. CH, CO.CHBr.CO Et. S.G. 28 1-511.

Formation. By addition of Br (I mol.) to an ethereal solution of aceto acetic ether (Duis-

berg, B. 15, 137.; A. 213, 138). Properties.-Pangent brown liquid; al. sol. water, v. sol, ether and alcohol. Gradually decomposes on standing, giving off IIBr. Fe Cl. turns its aqueous solution deepered. It is dis-solved by baryta-water, and on adding Cu(OAc), the solution gives a sup-green crystalline po-Cu(C, II, BrO), which on recrystallisation from alcohol or CS, separates as dark-green needles,

Reactions. 1. Alcoholic NII, forms sucoinyl-succinic ether. 2. Na added to its ethereal solution also produces succinvisuccinio ether (dilivdride of di-oxy-terephthalic ether). 8. NaOEt produces succentyl-succenic neid (Wedel, A. 219, 92). These reactions may be represented thus; 2011, CO, CHBr.CO Et

CH CO CH CO Ex

= 211Br + CO2Et.CH.CO.CH.

CH;C(OH).CH.CO,Et

CO,Et.CH.C(OH);CH

Anilide CII, CO.CHBr.CO.NHPh. [138 Bronine added to the audide of aceto-acetic ether in chloroform forms an additive product which on warming yields bromo-aceto-acetic anilide (Knorr, 4, 236, 79). Pearly plates (from alcohol); sl. col. water. Cone. II SO, produces bromo-oxy-methyl-quinoline.

Methyl-bromo-aceto-acetic ether CH, CO.CMeBr.CO.Et. Obtained by bromination of methyl-actto acctice ther. Liquid. Converted by heat into EtBr and C.H.O. (Pawlow, C. R.

Ethyl bromo-acoto-acetic ether

CH., CO.CEtBr.CO Et. S.G. 1-35. Obtained by adding Br to ethyl aceto acetic ether in ethercal solution (West I. A. 219, 102). Pumpent liquid. Fe₂Cl₆ turns its alcoholic colution violet. 100° it gives EtBr and C, H, O, or C, H, O, probably CH CO.C. CO.H. CH.CH. (cf. Demarcay, A. Ch. [5, 20, 105).

Iso-butyl-bromo-accto-acetic other CH, CO.C.(C. H., Br. CO. Et. From iso-butyl-aceto-acetic ether med 15 at -5 (Demanyay, Bl. [2] 31, 513; 33, 516; A. Ch. 5, 20, 433; G. R. 86, 1085, 1135). Alcoholic KOll convert? it, according to Demarcay, into hexoic, methylisobutyl-glyceric, heptoic, exphaptoic, and glycollic acids; according to Pawlow (C. R. 97, 99) alcoholic KOII forms an acid C.H.O. or CH, CO.C(CO.H): CH.Pr with evolution of Ethr.

Di-bromo-aceto-acetic ether CH, CO.CBr, CO, Et (?), S.G. 25 1-884. From acetoacetic ether (10 g.) in ether (10 g.) treated with Br (24.6 g.) (Duisberg, A. 213, 143). Fungent oil.

Reactions .- I. Aqueous Fe Cl. gives a deep

athyl carba mate: CBr. CH(OH).NH.CO.Rt. red colour.—2. Baryta-water gives a claret bronial-wrethous [182°] (Bushoff, B. 7, 632).

BEOMO-ACETIC REOMIDE v. BROMO-ACETIL tion. A sap-green pp. Cu(C,H,Br,O,), is, howcolour, but Ca(OAc), gives no pp. in this solu-tion. A sap-green pp. Cu(C,H,Br,O,), is, however, formed when cone. Cu(OAe), Aq is added to an alcoholic solution of the ether; it crystallises in needles (from alcohol).—3. Diluted with other and treated with Na. di bromo accto-acetic wther (SOp.) gives disoxy-terephthalic other (2 g.). NaOLi may be used instead of Na (Wedel, A. 219, 74).

Ethyl-di-bromo-aceto-acetic eth r C.H.Er.EtO, &c. CH.Br.CO.CBrEt.CO Et (2). 8.6. 164. A yellowish oil, got by bromination

of ethyl aceta acetic ether. Fe Cl. colours its alcoholic solution wine red (Wedel, 4, 219, 102). Pr Cl. colours its

Tri-bromo-aceto-acetic ether GILBr.CO.CBr.,CO.Et. 8.G. 83 2:114. From to to acetic other (10 g.) in chlor dorm (20 g.) by addition of bromine (38g.) (Dunberg, A. by Addition of bromine (38g.) (Dunberg, V. 213, 145). Yellow bound, slightly junepont. sl. sol. water. When heated, it gives off IIBr.

Beretions, - 1. Aqueons Fe Cl. rives after some time a slight red pp. 2. Gives in alcoholic solution with eneric acctate a green pp. $Co(C_sH_sBr_sO_s)_{\mathcal{F}}(W_t|\det_s^{\bullet}\mathcal{A},|249_t|95).$

Ethyl-tri-bromo-acoto-acetic ether C.H.Br.EtO, i.e. CHBr.CO.CBrEt.CO.Et (?). S.G. 186. Its alcoholic solution is turned winered by Fe Cl.,

Tetra- and Penta-bromo-aceto-acetic ethors. so called, are mixtures of tri- with per-bromoaveto neetle ether.

Penta bromo-aceto-acetamido CBr, CO.CBr, CO.NH. [c. 148]. From di-oxyamido pyridine and bromine water (Stokes a. V. Pechmann, Am. 8, 375). Needles or prisms; converted by boiling water into di brono-acetamide and CHBr. Alcoholic XII gives di bromomalouamide and CHBr.,

Per-bromo-aceto-acetic ether C_Br_inO, i.c. CBr_wCO.CFr_wCO.C.Rr_w [69° 70°. From hi-bream metage etic ether and broming at 80° for 2 mays. Colourless crystalling mass. Not affected by air. Gives no colour with Fe Cl, and no copper pp. (Wodel, A. 219, 97).

BROMO-ACLTO-AMIDO- v. Brown compo.

TRI-BROMO-ACETO-GUANAMIDINE Call BraNaOa. From bromine and squeous a etogoanamide (Nencki, B. 9, 236). Minute receibes, insal, water, alcohol, and ether. Split up by beiling with water into bromoform and eyanoric acid.

BROMOGACETOL r. Di-Brosio (Borani

BROMO-ACETONE CIL, CO.CH. Br. S.G. 1-99. Formation. I. By adding 1 mot, bromine to acctone, rather pure or mixed with water or with CS (Linnemann, 4, 125, 307; Lannerling, H. .. 6, 22). 2. By the action of a weak electric current on a mixture of acctone and HBr (Riche, C. B. 49, 276).

Preparation. 1. A or am of dry air saturated with bromine (138g.) is passed through 100 g. cooled acctone (Kinn, riing a. Wagner, A. 201, 27). 2. Bromine (1 mol.) is passed through a solution of 1 mol. acctone in 10 vol. water (Sokolowsky, B. 9, 1687).

Properties. - Yellow, very pungent-smelling oil, quickly turning violet when dry, more permanent when mixed with a little water. oxidises it to formic and acetic acids (Linnemann, Sitz. B. 68, 137). Aqueous K.t.O. forms a syrupy sold O_1H_1U , (7) Bromo-scetone com-bines with NaHSO₂.

u-Di-bromo-acetone CH, CO.CHBr, S.G. 2.5. Prepared by adding bromine (2 mols.) to acetone (1 mol.) mixed with a large quantity of water.

Liquid, volatile with steam. Not very pungent. Combines with NaHSO3.

s-Di-bromo-acetone CH.Br.CO.CH.Br. [24°], From di-jodo-acctone and AgBr at 150°; or

trom di chloro-acetone and aqueous KBr. Long needles with pangent smell (Völker, A. 192, 96).

Reduced by Zn and H SO, to account.

Tri brome acetone Clar CO CH. Formed by the action of alkalis upon hexa-brome-methylmethenyl di-ketone. NII, gives bromoform

(Combes, A. Ch. [6] 12, 211). Tetra-bromo acetone hydrate Callelle, O 2aq.

[43°]. From Ba (10 pts.) and acctone (I pt.) in the cold; when the resulting solid mass is recrystallised from dilute alcohol a mixture of tetra- and penta-bromo-actione is first deposited, ! and afterwards prisms of tetra bromo-nectoric hydrate Gall, Br.O 2aq. The hydrate is insol.

water, sol. alcohol (Mulder, J. 1864, 330). Perhaps this body is (C.H.Br.O)HOEt.

Penta bromo-acetone C. ItBr O. [76°]. Formation. I. From Br (12 pts.) and acetons (1 pt.) (Mulder, J. 1861, 330; if. Steiner,

B. 7, 505, 1281). 2. Separates on adding water to an alcoholic solution of 'phlorobromin,' that has stood some time (Benedikl, C. C. 1878, 101; A. 189, 168).- 3, From potassic cirriconute and Tr (Cahours, A. 64, 351; Grimaux, J. 1874, 522) .- 4. From chelidonic acid and Br (Wilde, A. 127, 167) .- 5, From agreous pyrnvic acid and Br at 100° (Wichel-

haus, A. 152, 260). Properties. - Trimetrie needles (from dilute alcohol); a:b:c = 6:18:1: 686 (Ditscheiner a. Friedlamher, Z. Kryst. 3, 103). Converted by aqueous or alcohotic NH, into di-bromo-acet-

amido. Aqueons KOH forms broun form, HBr. CO, and formic neid. Hexa bromo acotone CBr, CO.CBr, [112]. Formation .- I. By the action of Br on an

chloride or hydroloromide, or on bromo diohromazin' the first product of the action of bromine on these salis. 2, From brommilie acid and Br (Huntzsch a, Schnifer, B. 20, 2010). 3. From di amido-guniacol hydrochloride and Br

(Herzig, M. 3, 825). Properties. - Monoclinic prisms (front chloro-

form) insol, water. Decomposed by alcohol.

Reactions. 1. Boiling NatollAq or water at

180° form bromeform and CO. 2. Boiling conc. IINO, loss no action, but at 150° it produces bromo pierin CBr, NO, ... 3. Pry NII, gives rise to tri-bromo-acetamide and CBr, H. 4. Sodium

malgam reduces it to iso-propylalcohol.

BROMO - ACETONITRILE v. Nitrile or Ввомо-аскти астр.

a.BROMO ACETOPHENONE C.H. BrO i.e. O.M. CO.CH.Br. Thenyl bromo-methyl ketone. Phenacyl bremide. [50'].
Formation.—1. From acctophenous and Br

Emmerling a. Engler, B. 4, 118; Hannius, B. 0, 2006; Stacdel, B. 13, 837),—2. Formed,

ogether with CO, and HBr, by boiling a-phenyloxy-8-libromo-propionic acid with water Böttinger, B. 14, 1238).

Preparation.—1881 pts. of momine are slowly run into a mixture of 100 pts. of acetophenone and 500 pts. of glacial acetic acid: After stand-

ing an hour it is gently heated on a water bath till colourless, when it is at once poured into a large quantity of cold water; the yield is 133 pts. or 80 p.c. of the theoretical (Möhlau, B. 15, 2464).

Properties .- Trimetric prisms (from dilute alcohol); pungent; v. sol. alcohol and ether. insol, water.

Reactions. - 1. KMnO, forms benzuic acid.-2 Alcoholie NH₃ forms iso-indole C_sH₁N_{*}-

3. With sodie-malonic ether it forms C. H. CO.CH. CH(CO.H) .- 4. With socio-aceloacetic ether it forms acetophenone aceto acetio ether (v. p. 36) .- 5. Aniline forms phenylamido-acetophenone (Mülilau, B. 14, 172). 6. Reacts with hydroxylamine hydrochloride with production of iso-nitroso acctophenone-oxim, C.H. C(NOID.CH NILOH [1633] (Schramm, B.

16, 2183). - 7. An alcoholic solution of phenyl hydrazine reacts thus: I'ii.CO.CH Br + N.H.Ph = 11Br + 11,0 + 19i.C.H.,N.,Ph. The = IIBr + II,O + IIh,C,H,N,Ph. The product forms yellow needles (from alcohol) [1372]. It is very soluble in other, chloraform, and CS., less in abohol or light petrolenm. composed by acids (O. Hess, A. 232, 234).

w-Di-bromo-acetophenono Calla CO.CHBr. [374]. Prepared by adding bromine (2 mols.) to a cold solution of acetophenoue (1 mol.) in acetic acid, warming to 65°-70°, and pouring into water; the yield is c. 80 p.c. of the theoretical

(Engler a. Hassenbaum, B. 18, 2240; cf. Hunnius, B. 10, 2010; Fittig a. Wurster, A. 195, 161). Properties, Trimetric tables (from CS.); insol, water, Oxidised by KMnO₁ to benzoic achl.

Fractions. 1. Alcoholic KUAc Ph.CO.CII(OAc) - 2. Alcoholic hydroxylamine forms phenyl-glyoxim Ph.C(NOH), CH(NOH) [1523],- 3. By treatment with NII, a portion breaks up into benzamide and Cti Br. who snother part yields is sindilencing C_1, H_1, N, O_2 whilst w.TRI - BROMO - ACETOPHENONE - o. CAR-BOXYLIC ACID CBr, CO.C. H, CO.H. [160].

aqueous solution of triumido phenol hydroa'kalis into CHBr, and phthalic acid (Gabriel a. Michael, B. 10, 1551, 2199; 11, 1007). BROMO-ACETOTHIENONE v. THENTL BROMO-METHYL KUTONE and BROMO-THIENYL

From phthalyl-acetic acid and Br. Resolved by

BIGTHYL RETONE. HEXA-BROMO-ACETYL-ACETONE v. 11exa-UROMO-DI-METRYL-METRYLENE DI-KUTONE.

. BROMO-ACETYL-BENZENE is Bromo-Aceto-PHENONE (q, v_*) .

BROMO-ACETYL BROMIDE CH Br.CO.Br.

(150°). S.G. № 2·317. Formation. -1. From AcBr and Br at 100° (Gul, A. 129, 54). -2. Fr. m AcBr and PBr, at 150° (Samosadsky, Z. (2) 6, 105). -3. From

AcCl (64 pts.) and Tir (120 pts.) at 100 (Hübner, A. 124, 315; Naumann, A. 129, 257; Gal, A. 132, 179) - 4. By direct combination of u-dibromo ethylene Clir.; CII, with oxygen (Demole, C. R. 86, 512). Properties.-Pungent liquid: blisters the skin.

Reactions.-1. Aqueous Na CO, forms sodium bronm-acetate and sodinm glycollate.-2. Distillation with NaOAc produces Ac,O, bromoacctic anhydride and giveollide .- 3. gives a compound whence water liberates methyl-isopropyl-carbinol, acetone, and methyl ethyl ketone (Winogradow, A. 191, 127).

Di-brome-acetyl bromide CHBr. CO.Br (194°). From the proceding and Br at 150° (C.). Formed also by the action of oxygen on fribromo-ethylene (D.). Faming liquid; converted by alcohol into di-bromo-acetic other.

Tri-bromo-acetyl bromide CBr, CO.Br. (220°-225°). From the preceding and Br at 200° (G.). Slowly converted by water into tri-

bromo acetic arid.

BROMO-ACETYL CHLORIDE CH.Br.COCL S.G. 2 191. Purport liquid, produced by the action of PCl, on bromo acetic acid (Wilde, A. 132, 1711.

BROMO-ACETYL CYANIDE CII Br.CO.CN. [79°]. Formed, together with CH CN.CO.Br by the action of AgCy on bromosac tyl broadde (Hübner, A. 131, 68). Monoclinic taldes, decomposed by water into HCy and CH Br.CO II,

BROMO-ACETYLENE Claricil. Former by the action of absoluble KOH on CH Br.CBr, (Rebant, A. 125, 81), on CH Br.CHBr. (Vexe-jeff, Z. 5, 644), or on CHBr:CHBr (Sawitsch, A. 119, 183; Fontaine, C. I.: 70, 1361; Salcocoff. Bl. [2, 45, 215).

Preprestion. Acception dibaunide is heated with NaOH and dilute alreaded in an apparatus filled with nitrogen. The gas is condensed by a

freezing mixture (8.).

Properties. Grs at ordinary temperatures; m. sol. water. In the liquid form it is decomposed by light into a tri-bromo hencene and other products. It takes fare in mir; when slowly mixed with air bromo-actic acid is formed. Ammonineal cuprous chloride gives red pp. of express acetylide. Hr forms CHille Clic

B. BROMO B ACETYL PROPIONIC ACID C.H.BiO, i.e. CH_eCO.CHEr.CH. CO.H. Transport

levulie aci l. [59] Cry tallised from CS.

Formation, A. By biomination of a solution of Seacetyl propionic acid in cene, HCl below 0 : 2. By the action of water upon (a)-ring 'renlactone dibromble (Widtf, A. 229, 266; r. Di -BROWN ON VALUE RELEASED.

Properties. White needles (from CS). Sol.

alcohol, ther, and water. Revillors. Hy the action of aqueous Na CO, it yields exyste tylpropromie and and aretacrylic acid. By heating with roots, aqueans NII, at 1105-120° diamethyle ketims (tetra methylpyrazine) C.Me, No is formed, with evolution of With aniline it gives displayly tetra-CO. methyl-di-hydro pyrazine C.Me.N.Ph, (Wolff, B. 20, 425).

Ethyl other E(A'. (210). S.G. P 1 130. From Br and ethyl β-accept propionate (Counal a. Guthzeit, B. 17, 2285), Maining ether and NaOEt convert it into methyl propel ketone tricarboxylic ether (CO Et) CH.CHAc.CH CO. Lt .

(c. 283°) S.G. 1-1 007.

Di-bromo Bacetyl-propionic acid C.H.Br O. 11152. Solidines about 91 From Bacetylpropionic acid, chloreform and Er. From bromo-8-acctyl propionic acid and bromine. Long thin needles (Wolff, A. 229, 266; Hell a. Kehrer, B. 17, 1981).

Tri-bromo-S-acctyl-propionic acid C 11 it O. From S-acetyl-propionic acid, Br and chloroform (Wolff, A. 229, 267).

BROMO-pseudo-ACETYL-PYRROL v. BROMO-PYRRYL-METHYL KRYONE

BROMO-ACETYL-UREA P. UREA.

BROMO- ACIDS v. BROMO- COMPOUNDS.

BROMO-MET-ACROLEIN(C.H.DrO),(?),[78°]. Aerolein takes up Br forming di bromo propionio aldeliyde, a liquid which polymerises, becoming a grammy mass, which may also be obtained by the action of Br on metherolem. NaOEs converte this guinny metaerelem dibromide into bromo mel aerolem (Grimaux a. Alam, Bt. [2] 36, 136). Needles (from alcohol). Itas no smell; does not reduce Felding's relate in When distilled with H.SO, (I vel.) delated with water (I vol.) it gives off extremely pangent vapours which may be condensed to a liquid, sol, water, which is probably brome acrolem. By heating with NaOl3 it is converted into C,H BiO, [140].

a BROMO-ACRYLIC ACID CHI HO, i.e. CH (CBr CO II, 170). From as or aboli bromopropionie ned and alcoholic KOH (Plulippi a. Tellens, A. 171, 333; Warmer a. Tellens, A. 171, 340; Erlenmeyer, D. 14, 1867).

Properties. Rectangular momelinic plates, v. sol, water and alcohol. Decomposed by distillation or exposure to light. Combines with HBr forming ad-di brome propionie neal.

mm. With sodium makonio ether it gives CO D. CJCH J. CH(CO Et), identical with the compound from as distagnes propriete ether and disodium malonic other (Michael, J. pr. [2]) 35, 131).

B. Bromo-acrylic acid CHB: CH.CO.H. '1160]. From broughide or from trechloro ethylideno trid rome hetate by robe tion with Zu and HCl (Waitheli, J. 193, 55). Formed also by the addition of HBr to propiolic acid (Bandenwski, R. 15, 2702; Stolz, B. 19, 5400. Plates or modlen sol, water and chloroform; expludes ore locations.

Acryl colloids. This name is applied by Wiener a. Toller to three bodies having the composition (C.F.O.).z.

a-Acryleallord. Is formed in the preparation of a bromonerylia acid from abdibrougo propionic goal, expecially who a theartion teromes violent. Insol, water, alcohol, and ether, sol. NH, Aq and not reppet, by HCl.

B-Acryl collord is formed when a bramo. neryle are best left over H SO, It is a porous mas, sol. NH, vq and reppel, by HCl.

a story to a laid is formed with reparation of Lilly, by Leating ethyl a bromo-nerylate. Anonydams; in ol. NH Aq, but becoming canasis therein.

as Dishromo-acrylic acid (1116. Chr.CO.H.

S. 5/19 at 18%

method. 1. From 25 sea bromie neid and cold barria water (Jackson a. Hill, B. 11, 1673; .:m. 3, 111; 4, 169, 273; 1 all a. Andrews, P. An. A 16, 192; 17, 153), 2, From alcohatie KOH and ang tri-bromo per pionic seid (M.chael a. Nort n. Am. 2, 18; Mauthuer s. Suida, Silz, B. 83, 273; M. 1, 104). 3, From aß tri la amo propionie med and alcoholic KOH.

Proporties, Yearly plates (from alcohol). d, other and chincoform, st. sol. benzene and CS., Bodingbaryta water form bromo acetylene, > CO, formic, malonic, and brome propiolic soids: Heated with Br in a sealed tube it forms tetrabromo-propionio acid.

Salts .- AgA': slender needles .- PbA', aq: pearly plates, sl. sol. cold water .- BaA', aq. 6.28 at 18'.—CaA', 3aq : long needles.—KA'.

BB - Di bromo - acrylic acid CBr. CH CO. H.]. S. 3 06 at 20°. From tri-bromo-succinic acid by heating with water (Fittig a. Petri, A. 195, 70). Formed also by the union of 1413r with brome-propiolic acid (Hill, B. 12, 560;

Hill a. Mabery, P. Am. A. 16, 211). Properties. Large plates; boils with partial decomposition at 243-250°. V. c. sol. alcohol, and other; m. sol. cold water. Does not combine with III:r in the cold. Does not combine with Br in the cold, but at 100° it forms tetrabrome-propionic acid (Mabery a. Robinson, Am. 5, 251).

Salts. BnA', 2aq: S.1264. CaA', 3 aq. Ethyl ether Eth'. (213). Tri-bromo-acrylic acid Clir. CBr. CO.H. [118°]. S. I 37 at 20°. From αβββ-tetra-bromopropionic acid and alcoholic KOII at 60' (Mauthner a. Suida, M. 2, 109). Formed also by treating brome-propiolic acid with brominewater (Hill, Am. 3, 178); and from di-bromoiodo-acrylic acid and Br at 100' (Mabery a. Lloyd, Am. 4, 92). Monoclinic prisms, a:b:c

·502:1: 559 (Melville, P. Am. A. 17, 151). Triclinic pyramids, a:b:c 1·128:1;1·150; α . 83° 13', β 62° 26', γ · 91° 14' (Becke, M. 2, 111). V. sol. alcohol and other. Does not combine with bromine, even at 200?. The Ba salt yields tri-bromo-ethylene when boiled with water.

Salts. BaA'₂5aq. BuA'₂3aq. S. (of BaA'₂) 80·6. CaA'₂3aq. AgA'.

BROMO ADIPIC ACID C.H.BrO. From adipic acid and Br (1 mol.) at 160 ' (Gal a. Gay-Lussao, C. R. 70, 1175). Dark-brown mass, smelling like camphor; converted by hoiling alkalis into adipomalie (or oxyadipie ?) acid (v. p. 61).

(a) di-bromo-adipio acid Call Br.Oc. From adipio acid Br (2 mols.) at 160 (G. a. G.). Powder; decomposed by water. Water at 150 Iornis adipotartario (or di-oxy adipie ?) acid (v. p. 61).

(8) dl - bromo adipio acid Call Br.O. [115°-122°]. Formed together with bromo-hydronuconic acid, by the action of Br on a colution of hydromucomic agid (Limpricht, A. 165, 265). Needles. Converted by moist Ag.O or baryta into di oxy-adipie acid.

(7) · di · bromo · adipic acid C.H.Br.O. [175° Buo]. Formal by adding Br to a warm solution of hydromuconic acid in glacial HOAc (L) Small needles. Converted by moist Ag.O into ameonic acid, and by sodium-amalgam to hydromuconic acid.

Tri-brome adipicaeld C.H.Br.O. [1770-1800]. Formed by treating a hot solution of hydromuconic acid with excess of Br (L.). Small needles. Converted by boiling baryta-water into tri-oxy-adipio acid.

Tetra-bromo-adlpio acid CallabraCa. [2115]. Formed by heating hydromuconic acid with Br and water at 100'. Crystalline. V. sl. sol. water, v. sol. alcohol. H. W.

BROMO-ÆSCULETIN v. ÆSCULETIN, p. 65.

BROWN ALDRES . DR. W. HRONG LORING MAPS BYDE

BROMO-ALIZARIN v. BROMO-DI-OXY-ANTHRA-QUINONE.

DI BROMO-DIALLYL v. Di-BROMO-HEXIN-ENE.

BROMO-ALLYL ACETATE C,H,BroAc i.e. CHBr:CH.CH.,O.Ac (?). (184°). S.G. 12 1-57. From brome-allyl bromide (\$\beta\$-epidibromhydrin) and alcoholic KOAc (Henry, E. 5, 453). Iragrant liquid; not attacked by PCl. H. W. a-BROMO-ALLYL ALCOHOL C.H BrO is.

CH CBr.CH.OH. (152). From α-bromo-allyl bromide (a epidibromhydrin) and water at 130? (Henry, B. 14, 403). Liquid; converted by aqueous KOH into propargyl alcohol.

β-Bromo-allyl alcohol CHBr:CH.CH.OH(?). (155°). S.G. 121B. From β-bromo-allyl acetate (v. sup.) by distillation with solid NaOII (Henry, B. 5, 453). Is perhaps identical with the preceding.

DI-BROMO-DI-ALLYL-AMINE C.II.Br.N i.e. (C.H.Br),NH. From s-tri-bromo prepano (tribromhydrin) and alcobolic NII, at 100 (Maxwell Simpson, P. M. [4] 16, 257). Also from bromo allyl bromide and alcoholic NII, (Reboul, A. Suppl. 1, 232). Alkaline liquid, v. sl. sol. water. Converted by alcoholic NII, at 250° into methyl-pyridine - B',H,PtCl, -B'HgCl,

BROMO.ALLYL BROMIDE v. Di-BROMO-PROPYLENE.

BROMO-ALLYLENE v. PROPARGYL BROMIDE. Brome diallylene v. HEXONYL BROWLDE. BROMO-ALLYL ETHYL OXIDE v. ETHYL

BROMO-ALLYL OXIDE. BROMO ALLYL NITRATE

CHBr:CH.CH.,O.NO, (?). (140° 150'). S.G. 13 1.5. From & bromo-allyl bromide and AgNO, (Henry, B. 5, 452).

BROMO - ALLYL OXIDE C.H.Br.O i.c. (CHBr.CH.CH.).O (?). Bromo allyl ether. (c. 2148). S.G. 27 1-7. Formed together with propargyl alcohol from B bromo-allyl alcohol and solid KOH (Henry, B. 6, 729).

BROMO ALLYL THIO-CARBIMIDE C₃H₄BrN.CS. (c. 200°). From bromo-allyl bromido and alcoholic potassium sulpho-

eyanide (Henry, B. 5, 188). BROMO-ALLYL-THIO-UREA

(CaH4Br)NH.CS.NH, [III°]. From the proceding and ammonia (II.).

TRI-BROMO-ALOIN r. ALOIN, p. 141. BROMO-AMIDO-ACETOPHENONE [5:2:1] C₆H₄Br(NH₂).CO.CH₄.

Acctyl derivative CaH.Br(NHAe).CO.CH4. [160°]. Obtained by bromination of acetyl-o-ami lo-acetophenone in acctio acid solution (Bacyer a. Bloem, B. 17, 965). Slender felted colourless needles, sol. hot, al. sol. cold, accohol; v. sl. sol. cold water. By KMnO, it is oxidised to bromo-isatin [255°]. By beiling with KOII it yields bromo-indigo.

wwm.Tri-bromo o amido-acctophenone [5:2:1] CallaBr(NII.).CO.CHBr₂. [o. 115]. Fine leited orange yellow needles. V. sol.alcohol and other, orange yellow needles. V. sol. alcohol and other, sl. sol water. Obtained by saponification of the acetyl derivative by boiling it with a mixture of alcohol and aqueous HBr.

Acetyl derivative

CallaBr(NIIAe).CO.CHBr2: [185°], yellowish granular crystals, v. sol. ohlorolorm, al soi.

slechol. Formed by the action of bromine vapour on dry seetyl-o-amido-acetophenone mixed with a little iodine. By boiling with HCl it yields a-di-ohloro-m-bromo-o-amido-acctophenone. By KMnO, it is oxidised to bromo isatin. By boiling with dilute NaOII and exposure to the air it gives bromo indigo (Bacver a. Blocm, B. 17, 966).

DI. BROMO-AMIDO. ANTHRAQUINONE

C, H, Br (NH)O . [170° nucorr.]. Prepared by reduction of dibromo nitro anthragminine with otannous chloride (Claus a. Diernfellner, B. 14, 1334). Slender red needles. Sl. sol. all solvents. Has no basic properties.

BROMO-AMIDO-BENZENE v. BROMO-ANI-LINE.

(1, 2, 4).BROMO.AMIDO.BENZENE

PHONIC ACID C.H. BrNSO, i.e.
C.H. Br(NH.)SO, H (1:2:1). Brown aniline sulphonic acid. S. 1 19 at 11° (S.); 2 61 at 20°

(A.); 131 at 45 (bab.

Formation. -1. By reduction of (1, 2, 4)-bromo nitro benzene sulphonio acid (Goslich, A. 180, 100). 2. By sulphenation of a bromo-aniline (Andrews, B. 13, 2126). 3. From bromobenzene p-sulphonic acid by nitration and reduction. 1. From very dilute aqueous amidobenzene mesulphonic acid and bromine-water (Langfurth, A. 191, 176). 5. From (1, 3, 4, 6) diromo mamido beneene sulphanie zeid, faming HCl, and red P at 150 (Limpric st, B. 10, 1542). 6. By heating the same acid with water at 250

(L.).
Properties. - Anhydrous needles (from cono. squeous solution) or four- an I six-sided columns containing an (from dilute solution). Sl. sol. alcohol. Reduced by III and P, or by water at 120°, to amido bemo ne m sulphonic acid.

Salt s. .. K V13 aq (La.) ... KA' aq (Spiegelberg, A. 197, 257). - BaA', 20. -BaA', 301. S. (of BaA') 6·22 at 17'. -PbA', . CaA', 220. -AgA' 13nd. (1, 1, 3). Brome amide beazene sulphonic s zid

O.H. Er (NIL)SO.H (1:1:3]. S. 47 at 15".

From itim. -1. By heating p-brono anilino ethyl-sulphate (Nölting, B. 8, 1095) ... 2. By adding bromine to a cold represus solution of barium aniline o sulphonate (Limpricht, .t. 181, 196) .- 3. By nitration and reduction of brenosbenzeno m-sulphonic acid (Thomas, A. 186, 126). 4. From acetyl-p-bromo-aniline and funning H₂SO, (Borns, A. 187, 368).

Properties .- Slender silky needles (containing aq) or large efflorescent prisms (with 2 aq). Sl. sol. oold water, v. sl. sol. sleohol. Converted by Br into tri-bromo-amiline. HI and P form : aniline a sulphonic acid.

Salts. NH,A'. - KA'. - BaA', aq. - ChA', aq.

PbA', 2aq.

(1,4,2)-Bromo-amido-Leazene snlphonic acid O. H. Br(NH.)SO, H [1 4.2'. S. 1 11 at 18". From bromo benzene o sulphonic and by nitration and reduction (Sahlmann, A. 181, 203, 186, \$18). Needles (from conc. aqueous solutions) or rhombohedra (1) containing 2aq (from dilute aggeous solution); v. sol. hot water, insol. alcohol and ether. Reduced by HI and P to aniline m-sulphonic acid. Salts .- BaA', 2aq: needles, v. o. sol. water. -PbA'2--AgA'.

Bromo amido-benzene salphonio acid S. .74 at 8°. C.H.Br(NH,)(SO,H) [1:x:2]. Formed in smaller quantity in the preparation VOL L

of the preceding (B.). Priams; sl. sol. sold water.—BaA', raq: lamins, v. sol. water, Bromo-amido-bensens sulphonic acid C.H.Br(NH.)(SO.H) [1:3:x]. From acetyl mobromo aniline and fuming H.SO. (Borns.B. 8, 1072). Needles. -- BaA', 2nq.

(1.2.3.5)-Bromo-amido-benzene disulphonie acid C.H.Br(NH.)(SO,H), [1:2:3:5]. Formed by bromination of (1,2,4) amido benzene disulphonic acid (Zander, A. 198, 1). Needles (containing aq); v. snl. water, el. sol. al-abol.

Sults.-The acid sults are less soluble than the normal ones. -(NII_s), A" 2aq. BaA" 3aq. --Bull A", 5aq. - PbH A" 5aq - KA" 2nq.

Bromo-amido-benzene disulphonic acid C_all.lir(NH₁)(8O₁H)₂ [1:4:3:5] or [4:1:3:5] or [2:1:3:5]. Formed by adding browing to an none our solution of (1 or 4, 3, 5) amide henzens disulphonic acid (Heinzelmann, A. 188, 179). Prisms (containing 2 jaq); v. c. sol. water .-

BaA"Saq. 1'bA"3aq. Bromo-di-amido-benzene sulphonio acid C.H Br(NH) SO [H[1:2:6:4], Brome in phenylene diamine rulphonic acid. S. 51 at 17°. Got by reducing C.Br. (NO) SO, H with SnCl. (Bassmann, A. 191, 214). Long white needles (containing sq), turns yellow in sir. pure it is sl. col. water, when impure it is very soluble. Insol. alcohol. Paper moistened with its solution turns red in air. The aqueous solutions of its salts turn blue or red when evaporated. Converted by dinzo-reaction into p broma benzene sulphome neid. « BuA , nq.

(1,3,2,5) Di-bromo-amido-benzene enlphonis acid C.11,Br (N11.)(SO,11) [1:8:2:5]. Formed by adding bromine to an aqueous solution of aniline p sulphonic acid (Schmitt, A. 120, 138; Lenz, B. 8, 1066; A. 181, 24). Formed also by brominating (1,2,4,) amido henzene disulphonia acid (Zander, A. 198, 16). Needles or prisms (containing 2aq); v. sol, water and hot alcohol; ppd, by one H SO, from its aqueous solution. Br forms tri-brome milline, BaA', 2aq. -BuA', 3 | aq. S. (of BuA',) 16 at 11°. PbA', 2aq. -A: A'.

Reactions. FCl, forms a product (probably C.H.Br (NH.POCL)SO Co from which alcohol produces C.H.Br. (SO,Cl)NH.PO(OEt). [170°] (Laur. J. pr. 128, 256). V. also of Methyle-AMILE BLAZENE SULPHONIC ACED.

(1.3.4.6) Di brome amido-beazene sulphonie acid C.H.Br (NH.)SO.H [1:3:4:6]. S. 252 at 10; 249 at 7; 191 at 4° (Berndsen, A. 177, 81; Beckurts, A. 181, 213; Beinke, A. 186, 286; Knuth, A. 186, 301; Laugfurth, A. 191, 180; Bassmann, A. 191, 229, 238; Spiegelborg, A. 197, 206).

Formation, -1. From amido l'enzene mesulphonic acid and bromine. -- 2. From (1, 2, 4)bromo amida benzene sulphonic acid and Br --3. From the corresponding Call Br (NO2)SO2H .-1. From (1,3,5,2,6)-tribromo nitro-benzene sulphonic acid, Sn. and HCl. 5. From tribromoamido benzene sulphonio acid by treating with water at 150", or by treating with Sn and HCl.

Properties .- Needles (from hot water) : sl. sol. water, invol. alcohol.

Reactions .- 1. Water at 250° forms o-bromoaniline sulphonic acid and aniline mesulphonia acid. -2. Diazo reaction gives (1,3,4)-di-bromo-benzene sulphonic acid. - 8. Warmed with strong alcohol and KNO, a yellow crystalline body, poseibly C.H.Br. (SO.H).N.H.C.H.Br. SO.K. is formed. Heatod in sealed tubes with alcohol, it splits up into dibromo-benzene eulphonis scid and amido-dibromo-bonzene sulphonio acid

(Bässmann).

Salts. - BaA', 6aq. S. (of BaA',) 2.99 at 7°; 8·12 at 9°; 3·9 at 23°. — KA'aq. — NH,A'. CaA', 2aq. - CaA', 5aq. - PbA', S. 2.9 at 22,5°. (1,2,4,5)-Dl-bromo-amido-benzeno oulphonic

acid CallyBr2(NII2)(SO4II) [1:2:4:5]. S. 109 at 10°; 153 at 24°. From (1,2,4)-di-broam benzeno sulphonic acid by nitration and reduction (Spiegelberg, A. 197, 279). Trimetrie tables; v. sl. sel. alcohol. - NIIA' aq. - KA' 2aq. -BaA', aq: S. (ol BaA',) '67 at 11'. - 1'bA', aq: S. (of PbA',) '11 at 11°. - CaA', 3aq. - CuA', 4nq.-

AgA'. S. 053 at 11°.

(1,3,4,5)-Di-bromo-amido-benzenc sulphonic aold C,H,Br,(NH)(SO,H) (1:3:4:5). S. 3:13 at 10-5°. From umido-benzene o-sulphonic acid by bromination (Limpricht, A. 181, 198; L. 8, 1429), or from (1,3,5)-di-bromo benzene sulphonic need by nitration and reduction (Leuz, A. 181, 36). Trimetric tables (anhydrous) or 4-sided prisms (with aq). Converted by Br into tri-bromo-anilina.

Salts. KA'nq .- NaA' aq. S. (of NaA') 3.7 at 120.—BaA', 1 a. S. (of BaA',) 20 at 110.

Di-bromo-amido-bonzens sulphonic acid C_sH_sBr_s(NH_s)(SO_sH) (1:4:27:6]. S. ·62 at 10·5°. From (1.4.2) di-branco-benzeno sulphonie acid by aitration and reduction (Borns, A. 187, 362). Needles or prisms. - KA', ... l'aA', aq.

Di-bromo-amido-bonzono disulphonio acid CHBr_(N11_)(SO_H), [1:4:37:2:67]. From p-dibromo-benzene disulphonic acid by nitration and reduction (Borns, A. 487, 367). Crystals; v. sol. water. K.A", BaA".

Di-bromo-amido-benzeno di-sulphonic acid C,11Br,(NH.)(SO,11), [1: 4or6:2:3:5', From (1,2,4) audido benzene disulphonia acid and Br (Heinzelmann, A. 188, 182). Prisues (contain-

Di bromo di amido-bonzons sulphonic acid C.HBr.(NH.).SO.H [1:3:2:6:1]. One of the produots of the reduction of C.Br. (NIL) SO.11 (Bässmann, A. 191, 214, 218). Tublets (containing aq), v. sl. sol. water,

Tri-bromo-amido bsuzeno sulphonic acid C.HBr. (NH.)SO., H, [1:3:5:4:6]. S. 13:7 at 14°. 16:6 at 7° (B.).

Formation.-1. From amido-benzene m sulphonic acid and Brt (Berndsen, A. 177, 86). 2. From the corresponding nitro-acid, by Su and HCl, somo di bromo amido benzene sulphonic acid being also formed (Rencke, A. 186, 282; Knulb, A. 186, 298; Januarath, A. 191, 198).— 3. From (1,2,4)-bromo amido benzene sulphonio acid by bromination (Spiegelberg, J. 197, 275). Properties. - Thin needles (containing aq).

Sol. cold water and alcohol. Heated with water at 145° it becomes Call Br. (N11.)SO. H.

Salts. -BaA', 9aq. S. (dry). 43 at 7° (L.), 34 at 3° (Bässmann, A. 191, 221). -KA'nq. 935

at 4° (B.).—PbA'.9aq. S. (of PbA'.) 73 at 14°. Tri-brome-amide-benzene sulphonie acid C₄HBr₂(NH.)(SO₄H) [1:2:3:4:5]. From (1, 2, 3, 5).

and reduction (Lenz, A. 181, 48). Tuffs of alen der needles (containing aq), v. sol. water and 98 p.o. alcohol. -BaA', 1 aq.

Tri-bromo-amido-bonzene sulphonie acid C_HBr_(NH_c)(SO_tH) [1:2:5:6:4]. From the cor-responding nitro-acid (Spiegelberg, A. 197, 288). Long prisms (containing 13aq) or slender needles (with aq). V. sol, water and alcohol,—KA'aq. S. (of KA') 200 at 1°.—NH₄A'.—CaA', 33aq.—BaA'... S. '006 at 1°.—PbA', 2aq. S. 40 at 35°.—AgA', 3aq. S. (of AgA') 46 at 10°.

The home of a midd, heaven a sulphonic and

Tri-bromo di amido-benzene sulphonio soid C.Br. (NH.) SO H [1:3:5:2:4:6]. A product of reduction of C.Br.(NO.) SO.H (Büssmann, A. 191, 249). —BaA , I , aq.

Tetra bromo amido benzene sulphonic acid C.Br. (N11.)SO.H. [1:2:3:5:4:6]. S. 2·25 at 11° (Beckurts, A. 181, 223). Got by roducing C.Br. (NO)SO, II with Su and 11Cl, not allowing the temperature to rise to 100°, or Br, will ba displaced by H. Needles (containing 2ng), V. sol, alcohol and water,

Salts .- (Langfurth, A. 191, 201) BaA'2aq. S. (of BaA',) '4 at 13', - CaA', 7aq, KA' 13aq. S. (of KA') 171 at 15'.

Tetra-bromo-amido-benzene sulphonic acid C, Br, (NH,)(SO,H) [1:2:3:4:5:6]. S. 03 at 11°. From the nitro acid (Spiegelberg, A. 197, 302). Needles (continuing 2aq). V. sol. alcohol. KA'uq. S. (of KA') 105 at 11. CaA'. 2aq. S. (of CaA',) .107 at 11.50, BaA', aq. S. (of BaA',) 0155 at 11.5

(a) · BROMO · o · AMIDO · BENZOIC ACID C_bII [Br(NII]CO]H [1:2:3]. Bromo-anthranilic acid. [170]. From the corresponding nitrocompound by Sn and HCl (Hübner, A. 222, 104; cf. A. 143, 211; 149, 134). Needles; m. sol. water. Sodium amalgam reduces it to o-amidobenzoicacid [114"]. Nitrous acid lorus the diazoderivative C, H, BrCO, H.N, NH.C, H, BrCO, H.-Salts .- AgA'. BaA' nq .- CuA'. Acetyl derivative

Callabr(NHAc)(CO H) [17:2:3]? [215°]. Obtained brominating acetyl-o-amido-benzoic acid (Jackson, B. 11, 879).

(B)-Bromo-o-amido-bonzoio acid. Bromo-anthranilic acid. C. H. Br(NH.)CO.11 [1:4:3]. [2089].

Formation. -1. By reducing (1, 4, 3) bromonitro-benzoic acid (Hubner, Oldy a. Philipp, A. 113, 212; Mecker, Hubner a. Petermann, A. 149, 133). 2. By boiling bromo-isatoic acid with conc. HCl (Dorsch, J. pr. [2] 33, 35).

Properties. -V. sol. acetono, sol. alcohol. ether, chloroform, benzeae, and glucial acetic neid, sl. sol. boiling water. Sodium-amalgam reduces it to o-unido-benzoic acid [1410].

Salts. -BaA' 4aq: needles, v. sol. water. Amide. C₈H₈Br(NH) CO.NH. [177°]. From brome isatole acid and NI! Aq. Flat needles. V. sol. nicohol, acetone and glacial acetic acid, m. sol, water and benzene. Insol. other.

(1,2,4)-Bromo-m-amido-beazoic acld C. II,Br(NH.)CO.H [1:2:4]. [225°]. duoing the nitro acid by Sn and 11Cl. Small colourless needles (from water), becomes reddish in air (Hübner, A. 222, 179; Burghard, B. 8, 558; Raveill, B. 10, 1707). Reduced by sodiumamalgan to m-amido-benzoic acid. - HA'HCL-CuA'2.--PbA'2.
(1,8,5)-Bromo-m-amido-benzoio acid

tri-bronto-benzens sulphonio acid by nitration C.H.Br(NII.)CO.H [1:3:5]. [215°]. From the

corresponding nitro- acid by Sn and HCl (Hesemann s. Köhler, 4.222, 169). Needlas (from alcohol). Turns red in light. Salts.—HA'HCl.— (HA')₂H₂SO₄.—BaA'₂4aq.—CaA'₂5jaq. (1,4,2)-Bromo-m-amido-benzolo acid

C.H.Br(NH.)CO.H [1:4:2]. [180°]. From (1:4.2)-bromo-nitro-benzoic scid, Sn, and glacial HOAc (Burghard, B. 8, 560). Flat needles (Smith, B. 10, 1706).

(1,2,4,5)-Di-bromo-o-amido-beazoic acid C₄H₂Br₂(NH₂).CO₂H [1: 2or6:5:4]. Di-bro Di-bromo-

anthranilic acid. [226°-228°].

Formation.—1. I rom di-bromo-nitro-benzole acld (Smith, B. 10, 1706). - 2. From o-nitro-tolueno and bromine at 170° (Wachenderli, A. 185, 281; Grieff, B. 13, 288).-3. From isutoio

acid (q. v.) and bromine (Dorsch, J. pr. [2] 33, 37).
Properties.—Clumps of needles (from alcohol). Long needles (when sublimed). Sol. alcohol, acotone and glacial acetic acid, al. sol. chloro-

dom, benzene, ether, and water.

Amide Call Bra(NH.)CO.NH. Poarly tablets (from alcohol-acetone). Formed from di-bromo isatoic acid and NH3, Aq at 100 '.

Di-bromo-o-antido-benzoic acid CaH.Br. (N1L)CO.H [41:2or6:5:4]? Disbromoanthranilic acid. 225°). S. 1; S. (alcohol) 2. From the mitro- acid by reduction (Hillmer, A. 222, 189). Colourless needles. Reduced by sodium amaleum to o amido-henzoic acid. BaA', 4nq. - CuA', 4 jaq. - SrA', 2nq. - CuA', This acid is probably identical with the preceding.

(1.3,4,5).Di-bromo-o-amido-beazoic acid C.H.Br.(NII.)CO.H. [1:3:4:5]. Di bromo anthranilic acid. [225]. S. 1; S. (HOAc) 3. By reduction of the nitro- acid by Sn and HCl (Hesemann a. Köhler, A. 222, 175). Reduced by sodium amalgam to o amido-benzoic ucid. Needles (from alcohol).-BuA', 4aq. -CaA', 4aq. -CuA'2

(1,3,5,4)-Di-bromo-o-amido-benzoic acid C. H. Br. (NH.) (CO.H). Di-bromo anthranilic acid. (1966). From benzoic acid by bromination, nitration, and reduction (Augerstein, A. 158, 16). Needles (from dilute alcohol). Reduced by sodium-amalgam to o amido-lenzoic acid.

Di-bromo-p-amido-benzoic acid

C, H, Br, (N11,)CO, H [1:3:2:5]. Obtained by adding bromine-water to an acidified solution of p-amido-benzoic acid (Beilstein a. Geitner, Z. [2] 1, 505; A. 139, 1). Needles (from alcohol). N11,A 2nq. - NaA' 5aq. - CaA' 2 6nq. - BaA' 2 4nq. Tri-bromo o amido benzoie acid

C. HBr. (NII.)CO. H [1:2:x:4:5]... Tri-bromo. anthranilic acid. [c.119°]. From isatoic acid and bromino (Dorsch, J. pr. [2] 33, 37).

Properties .- Slender it alles. May be sub-Very soluble in glacial acetic acid, alcohol and acctone, sol. ether and chloroform, less soluble in benzene, sl. sol. hot water.

Tri bromo m amido benzoio acid C. HBr. (NH2)CO. H [1:3.5:4:6]. [170°]. From m-amido-benzoic acid and bromine water (Beilstein a. Goitner, Z. [2] 1, 505; A. 139, 6; Vollbrecht, B. 10, 1708). Needles; m. sol. hot water. On dry distillation it gives tri-bronto-aniline.—NaA'4aq.—BaA', 6aq.

Tri-bromo-di-amido-benzoic acid C.Br.(NH2)2(CO2H) [1:3:5:2:4:6]. From s-diamido-benzoic acid and bromine-water (Griess,

A. 154, 832). Long needles (from dilute alco-hol). Sl. sel. cold water.—AgA'. Tetra-brome-o-amido-benzoio acid

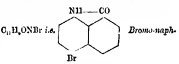
C.Br.(NH.,)CO.H [1:2:3:4:5:6]. [115°]. From isatolo acid and excess of Br (in glacial acetic acid) (Dorsch, J. pr. [2] 33, 38). White needles. At 100° it sublimes in long slender needles

BROMO - AMIDO - HYDROCARBOSTYRIL v. p. 161.

BROMO-AMIDO HYDROCINNAMIC ACID

v. Bhomo-amido-8-phenyl-propionic acid. BROMO AMIDO NAPHTHALENE t. BROMO-NAPHTHYLAMINE.

BROMO-AMIDO-NAPHTHOIC ANHYDRIDE



thostyril. [257°]. By reduction of bromo-nitro-(a)-maphthoic acid with FeSO, and aqueous NH. and ppn. with acctic acid the amide-acid is obtained, and by boiling with alcohol it is converted into the anhydride, which crystallises out on cooling in brown needles (Ekstrand, B. 19.

Di-bromo-amido-unphthoio anhydrlde

$$C_{10}H_4Br_2$$
 $\stackrel{CO}{\underset{NH}{\downarrow}}$. Di-bromo-naphthostyril. [270°],

Prepared by heating amido naphthoic anhydride

 $C_{10}H_{\bullet}$ is suspended in water with a large

excess of bromine. Yellow needles (from alcohol). M. sol. hot acetic seid.

Acetyl derivative C, H,ONBr Ac [1850]; small yellow needles (Ekstrand, B. 19, 1136). BROMC-AMIDO-(0)-NAPHTHOQUINONE

 $C_oH_4 < \frac{CO.C_1NH_2}{CO.CBr}$, [205°]. Formed by boiling

bromo amido-(a) najdithoquinone imide $CO = C(SH_2)$ $C_sH_{\underset{\mathbf{S}_1}{\longleftarrow}}\mathbf{C}_{(\overset{\mathbf{N}}{\mathbf{N}}\overset{\mathbf{H}}{\mathbf{H}}),\overset{\overset{\mathbf{G}}{\mathbf{C}}\overset{\mathbf{F}_T}{\mathbf{F}_T}}$ with dilute acids. Orange

silky needles. Sublimable. By boiling with dilate alkalis it is converted into bromo-oxy-(a)naphthoquinene.

Accty: Verivative: [137°]; sulphur-yellow needles (Zincke a. Gerland, Β. 20, 1514). BROMO - AMIDO - (σ) - NAPHTHOQUINONE -

CO C(NII₂) [200]. Obtained C(NH).CBr

by adding bromine (5 c.c.) to di-amido-(a) naphthol or its stanno-chloride (10 g.) suspended in Sectic acid. Orange-yellov needles (from alcohol). V. sol, hot atcohol and hot benzene. By SnCl, it is reduced to brome di amide-(a)-naphthol. By boiling with dilute acids it is converted into bromo amido (a) naphthoquinone CO.C(NII2)

C,H,< . By boiling with dilute NaOH

it is converted into brome oxy-naphthoquinons-

imide $C_4H < \frac{CO - C(OH)}{C(NH).CBr}$. The latter body

AMPROVED THE ANALOG THE PROPERTY OF THE PROPER

when treated with conc. HCl or alcoholic NaOH is converted into bromo-oxy-naphthoquinone C.H. (CO.C.(OH) which is also formed by boil-

ing the bromo amido naphthoquinone with diluta alkalis (Zincke a. Gerland, B. 20, 1510).

BROMO-o-AMIDO-PHENOL

C,H,Br(NII.)(OH) [1:3:1]. [128°]. Formed by reducing bromo-nitro-phenol with Sn and HCl (F. Schütt, J. pr. (2) 32, 61). Thin plates (from CS4). Needles (from alcohol). Sol. ether, benzene, hot water, and hot chloroform. turns the sequenus solution cherry-red.

Salts .- B'HCl*. Very soluble plates. B'HBr. Crystallises very casily. - B'.H.SO. Acelyl derivative C.H.(OH)(NHAc)Br.

[178]. Plates or needles (from water).

Methyl ether Call Br(NII.) (OMe). [982]. From the nitro- compound, Sa and HCl (Staedel, 4. 217, 59). Plates (from benzenc). benzene, other or hot alcohol, insel. water. Salts. - B'HCl. - B', ll, SO, - B', ll, C,O, Ethyl other [57]. Broad needles (from

alcohol). V. sol, benzene, alcohol or ether. Salts. -B'IICl.--B',H,SO, -B',H,C,O, Bromo-m-amido-phenol C,H,Br(NH,)OH

[x:1:3]. Ethylether C.H.Br(NIL)OEt. Liquid; v. sol. alcohol and other, v. sl. sol. water. hydrochloride, sulphate, and oxalate crystalliso in white plates C. H Br(OEt)NH, HCl, SuCl, (Lindner, B. 18, 612).

Bromo p amido phenol

C, II, Br(NIL)(OII) (1:3:6]. [158°]. Prepared by reducing brome nitro-phenol, or its benzyl derivative, by Sn and HCl (O. Hölz, J. pr. '2) 32, 65). Short needles (from dilute alcohol). Sol. ether, benzenc, and chloroform, st. sol. cold water.

Salts. B'HCl: silvery plates. -(B',HCh SuCl.,-B'Hbr.

Accial derivative Call (OH) NHA0 Br. [157']. Thick nocdles (from hot water). Soluble in alkalia.

Methyl other.

From the nitro-compound, Su and 11Cl (Staedel, A. 217, 68). O.l. Tusol, water, v. sol. benzene, alcohol, or ether. B'HCl. -B',H2SO, -B',H2C2O4. Ethyl ether

From the nitro compound, Sn and HCl. Oil.

B'HCl.—B',H,SO, ... B' H,C,O,

DI. bromo-o-amido-phonol C.H.Br (NH.)(OH) [1:3:5:6]. [927]. Formed by reducing di bromo o nitrophenol (Hölz, J. pr. 2) 32, 69). Slender yellow needles (from dilute Sparingly soluble in water, v. ool. alcohol). alcohol, ether, benzene, and chloroform.

Salts. - B'HCl; plates. - (B'HCl) SuCl2: needles. - B'HUr: needles.

destyl derivative C.H.(OII)(NIIAo)Br., 16°]. Yellowish teedles (from hot water). [186°]. Yo Sol. alkalis.

Methyl ether O.H.Br. (NH.) (OMe). From the nitro-compound by Sn and HCl (Staedel. A. 217, 63). Oil. Sol. alcohol, ether, or dilute acids, insol. cold water, sl. sol. hot water. RHCl H. R. H. SO. H. 1273 — P. H. C. O.

B'IICl.—B',H SO, [177°].—B',II,C,O, Ethyl ciher C,II,Br,(NII,)(OEt). [92°]. Quadratic crystals (from alcohol). V. sol. alco-M.bromo.o.amido.phenol.

Ethyl ether C. [1]. Br. (NII.) OEt [1:3:4:5]?

[58°]. Amido-phenetol, C.H. (NH.) OEt (10g.), boiled with glacial acetic acid (100g.) and bromin (117g.). The product is poured into water an the oil that separates is distilled with steam an recrystallised from alcohol. Glittering prisms grouped in tufts. Resembles di-bromo-o-tolui dine in being but feebly basic (Möhlau a. Ochmi chen, J. pr. 132, 479).

Di bromo-m-amido-phenol.

Ethyl ether C4H2Br2(NH2)OEt. From the nitro-compound. Oil. The hydrochloride, sulphate, and oxalate crystallise in needles.— C, II., Br. (OEt)NH., HCl, SnCl.: glistening plates (Lindner, B. 18, 613).

Di.hromo-p-amido-phenol.

C_oH.Br. (NH.)OH [1:3:5:2]. [180°] (M. s. B.); [190°] (L. s. G.). Formed by reducing the nitro-compound (Möhlau, B. 16, 2845; Hölz, J. pr. [2] 32, 67; Möhlau a. Böhmer, J. pr. [2] 21, 470; Lellmann a. Grothmann, B. 17, 2731). Formed also by reducing di-brominated p-diazophenol (q. v.) with tin and HCl (Böhmer, J. pr. 132, 469):

 $C_e H_2 Br_2 < \frac{O}{N_2} + 2H_2 = C_e H_2 Br_2 (OH) \cdot NH_2 + NH_4$

Properties .- Ppd. by NaIICO, from solution of its hydrochlorido as microscopically small needles grouped in tufts. Turns blue in air. SI. sol. ether, v. sol. alcohol, m. sol. het water.

- B'HCl : glittering plates .-Salts. (B'HCI) SuCl. - B'HBr.

Reactions. - When NoO, is passed into an alcoholic solution of its hydrochloride, yallow crystals of diazo-dibromo-phenol (q. v.) are obtained, but this body is isomeric with that by tha reduction of which the dibromo-amido-phenol was prepared. An 'intra-molecular changa must therefore have occurred somewhere.

Acetyl derivative C.H.Br. (NHAc)(OH). [174]. Glittering plates (from dilute alcohol). Mothyl ether [127°], Sn, and HCl. White porcelain-like mass. Extremely sol. ether, benzene, or alcohol (Staedel, A. 217, 70; Staedel a. Damm, B. 11, 1749). Salts.—B'IICI.—B'_cH₂SO₄.—B'_cH₂C₂O₄ [195°]. Ethyl ether [67°]. Needles (from alcohol). V. sol. alcohol, ether, or benzene. Salts .-

B'HCl.—15',11.SO,.—B',2H,C,O,.
Tri-hromo-o-amido-phenol.

Ethyl ether C. HBr. (N11,)OEt [1:3:4:5:6]? [77°]. I rom amido-phenetol (5 g.), glacial acotio acid (50 g.) and bromino (17.5 g.). The product is poured into water, and the pp. orystallised from alcohol. Long silky needles. Nearly insol. boiling cono, HCl. It is totally decomposed whon heated to a temperature a little above its melting-point.
Tri-bromo-m-amitio-phenol

[115°]. Formed O.HBr.(NH.).OH [1:3:5:2:6]. by reduction of tri-bromo-m-nitro-phenol (Daccomo, B. 18, 1168). Colourless silky needles.

Sol. alcohol, other, benzene, and hot, el. sol. cold, water. Fe,Cl, gives a green colouration.

Ethyl ether C,HBr,(NH,)OEt. Solid, v. sol. alcohol and ether, sl. sol. water.—B'RCI: white needles, sol. alcohol.—B'H. SOl.; white needles, sol. alcohol. - B'HCISnCl; white

needles (Lindner, B. 18, 614). BROMO-AMIDO-DIPHENYI

Acetyl derivative O. H. BrNHAc. [247]. From Br and p. amido. diphanel

HOAc (Hübner, A. 209, 848). Needles (from alcohol).

Di-brome-di-amide-diphenyl C., H.Br. (NH.).

P. From di-brome-di-nitro-diphenyl by Sn and HCl (Fittig, A. 132, 207). Insol. water.

Di.bromo-di.amido-diphenyl

(2:4:1] C.H.Br(NH.).C.II.Br(NH.) [1:2:4]. Dibrono-benzidine. [152²]. Obtained from [3:1] C.H.Br.NH.NII.C.II.Br [1:3] and IICl (Gabriel, B. 9, 1407). Trimetric crystals, m. sol. cold alcohol.—B"2HCl.—B"H,PtCl.,
Di-bromo di amido diphenyl C,,11,Br,(NH,),

[1:5:2].. (108° corr.). Small plates. Formed by the action of an alcoholio solution of SnCl, and H₂SO₄ on p-bromo-benzene-p-azo-bromo-benz-ene. Treated with nitrous acid in alcoholic solution it gives an azimido body C12H,N2Br2 which forms glistening violet needles, [206], al. sol. alcohol (Schultz, B. 17, 465).

Tetra-hromo di anido diphenyl C, H, Br, N, Tetra-bromo-benzidine. [286]. Prepared by bromination of benzidino (Claus a. Risler, B. 14, 86). Slender colourless needles. Sol. alco-

hol, ether, CS., C.II.; insol. water and acids. BROMO-AMIDO-PHENYL-ACETIC ACID CaH, (Br)(NH,)(CH, CO, H)[1:2:5], [136], Formed by saponification of the acetyl derivative of its nitrile, or by bromination and saponification of the acetyl derivative of p-amide phenyl acetic acid (Gabriel, B. 15, 840). Colourless plates. Sol. alcohol and ether, insol. CS₂.

Acetyl derivative [165°], Nitrile C.H.(Br)(NH,)(CH,CN). Acetyl derivative C, II, Br(NIIAc)(CIL, CN) [129°]. colourless needles. Sol. alcohol, sl. sol. cold water. Formed by bromination of the acetyl derivative of p-amide phenyl-acctenitrile (Gabricl, B. 15, 810).

Bromo amido-phenyl-acetio acid C.H.Br(NH,)CH, CO.H [1:20r6:4]. From the nitro- compound [114°] by Sn and ! HCl (P.P. Bedson, C. J. 37, 98). Silky needles. (from water). Sol. alcohol and Click. Sl. sol. ethor .- B'11Cl aq: turns red in air.

(a) Bromo amido phenyl-acetio acid C.H.Br(NH.)CH. CO.H. [167°]. From the nitro- acid [169°] by Sn and HCl (Bedson). White needles (from water), reddens in air. Sol. alcohol and chloroform, sl. sol. other.

Salt .- B'HClaq: white needles (from water). (8) Bromo amido-phenyl-acetic acid

C.H.Br(NII.)CH2.CO.H. [186°]. From the nitro-soid [162°] by Sn and HCl (Bedson). The hydrochloride is less soluble in water than those of the two preceding bodies.

(1, 3, 2, 5). Bromo-di-amido-pheayl-acetic acid C₄H₂(Br)(NH₂)₂.CH₂.CO₂H₂[1:3:2:5]. [0. 200°]. Long colourless need s. Formed by reduction (1, 3, 2, 5) bromo nitro amido pheayl acetie acid (Gabriel, B. 15, 1995).

BROMO-AMIDO-PHENYL ETHANE

C,H,BrN i.e. C,H,Br.CH,CH,NII, Dromophenyl ethyl amine (253°). From phenyl pro-plonamide, KOHAq, and bromine (Hofmann, B. 2740). Peerly plates (from water).— B'IICl. (1,2,5) BROMO-AMIDO. β - PHENYL - PRO-

PIONIO ACID C.H. (Br)(NH.).C.H. CO.H [1:2:5]. Bro no amido hydrocinnamic acid [105°]. Formed by bromination of the acetyl derivative of pamido-phenyl-propionio acid, and deep violet.

saponification of the product (Gabriel, B. 15, 2292). Long glistening crystals. Sol. most ordinary solvents and in acids and alkalis.

Acetyl derivative: [160°]. Long colour-less needles, soluble in alcohol, ether, and benzene.

Bromo-m-amido-phenyl-propionio acid

C.H. (Br)(NII.).C.II. (CO.II [2:1:5]. [117°-119°]. Long prisms. Sol. water, alcohol, ether, and C.H. Prepared by reduction of p-bromo-mnitro-hydrocinnamic acid with tin and HCl .-A'HCl: glistening soluble needles (Gabriel a. Zimmermann, B. 13, 1681).

(es.) D1-BROMO O AMIDO - PHENYL-VALE.

RIC ACID

C.H.Br.(NH.).CH..CH..CH..CH..CO.H. with nql. Long colourless needles (containing aq). Formed by reduction of an alcoholic solution of di-bromo-amido-phenyl-di-bromo-valerio acid C,H,Br,(NIL).CllBr.CllBr.Cll,.Cll,.Cll,.CO,H with zine and HCl. V. sol. ordinary solvents, insol, cold water. It could not be converted into an inner anhydride, even by dehydrating agents.

Ethyl other CulligBr (SII)CO Et; thick oil; its hydrochloride forms white needles (136°). Acetyl derivative Ciellie Br. (NHAc)CO.II [206°]; aggregates of thin white needles; v. sol. alcohol, ether, &c.

Acetyl-ethyl-ether

Ciallia Bra (NIIAc) CO Let [139°]; colourless orystals , v. sol. alcohol, ether, and acetic acid, more sparingly in benzene, insol, water and ligroin; in small quantities it can be distilled undecomposed (Diehl a. Einhorn, B. 20, 380).

δγ·cso-Tetra-hromo-o-amido-phonyl valerio acid C, H.Br. (NH). CHBr.CHBr.CH, .CH, .CO, H. [167°]. Formed by bromination of o. amido. styryl-propionic acid discolved in chloraform. Small microscopic needles, V. sol. alcohol, ether, and acetic acid, insol, water and CS.

(Dich) a. Einhorn, B. 20, 379). BROLIO-AMIDO-QUINOLINE

C₀H₂N(Br)(NH₂). [164°]. Large monoclinis prisms, or long colourless needles (containing aq). Sol. alcohol and ether,

Salts,-B'HNO,: glistening yellow needles. - BHCl: soluble red prisms. B'H Cl PiCl: microscopie orango needles.

Acetul derivative C.H.N(Br)(NHAe). [105°]. Thin colourless plates (La Coste, B. 15, 1920i

D1-BROMO-AMIDO-RESORCIN
C₄HBr₂(NH₂)(OH)₂ [1:2:4]. Distribyl ether
C₄HBr₂(NH₂)(OE4)₂ [1:2²); plistening needles or plates (Will a. Pukall, B. 20, 1126).

BROMO-AMIDO-STYRELIE

C.11, Br(NII,).C.11.

Acetyl derivative [183]; felted noedles, col. alcohol, other, and actic heid; formed by bromination of p-amide cinnamic acid dis-solved in AcOH (Gabriel a. 1) rzberg, B. 16, 2043).

BROMO-AMIDO-SUCCINIC ACID

C.H.Br(NII.)(CO.H). [140]). From di-bremo-succinic acid and alcoholic NII. (Claus, B. 15, From di-bremo-1850). Radiating needles; v. sol, water and alcohol.—AgA'.

BROMO-AMIDO-THYMOL

C.HBr.McPr(NH,)(OH). Prepared by adding Na,CO, to dilute solution of its hydrobromide (v. Long colourless prisms, rapidly turning

moderately strong HBr to thymoguinone-chloroimide (q. v.) a yellow flocculent pp. is formed. Ether extracts brome-thymoquinenes from this, leaving the above salt, which is soluble in water and alcohol, but is thrown down as needles when HBr is added to its concentrated aqueous solution (Andresen, J. pr. 131, 182).

... v D BBL.

By adding

Bromo-amido-thymol [o. 90°]. From bromonitro-thymol, zinc-dust, and 11Cl (Mazzara a. Discalzo, G. 16, 196). Scales. Converted by nitrous acid gas into bromo-thymoquinone [48°].

BROMO-ANTDO-TOLUENE v. BROMO-TOLU-IDINE.

Bromo-di amido-tolucae v. Bromo-Tolylene DIAMUNE.

BROMO - AMIDO - TOLUENE SULPHONIC ACID C. II. (CII.) (NII.) 15 SO. II [1:2:3:5]. Bromotolaidine sulphonic acid. From o-tolaidine sulphonic acid and bromine-water (Nevilo a. Winther, C. J. 37, 627). Prisms.

Reactions. -1. Converted by diago-reaction into a bromo tolurno sulphonic acid whose sulphochloride melts at [56"] and whose amide at 1147°L 2. Heated with HClat160 'it gives dibromo-o-toluidine [410] and two mono-bromo-toluidines, an oil and a crystalline body, [549-579]. One of these bromo-toluidines must be obtained from the acid by displacement of SO₅H by II, and must subsequently give rise to the other mono and the di-bromo tolaidine. Inasmuch a : oily brome telefiline, C.H.Mc(NIL)Br [1:2:3], heated with HCl nt 160° gives n substance [10° 47°] and crystals [53° 55°], while the crystalline bromo-toluidine [54°-57°] is not affected by this treatment, we must suppose the sulphonic scid to be G,U,(GH,)(NH,)BrSO,H [1:2,3;5] rather than [1:2;5;3]. When it is heated with HCl the oily broug-tolniding is first formed, but this being unstable changes into its crystalline isomeride and the crystalline dibromo-pluiding.

Brown amido tolueno sulphonic acid C, II, Mc(NII,) Br(SO, II) [1:4:2:3 or 5]. S. 532 at 21". From o bromo toluene by sulphonation, nitration, and reduction (Schäfer, A. 174, 360). Trimetric lamine (from hot water) or nodules (from alcohol). Converted by bromine-water into tri-bromo tolnidine [823].

Salts. BaA', aq. PbA', aq. Bromo amido tolucuo sulphonio acid C.H.Mc(NH.)Br(SO,H). [1:57:4:3]. From (1,4,2)-bromo-tolucne sulphonic acid by intration, and reduction (S.). Needles (from water). V. sl. sol. water. Ba A', 4nq.

Bromo amido toluene sulphonio acid

C.H.Mc(NH.)Br(SO,H) [1:22,4:6]. S. 3-2. From (1,1,3)-brome telucie sulphonic acid by nitration and reduction (Schäfer, A. 174, 360). Prisms (from water). Browning water gives tri-bronno-tolnidine (72).— BaA\$ 2aq. - NaA', 2aq.

Bromo amido tolneno sulphonic acid C.H.Me(NH.)Br(SO.H) [1:4:x;2]. S. 23 at 20. From boiling aqueous p-toluidine o-sulphonic acid and bromine (Jenssen, A. 172, 230; B. 7, 55). Needles (from water). V. sl. sol. boiling water .-- KA' aq. -- BaA', 7nq.

Bromo amido-toluone sulphonic acid C.H.Me(NII.)Br(SO.H). S. 19 at 20°. From o-toluibue sulphonic acid by conversion into o-

duction (Pagel, 4. 176, 800). Minute this prisms. BaA's aq.

Bromo-amido-toluene sulphonic acid C.H.Me(NH2)Br(SO3H) [1:4:57:3] Formed, together with di-bromotoluidine, by passing brominc-vapour into cold aqueous p-toluidine m-sulphonio acid (v. Pechmann, A. 173, 210; Limpricht, B. 7, 452). Clumps of needles. V. e. sol. water. Converted by diazo- reaction into (3 or 5, 1, 2) bromo toluene sulphonio scid. -KA'.-BaA', 2aq -PbA', -AgA'.

Di-bromo-o-amido-tolueae snIphonio acid C,HMe(NH,)Br,SO,H [1:2:7:7:4]. S. 64 at From o-tolnidine p-sulphonio acid C.H.Me(NH2)(SO3H) [1:2:4] and bromine (Hayduck, B. 7, 1353; A. 172, 211). Capillary needles containing aq (from water). - BaA', 9aq.

Di bromo amido tolucne sulphonic acid C.HMe(NH.)Br.(SO.H) [1:2:37:7:57]. From otolnidine by sulphonation and bromination. Needles (containing aq). V. sol. hot water and hot nicohol. Gives tri bromo-toluidino [112°] when distilled with KOH .- BaA', 4aq .-PbA', 3aq (Gerver, A. 169, 380).

Bromo-di-amido-toluene salphonic acid C.HMe(NH2)2BrSO4H [1:27:67:37:4]. From tolneno p-sulphonio acid by nitration and reduction (Seliwnnert, A. 186, 360). Tables; sl. sol, water. - KA'2 nq.

DI-BROMO AMYL ALCOHOL C, H, Br, O i.e. CH, Br, CHBr, CHELOH. Di-bromo-di-cthyl-carbinol. From vinyl-di-ethyl-carbinol and bromine

(Wagner, J. R. 16, 320). Non-volatile liquid. BROMO-o-AMYL-ANTHRACENE

nation of anyl authracene in CS... Picricacid compound. [110°]. Orangeyelfow needles (Liebermann a. Tobias, B. 14, 797).

BROMO-sec-AMYL-BENZENE C, 11, Br i.e. C.H. CHELCHBr.CH₃(?). (c. 79°) at 40 mm. 8.G. 21 P28. Oil. Obtained by brominating sec-unyl-benzeno (Dufert, M. 4, 620). Decomposed slowly by boiling water into HBr and pentenyl-benzene.

γ-δ-di-bromo-amyl benzeno [53°-54°] Ph.CHBr.CHBr.CH, CH, CH, From phenylamylene (q. v.) and bromine. Needles or plates. γ-δ-di-bromo-isoamy I-bcuzeno

Ph.CHBr.CHBr.CH(CH₂), [128°-129°]. From phenyl-iso-amyleno and Br (Schramm, A. 218, 394). Needles (from alcohol).

Tri-bromo-isosmyl-benzene C, H, Br. (140°) Obtained by brominating iso-amyl-benzene at 100 (Bigot a. Fittig, A. 141, 161). Needles.

BROMO-IŞOAMYLENF, Callaisr. Pentenylbromide. (1003-1109). From 'isoamylene' by successive treatment with bromine and alcoholic KOIT (Bauer, A. 120, 167). Successive treatment with cone. Il, SO, and water forms amylene dibromide and a ketone C. H, O (Bouchardat, C. R. 93, 316).

ω Bromo-iso-amylene (111°) Pr.CH:CHBr. From isovaleric aldehydo by successive treatment with PCl, Br, and alcoholic KOH (Bruylants, B. 8, 406).

7. Brome-n-amylene CH2:CBr.CH2.OH2.CH brome toluene sulphonic acid, nitration, and re- (173°). S.G. 110. From mothyl propyl ketone

by successive treatment with POLBr, and alcoholic KOH (B.).

Bromo-amyleus C, H, Br. (111°). From dibromo-bezoic acid CH, CHBr. CBrEt. CO, H and cold aqueous Na_.CO₃ (Fittig, A. 200, 36). Brome-amylene C₃H₃Br. (115°). From iso-

valerylene and HBr (Reboul, Z. 1867, 173).

Bromo-amylene C, H, Br. (c. 106'). S.G. 15 1.173. From isoprene C.H. and HBr (Bonchardat, J. 1879, 577).

DI-bromo-amylene C.H.Br., (c. 170°). From

isovalerylene and bromine (Reboul, A. 135, 372).

Di-bromo-amylene C₃H₃Br₂i.e. Pr.Clir.CliBr. 175°). From isopropyl-acetylene and Br (Bruylants, B. 8, 407).

Di-bromo-amylene C.H.Br., i.e. Pr.CBr:CHBr.

(190°). From n-propyl-acetylene and Br (B.).

BROMO-AMYLENE GLYCOL C.11, Br(OH). From amylene dibromide by successive treatment with AgOAc and solid KOH (Bauer, J. 1861, 664).

TBI.BROMO.p. ISOAMYL.TOLUENE

C₁₂H₁₅Br₃ i.e. C₅HMeBr₂.C₅H₁₁. From p-isoamyltoluene and bromine at 100°. Sticky liquid (Bigot a. Fittig, A. 141, 135).

TRI-BROMO-ANETHOL C. II, Br.O. From anothol and Br. Crystals (Cahours, A. 41, 60).

BROMO-ANILIC ACID v. DI BROMO DI OXY-

o-BROMO ANILINE Call BrN i.e.

C,H,Br(NH2) [1;2]. Mol. w. 172. [319]. (2299 i. V.). From o-bromo-nitro-benzene by reduction with tin and HCl (Fittig a. Mager, B. 7, 1175).

Acetyl derivative Call BiNIIAc. o. Bromo-acctanili le. [99°]. Long needles (Körner, G. 4, 330).

m. Bromo anilino CaHaBr(NHa) [1:3]. [180]. (251°). Formation. 1. From m brome nitro-zene (Fittig a. Mager, B. 8, 361). 2. m-nitrodiazo-benzene sulphate (from m-nitraniline) is treated with a hot solution of cuprous bromide and the crude m bromo-nitro benzene reduced with tin and HCl (Sandmeyer, B. 18, 1495).

Acetyl derivative C.H.BrNHAc. [88] Needles (from dilute alcohol) (Scheufelen, A.

231, 175).

p.Bromo-aniline C_all_aBr(NH_a) [1:4]. [62].
 (Mills, P. M. [5] 14, 27). [66] (Körner, J. 1875.

Formation .- 1. By reducing p-bromo-nitrobenzene with tin and HCl (Riche a. Bérard, A. 133, 52; Fittig a. Mager, B. 7, 1175; 8, 361).-2. By bromination of acetanilide and distillation of the product with KOH (Mills, P. 4. 4] 19, 21; Pr. 10, 589) .- 3. By distilling bromoisatin with KOH (Hofmann, A. 53, 42) .- 4. In imall quantity by brominating aniline with Br vapour or in benzeno solution (Kekulé, Z. 1866, 387).

in boiling, giving aniline, and di- and tri-bromo-

miline.

Reactions. -1. HCl at 160° gives aniline and li-bromo-aniline (Fittig a. Buchner, A. 188, 23). L Sodium acts on it in ethercal solution formng benzene-azo-benzene, aniline, and NaBr Anschütz a. Schultz, B. 9, 1398; Claus a. loques, B. 16, 909).—3. By dissolving in cooled uming HNO, it is converted into tri-nitro-aniline pleramide) (Hager, B. 18, 2578) .- 4. By nitra-

tion of p-bromaniline dissolved in 10 pts. of H.SO, bromo-nitro-aniline C.H.Br(NO.)(NH.) [4:3:1] is produced (Nölting a. Collin, B. 17, 266).—5. Cyanogen forms a compound C.H.Br.NH.C.NH.).C.NH.NHC.H.Br [245°]; white plates (from alcohol) (Sent, J. pr. [2] 85, 530).

Salts .- B'HCl; monoclinic .- B'.H.PtCl. B'HBr taq: monoclinic prisms (Stacdel, B. 16, 28).—B',H,SO4: lamine.—B',H,CO4.

Formyl derivative Cli Br. NII (CO.II). [119]. Long white needles. Insol. cold water, sl. sol. hot water, v. sol. alcohol and ether. Prepared by heating p-bromaniline with formic ether, or by brominating formanilide (Demustedt, B. 13, 231).

Thioformyl derivative

C.H.Br.NH(CS.H) [1907]. Needles. V. sol. hot ether and alcohol. Prepared by the action of P.S. on the preceding body (Dennstedt, B. 13, 236).

Acetyl derivative [166°]. From acetanilido in glacial HOAo and Br (Remmers, B. 7, 346; Gürke, B. 8, 1114). Also from p-bromo-aniline and AcCl (Korner, G. 4, 329), or acetamide (Kelbe, B. 16, 1200). Monoclinic prisms: a:b:e= 1:538:1:1:435 (Panebianco, U. 9, 357). M. sol. alcohol, v. sl. sol. water.

Isobutyryl derivative

C.11, Br. N11. C. II, O. [1282]. From isolutyrylanilino and Br vapour (Norton, Am. 7, 117). Needles (from alcohol).

Benkoul dericative C.H.Br(NHBz). [202']. From benzamilide and Br (Meinecko, B. 8, 564).

Oxalyl derivative C.O. (NH.C.H.Br), [above 200]. From Br and examilide in HOA. (Dyer a. Mixter, Am. 8, 351).

Di-brome-aniline C.H.Br (N11.) [1:3:4]. [79°]. Formation. - 1. By saponitying dibrominated acetanilide (Griess, A. 121, 266), or brominated o- or p-1. o.nc-acctanilide (Körner, G. 4, 329).— 2. Together with mono- and tri-bromo-aniline by brominating aniline (Kekulé, K. 2, 635). -3. By distilling di-bromo-isatin with KOH (Hofmann, A. 53, 47). -4. Together with mono- and tribrome-aniline by heating nitrobenzene with conc. HBrAq at 190° (Baumhauer, B. 2, 122; Z. [2] 5, 198). 5. By reducing di-bromo nitro-benzene (Wurster, B. 6, 1491).

Properties.-Needles or long plates. Salts Bilch B'll, ltCl .- B', ll, SO, Acetyl derivative Call, Br, NIIAc. [146] Benzoyl derivative C.Il. Br. NBz. [1:3:4]? [1312]. From benzanilide and Br (Hübner, B. 10, 1710). s-di-bromo-anilue Callatra(NIL) [1:3:5].

(57°). From p-nitro acctanilide, by successive conversion into nitro-ani ne, di-bromo-nitro-Properties.—Trimetrie crystals; insol. oold amiline, and di-bromo-nitro benzene, and reduc-water, v. aol. alcohol and other. Decomposed tion of the product (Körner, U. 4, 368; Langer, A. 215, 116). Necdles.

Dl - bromo aniline C.H.Br. (NH2) [1:2:4]. [80°]. From the corresponding di-bromo-nitro-

benzene [58°] (Korner, G. 4, 370).

Di - bromo - aniline C_bH₂Br₂(NH₂) [1:4:3]. [52°]. From the corresponding nitro-compound [85°] (Riche a. Berard, C. R. 59, 141; Meyer a. Stuber, A. 165, 180).

Tri-bromo-aniline C.H.Br.(NH2) [1:8:5:6]. [119°]. (300°).

-, we worken or broming on aniline or on an aqueous solution of a salt of aniline (Fritzsche, A. 44, 291; J. pr. 28, 204; Hofmann, A. 53, 50).-2. By the action of Br on o- or p-bromo aniline (Körner, G. 4, 305). 8. By reduction of tri-bromo-nitro-benzene (K.).

4. Together with aniline and di-bromo-aniline by the dry distillation of p-bromo-aniline or by

heating it with HClAq at 160° (Buchner, P. 8. Preparation.-Bromine (500 g.) is gradually

added to aniline (100 g.); as soon as a solid mass is formed glacial acetic acid is added and the addition of bronsino continued until the mass The product is washed has a red colour. with dilute alcohol and water and crystallised

from alcohol (H. Silberstein, J. pr. [2] 27, 101). Yield 80 per cent. Properties .- Long colourless needles.

aniline half dissolved, half suspended in alcohol. gives a yellow pp. of tribromo diazo benzone nitrate .- 2. Converted by diazo-reactions into s-tri-bromo-benzene and into u-tetra-bromo-benzene .- 3. Conc. 11NO, gives di-bromo-di-nitromethane, tetra bromo quinone, u-tetra-bromo-

Reactions .- 1. N.O. passed into tri-bromo-

benzene, picric acid, and oxalic acid (Losanitsch, B. 15, 472). Sults. The salts are very unstable, and cannot be formed in agreens solution. B'HBr: small white needles [1903], insol, other and

benzene; decomposed by water .- B'HCI: small white needles (Gattermann, B. 16, 635). Acetyl derivative C.H.Br. NIIAc. [232]

(Renmers, B. 7, 319).

Diacetylderivative C.H.Br. NAc. [123°]. Tri-brome aniline C It Br (NIL) [1:2:3:5].

[ahove 130]. From tri bromo nitro benzene [112°] (Körner, (1. 4, 328). Its salts are not decomposed by water.

Tetra-breme aniline C_HBr_(NIL.) [1:2:3:5:6]. [115°]. Fermed by brominating m-bromoaniline, or (1, 4, 3) di bromo aniline (Körner, G. 4, 328; Wurster a. Nölting, B. 7, 1564). Is a by-product in the action of bromine on nitrobenzene in presence of Fe Br. (Schenfelen, A. 231, 161). Needles, Converted by the dinze-- reaction into u-tetra-bromo-bensene.

Penta-brome-aniline C. Br. NH. [222]. Obtained by brominating (1, 3, 5) di bromo aniline (Körner, G. 4, 368). Large needles (from alcohol mixed with toluene)! BROMO-ANILINE SULPHONIC ACID v.

BROMO-AMIDO-BRNZENE SULPHONIC ACID. BROMO-ANISIC ACID v. Methyl derivative

of BROMO-OXY-BENZOIC ACID. BROMO-ANISIDINE v.

Methyl-Baomo-AMIDO-PHENOL.

BROMO ANTHRAGENE O, H, Br. Obtained by warming anthraceue dibromide. Needles. Forms a red pierie acid compound. (A. 1,2) - Di-bromo-anthracene C., H. Br., [221°].

From Br and authracene in CS. (Graebe a. Liebermann, A. Suppl. 7, 275). From triphenylmethano in CS, and bromine in sunlight (triphenyl-methyl bromide being also formed) (Allen a. Keillace, A. 227, 109; 228, 251). Golden bromine upon anthranol (1 mol.) dissolved in CS, needles (from toluone); may be amblimed. V. sl. Yellowish crystals. Insol. boiling aqueous, but sol. alcohol and ether. Alcoholic KOH at 100° forms authracene. Dilute HNO, gives anthra-

quinons. The compound with pieric acid is re Combines with Br forming O. H.Br. [170°-180° decomposed at 180° into tri brome anthracem HBr, and Br.

Di-brome-anthracene C, H, Br, [B. 1:2]? [192°] From di-bromo-anthraquinone, HI, and P at 150° (Miller, A. 182, 367). Golden tables (from alcohol). Oxidises to di-bromo-anthraquinone.

Tri-bromo-anthracene C, H, Br, (A. 1, 2, B.) [169°]. Formed by heating (A. 1, 2)-di-bromo-anthracene tetrabromide (G. a. L.). Yellow needlea (from henzene). Oxidised by HNO₄ to

bromo anthraquinone (Claus a. Hertol, B. 14, 979).—Bromide: C.H.Br., Tetra-bromo-anthracene

C₁,H₂Br₄(A. 1, 2, B. 1, 2)? [254°]. From di-bromo-authracene tetrabromide and alcoholic KOH (Anderson, A. 122, 304; G. a. L.). Yellow needlea (from xylene). Gives di-bromo-anthraquinone on oxidation. - Bromide: C,H,Br, [2120] (Hammerschlag, B. 10, 1212).
Penta-bromo-anthracene C₁₁H Br. [212°].

Formed by heating tetra-brome-authracene tetrabromide at 230° (II.). Yellow powder. Oxidisos to tetra bromo-authraquinone. [310°-Hexa-breme-anthracene C, H Br.

320°]. Prepared by the action of Br in presence of I on di bromo-anthracene (Diehl, B. 11, 178). Oxidised by K.Cr.O. and H.SO. to tetra-bromoanthragninous fc. 300 1.

Hexa breme anthracene Cull Br. 370°]. From Cull, Br, and alcoholic NaOlI (H.). Silky yellow needles (from kerosene). Oxidation gives tetra bromo anthraquinone [above 370°]. Hepta-bremo-anthracene Cull Br. [above

350°). Prepared by the prolonged action of Br at 200° in presence of I on di-bromo-anthracene (D.). Yellow needles. Insol. alcohol and other.

Ooto-brome-authracene Call Br. Formed by very prolonged action of iodine bromide at 360 on kepta bromo-anthiacene (D.). Needles. Insol.

most ordinary solvents, al. sol. nitro benzene and aniline. (A. 1) - BROMO - ANTHRACENE-(A. 2)-CAR-

BOXYLIC ACID C, II, C(CO, II) C, II, [266°]. Formed by the action of bromine upon anthra-

cene-(A.)-carboxylic acid in acetic acid. Long glistening yellow needles. Sublimable. Sol. alcohol, ether, and acetic acid, v. sl. sol, benzene and water. Its solutions have a blue fluorescence. Evolves CO2 at its melting - point leaving bromo-anthracene.

Salts .- AgA': yellow microscopic prisms. -KA': long very slender yellow needles.-BaA': yellowish needles (Behla, B. 20, 704).

DI-BROMO ANTHRACENE DISULPHONIC ACID O, H.Br. (SO, H), From di-bromo anthra-ceno and funing ILSO, (Perkin, C. J. 24, 19). Oxidises to anthraquinone disulphome acid.

Salts.-Na A' .- BaA". BROMU ANTHRANOL C. H.OBr i.e.

 $C_{e}H < CO > C_{e}H$, or $C_{e}H < CO > C_{e}H$

[148°-151°]. Formed by the action of (1 mol. of)

Di-brome-anthranol C, H,OBr, &c.

C.H. CBr. O.H. Anthraquinons - bromide. [157°]. Formed by the action of (rather more than 2 mol. of) bromins upon anthranol (I mol.) dissolved in CS. Large rhombic crystals. Insol. aqueons alkalis. By boiling with neetic acid it is readily converted into anthraquinone

(B. 1)-BROMO-ANTHRAQUINONE

C₁₄H,BrO₂ [188°]. Yellow needles. Sublimable. Prepared by heating o-bromo-henzoylbenzoic acid C.H.Br.CO.C.H.CO.H with H.SO. to 180°. On fusion with KOII it gives erythrooxy-anthraquinoue (Pechmann, B. 12, 2127).

(B. 2) - Brome - anthraquinone C, 11 Bro.. [187°]. Formed by exidation of tri-brome anthracene with Cro. and HOAc (Graebe a. Liebermaan, A. Suppl. 7, 290). Yellow needles; may be sublimed; sl. sol, alcohol, m. sol, het benzene. Converted by potash-fusion into alizarin.

Di-bromo-anthraquiaone C, II.O.Br. [265°]. Formation. — 1. By brominating anthra-quinone (Graebe a. Liebermann, A. Suppl. 7. 289; Diehl, B. 11, 181). 2. By oxidation of tetra-bromo-anthracene or di-chloro-di-bromoanthracene with CrO, (G.a. L.; Hamaiersching, B. 19, 1107).

Properties. - Boils with slight decomposition, st. sol. boiling alcohol, m. sol. glacial HOAc. Potash-fusion at 250° gives alizarin. According to Perkin (C. J. 37, 551; priv. com.) the di-bromo anthraquinone formed by the first method nielts at 215° (or 250° cor.) and differs from that formed by the second method, which melts at 275° (or 281.5° cor.) by giving, on potash-fusion, not only alizarin but also flavopurpurin and anthrapurpuria (tri-oxy anthraquinone).

Tri-brome anthraquinone C14H, Br, O2. [186] Prepared by the action of Br in presence of I upon anthraquiaoae or di-bromo-anthraquimme at 260° (Diehl, B. 11, 182). Yellow medles; insol. alcohol.

Tri - brome - aathraquinone C₁₄H Br₈O₂, [5°]. From penta brome-anthracene, CrO₂, and HOAc (Hammerschlag, B. 10, 1213). Flat

nsedles (by sublimation).

Tstra - bromo anthraquinous C, II, Br,O, [295°-800°]. Formed by oxidising hexa bromo. anthraceae [320°] or by brominating tri-bromo-anthraquiaoae [186°] in presence of I (D.). Yollow needles.

Tetra - bromo - anthraquinone C, II, Br, Oz. [above 370°]. Formed by oxidising hexa-bromoanthraceno [above 370°] (H.). Yellow needles.

Penta - bromo - anthragninons C, Il Br.O. Formed by oxidation of hepta-bromo-authracene (D.). Sublinees with difficulty without

melting; sl. sol. boiling toluene DI. BROMO. DIANTHEYL C., H., Br., [far abovs 300]. Yellow prisms (from toluene). Formed by bromination of dianthryl dissolved in CS. (Liebernana a. Gimbel, B. 20, 1855). DI-BROMO - APOPHYLLENIC ACID

O,H,BrNO, i.e. C,NH,Br(CO,H)(CO,Me) C,NH,MeBr(CO,H) CO . Methyl other of bromo pyridine dicarboxylic acid? From bromo-C.N.11.0.8 (Trzeinski, B. 16, 1057).—7. Potastaronine (a derivative of nareotins) and Br sum sulphocyanids forms sulphocyano-harbi-(v. Gerichten, A. 210, 91). Hard prisms (con turio acid C.N.11, SO, (T.).

taining 2sq); sol. hot water. Its solutions give no pps. with salts of Pb. Ag, or Cu. Heated with cone. HCl it forms CO₂, MsCl, and a brome-pyridine carboxylio acid (?) [199°].

Salt.—BaA', 3aq. DI BROMO APOPHYLLIN C, II, Br, N, O, [229°]. From bromo-tarconine or bromo-apophyllenic acid and bromine-water (v. Gerichten. A. 210, 91). Six sided tables (containing 4aq); sol, water, insol. other. Alkalis form a deep-red solution. HCl at 150° gives CO, McCl, di-bromo-pyridine, and di-bromo-pyridine mothylo-chloride.

Salts. -B"HCl .- B"2HCl .- P", H .P | Cl . aq. .-B"HBr. -B"2HBr.

DI - BROMO - ATROLACTIC ACID v. 88-Dr. BROMO G-OXY-A-PRENYL-PROPIONIC ACID.

BROMO - ATROPIC ACID Call, Bro. i.e. Ph.C.HBr.CO.H. Exo-bromo a phenyl-acrylic acid. (130). From the compound of atropio acid (1 mol.) with IIBr (2 mols.) by boiling with water (Fitlig a. Wurster, A. 195, 162). Slender needles (from water).

TETRA BROMO AURIN Cullin Br.Ox. Formed by brominating aurin dissolved in 110Ac (Dale a. Schorlemmer, C. J. 35, 152; cf. Zulkowsky, M. 3, 465). Bronzed crystals. Alkalis form violet solutions.

Salt .-- A"Ag: dark violet insoluble pp. Ethul ether A"Et,: [1109-115"]; microscopic reddish crystals; sol. alcohol, ether, and

BROMO-AZO BENZENE E. pp. 874, 379, TETRA-BROMO-AZOPHENINE C., pl. 11, Ir, N. [243"], Formed by heating p-nitroso-di-phenylamine with p-bromo-uniline and p-bromo unilina

hydroculoride at 100°. Greatly resembles azo-phenine (O. Fischer a. Hepp, B. 20, 2181). BROMO-AZO-TOLUENE v. p. 391. DI-BROMO-AZOXY-BENZENE v. p. 427.

BROMO AZOXY-TOLUENE v. p. 428, BROLIO - MARBITURIC ACID C. 11, BrN.O. i.e. CHBr CO.NH CO. Bromo-malonyl-urea.

Ureide of bromo malonic acid. From di-bromobarbituric acid by reduction with Na or Zn, or by evaporating with aqueous HCN (Baeyer, A. 130, 134). Small needles; sl. sol. cold

Salts .- NII,A'. - ZnA', 8aq. - ZnA', 6aq (Mulder, B. 12, 2309).

Di brema barbit irie neid C,11 Br. N.O. Dibromo-malonyd-urea. Formed by the action of Br on barbituric, nitro-barbituric, nitroso-barbituric, and hydurilic soids (Baeyer, A. 127, 199; 130, 130).

Preparation .- From di-bromo oxy methylnracil and faming HNO, (Behrend, A. 236,

Properties .- Trimetric crystals. Sol. water, y. sol. hot alcohol and ether.

Reactions .-- 1. Boiling water forms alloxan and HBr .- 2. Ill reduces it to hydurilic and barbiturie acids. -3. H.S forms dialuric acid. --4. Bromine water forms CO, and tri-bromo-acetyl urea. - 5. Alkalis form CO, tri-bromoacotyl urea, and bromo barbituric acid. - 6. Thio urea forms so called this pseudo-urio acid Di bromo-di-barbituric acid C_bH₃Br₂N₁O₅ aq. Prisms (Baeyer, A. 13O, 145; of. p. 44O).— C_bH₃Br₁N₁O₄HBr. Prisms, v. sl. sol. water. Di-BROMO-SEHENIC ACID C_cH₄Br₂O₅.

[43°]. From erucic acid O_nH_aO₂ and bromine (Haussknecht, A. 143, 40; Otto, A. 135, 226). Nodules. Converted by alcoholic KOH to bromoerucio acid, CallaBrO, and behenolic acid C_{in}H_nO_i Moist A_iC_iO forms oxy erucic acid and di-exy-behenic acid. Sodium amalgam forms erucic acid. Salte.—BaA'_x.—PbA'_x.

Di-breme-bohenic acid C_{in}H_nBr_iO_i, [54°]. From brassic acid and bromine (II.). Reduced

by sodium-amalgam to brassic acid. Alcoholic

KOH at 220° gives behenolie acid.

Tri-bremo-behenic acid C. H. Br.O... From bromo-erucie acid and bromine (H.). Tetra-breme-behenic acid C.21f, Gr,O2. [78]. From behenolic acid and bromine

lamine (from alcohol). BROMO-BENZALDEHYDE v. BROMO-BENZOIC

ALDREYDE. BROMO-BENZAMIDE v. Amide of BROMO-

BENZOIC ACID.

BROMO. BENZANILIDE v. Anilide of BROMO-BENZOIC ACID.

BROMO-BENZENE C₀1I₀Br. Phenyl bromide. Mot. w, 157. (156°), S.G. ⁹" 1-1911. µ_S 1-5736. R 25-81 (Bruhl). S.V. 119-9 (Schiff, B. 19, Vapour pressure : Ramsay a. Young, C. J. 47,616.

Formation.-1. From bromine and benzeue, cither holling or in sanlight (Couper, A. Ch. [3] 52, 309; Seliranm, B. 18, 606). The reaction is promoted by the presence of 1 (Ador a. Rilliet, B. 8, 1287) or Al₂Cl₄ (Greene, C. R. 90, 40; Leroy, Bl. (2) 48, 210).—2. Formed by running a solution of NaNO, into a hot solution of Cu,Br, and aniline in dilute H.SO, (Sandmeyer, B. 17, 2652). 3. By heating diagobengene with a large excess of strong HBr; the yield is 32 p.c. of the theoretical (Gasiorowski a. Waj'ss, B. 18, 1938). 4. From phonol and bromido of phosphorus (Riche, A. 121, 357).

Properties. -Oil; not attacked by KOII, KCy, or AgOAc.

Reactions. -1. Boiling with Al_Cl, produces benzene and di-bromo bonzenes (Dumreicher, B. 15, 1867). - 2. Converted in the animal system into p-bromo-phenyl-mercapturio acid Calla BrNSO, p-bromo phenol, brome pyrocateclim, and brome-hydroquinone (Batmann a. Prensse, H. 3, 156; Jaffe, B. 12, 1092).—3. Converted by cone. H₄SO₄ into di-brome-benzone sulphonic acid and brome benzene di-sulphonic acid (Herzig, M. 2, 192).—4. ClSO₂H forms C.H.Br(SO₂H) [1:4] and C₂H₂Br.SO₂C₄H₄Br theckurts a. Otto, B. 11, 2061).—5. Heating with McONn gives Call OMe and phonol (Blau, M. 7, 621). According to Mittien (B. 17, 2631) there is a second bromo-benzene [c. 62], o.Di-breme-benzene C_sII_sBr_s [1:3].

(221°). S.G. 2 2.003. Formed in small quantity in preparing p-di-bromo benzene from benzeue and Br (Riese, B. 2, 61; A. 164, 176). Formed also from o nitraniline vid o bromonitro-benzene (Körner, G. 4, 833). HNO, forms chiefly di-bromo-nitro-benzene [58°

m-Di-bromo-benzene C.H.Br. [1:3]. (219°). B.G. 188 1-955.

Formation.—1. From (1,8,4)-di-bromo-ani-line (Meyer a. Stüber, A. 165, 169).—2. From s-di-bromo-aniline.—3. From m-di-nitro-benzene vid m-nitro-aniline, m-bromo-nitro-benzene. and m-bromo-aniline (K.) .- 4. Together with the p- isomeride by the action of Br on benzene in presence of Al.Cl, (Leroy, Bl. [2] 48, 211).

Properties.—Has not been solidified. Yields

with HNO, two nitro-derivatives [62°] and [83°]. Sodium acting on an ethereal solution forms diphenyl, C₁₈H₂,Br₂ (220°) and C₁₈H₂₂Br₂ [250°], (Goldschmiedt, M. 7, 45).

p. Di-bromo-benzene C₂H₄Br₂ [1:4]. [89°].

(219°). S.V.S. 127.8 (Schiff).

Formation.-1. By bromination of benzene or bromo-benzene (Couper, A. Ch. [3] 52, 309; Richo a. Bérard, A. 133, 51; Riese, A. 164, 162; Jannasch, B. 10, 1355).—2. From p-diazo-bromobenzene perbromide by heating with alcohol (Griess, C. J. 20, 66).—3. From p-bromo-phenol and PBr. (A. Mayer, A. 137, 219) .- 4. From ben-

zeno, Br, and Fo,Cl_s (Scheufelen, A. 231, 188).

Preparation. -1. By treating benzeue with Br and a little 1 .- 2. Dromine (960 g.) is added slowly to benzene (210 g.) and Al₂Cl₆ (30 g.). On adding water in excess, the product deposited consists of crystals of the para-compound, with an oil, consisting chiefly of the meta-compound. This oil, however, contains some para-di-bromo-benzene in solution. separate this, the oil is treated with fuming ILSO, The meta-derivative is easily sulphonated in this way, but the para- is not affected. On adding a large excess of water, the solid C.H.Br. [1:4] is ppd., and the clear solution of the meta-sulphonate distilled with steam gives C, H, Br, [1:3] (Leroy, Bl. [2] 48, 211).

Properties .- Prisms or plates, isomorphous with p-di-chloro-benzeno (Friedel, Bl. [2] 11, 38). Sublimable. Sl. sol, alcohol.

Meactions .- 1. Heated for 30 minutes with Al_Cls (! part) gives Catt Br, CsH Br, [1:3], and two tribromobeuzenes, C, H, Br, [1:2:4] and [1:3:5] (L.) -- 2. IINO, forms a nitro derivativo [85°] --3. In ethercal solution sodium forms diphenyl, di-phenyl-benzene, and C₁, H₂, Br₂ [265] and C₁, H₂, Br₂ [300] (Riese, A. 164, 164; Goldschmiedt, At. 7, 42).—4. Me1 and Na form p xylene. -5. NaOEt at 190° gives CollaBroEt, bromo-benzene, and benzene (Balbiano, G. 11, 401).-6. NaOMe forms C, 11, BrOMe, C, H, (OMe), and C₈H₁(OMe)(OH) (Blan, M. 7, 621).-7. Converted by conc. 11.80, into tetra- and hoxabromo-benzene (Herzig, M. 2, 192).

c.Tri - brome - bonzene CallaBr. Mol. w. 315. [87°]. From (1,3,5,2)-di-bromo-nitro aniline vid tri-bromo-nitro-benzene, and tri-bromo-aniline (Körner, G. 4, 401). Tables (from alcohol); may be sublimed.

s-Tri-brome-Senzene C.H.Br., [1:3:5]. [119°]. (278°). Formation.—1. From ordinary tri-bromo-anilino (Stüber, B. 4, 956; A. 165, 173; Reinke, 4. 186, 271; Bassmann, A. 191, 206; Silber tein, J. pr. [2] 27, 104). -2. From (1,3,5)-di-bromo aniline (K.). -3. In small quantity by the action of light on bromo-acetylene (Sabanejeff, J. R. 17, 176).

Properties. - Needles (from alcohol); al. sol. alcohol. Boiling cone. H.SO, forms hexabromo benzenc (Herzig, M. 2, 197). NaOMe forms C.H.Br. (OH) and C.H.Br. OMe [770] (Blau, M. 7, 621).

u-Tri-bromo-benzene C.H.Br. [1:2:4]. [44°]. (275°). Formation.—1. From hexa bromobenzene and alcoholio KOH (Mitscherlich, P. 35, 374; Lassaigne, Rev. Scient. 5, 360.
 From (1,3,1)-di-brono phenol and PBr, (Mayer, A. 137, 224).—3. From (1,3,4)-di-bromoaniline (Griess, Tr. 154, 667). -4. From o., m., and p., di-bromo-benzene by nitration, reduction, and diazotisation (K.) .- 5. Got from each of the dibromobenzenes by heating with water and bromine at 250° (Wroblewsky, A. 192, 220).— 6. From benzene, bromine, and Fe₂Cl₂ (Schenfelen, A. 231, 188).—7. As a by-product, in the preparation of C. H.Br. by the action of Br. upon Call in presence of Al.Cl (Leroy, Bl. [2] 48, 213). Properties. -Needles; sl. sol. alcohol.

s-Tetra-bromo-benzene C_o1f.Br_{*} [1:2:4:5]. Mol. w. 394. [175]. Formation.—1. From pdi bromo benzene and Br at 150? (Riche a. Bérard, A. 133, 51).-2. From n-tri-bromo-benzeno and Br. 3. From nitro-benzene and Br. at 250° (Meyer, B. 15, 16).—1. From benzene (30 g.), Fe Cl. (5 g.), and bromine (210 g.); the mixture must be cooled (Schenfelen, A. 231, 187). Properties. - Long needles (from alcohol).

u-Tetra-bromo-bouzene C.H.Br. (1:2:3:51, [98-5]). (329). Formation.—1. From (1,3,5,2). tri-bromo-phenol and PBr₅ (Körner, A. 137, 218, Mayer, A. 137, 227). 2. From (1,3,5,2) tribromo-amiline (Wurster a. Nolting, B. 7, 1561; V. v. Richter, B. 8, 1428; Silberstein, J. pr. [2] 27, 118). 3. From C. [HBr. SO H by heating with HC! (Bits mann, A. 191, 221). 4. One of the products obtained by heating tri bromodiazo-benzene nitrate with benzene (Silberstein, J. pr. [2] 27, 110). Properties. Long needles, v. 8l. sol. abohel. 41NO, (S.G. 150) gives a nitro-derivative [962]; furning HNO, (8.G. 151) gives a di-nitro-derivative [228°].

Totra - bromo - benzene C. II.Br., [160°]. From p-nitro benzoio acid and Br at 280° (Halberstadt, B. 14, 914). Needles (from alcohol).

Tetra - brome - benzene C, H, Br., [138]. From p-di-brome - benzene and boiling cons. H.SO, (Herzig, M. 2, 195). One, if not both, of the two last mentioned bodies is, doubtless, a mixture.

Penta bromo-benzone C. HBr. Mol. w. 473. [260°]. Formation .- 1. From Br and nitrobenzene or di nitro benzene at 250' (Kekule, A. 137, 172). 2. From s-tri-bromo-benzene and fuming H.SO, at 100° (Bässmann, A. 191, 208).—
 From alizarin and BrI at 250° (Diebl, B. 11, 191). Properties. -Silky needles; v. sl. sol. alcohol, sol. benzene.

Hexa - brome - benzene CaBra. Per-brome-benzene. [above 315]. Formation.—1. By the action of Br containing I mon benzene, toluene, or bonzene azo-benzene at 250° 100° (Gessner, B. 9, 1505). 2. From tetra 'romo-quinone and PBr, at 280° (Rooff, B. 10, 103). - 3. From p-dibromo benzene (or tri bromo-benzene) and cone. H₂SO, (Herzig, M. 2, 192).-4. By heating at 360° per-bromo-methane, per-bromo-ethane, perbromo-butane, per-bromo-pentane, or per-aromo-hexane (Merz a. Woith, B. 11, 2235).—5. By heating benzene with bromine and AlaBra .- 6. By heating pentabromo-phenol with PBr, at 260° di amido-benzene sulphonio aci (Merz a. Weith, B. 16, 2890). -7. From bromine 188, 153). Deliquescent prisms.

(800 g.), Fe,Cl, and benzene (17 g.) in the cold. Good yield (110 g. instead of 119 g.) (Scheufelen, A. 231, 189). Properties.—Long needles (from tolueno). V. sl. eol. alcohol and tolueno.

BROMO-BENZENE O-SULPHONIC ACID C.H.Br.SO,H [1:2]. From o-amido-benzene sulphonio acid (Berndsen a. Limpricht, A. 177, 101; Bahlmann, A. 181, 203; 186, 315). Long deliquescent needles; v. c. sol. alcohol.—NILA',—RA'aq. — BaA', aq. — BaA', 2aq. — CaA', 2aq. —

PhA', 3aq. -AgA'. Chloride C. ~C_H_Br.SO_Cl {51°] Amide C.H.Br.SO NIL [1860] Bromo-benzene m-sulphonic ccid

C.H.Br.SO.H [1:3]. From amido-benzeno msulphonic acid (Berudsen, A. 177, 92). Formed also by brominating benzene sulphonic neid or its silver salt (Ross Garrick, Z. (2) 5, 549; Genz, B. 2, 405; Nölting, B. 8, 819; Thomas, A. 186, 133). Very deliquescent. KA'aq. S. (of KA') + 1 at 0'... PbA', 2aq...-PbA', 3aq....S. (of PbA') 578 at 7'...-BaA', 2aq. S. (of BaA') 3'5 at 14'.

CaA', AgA', Chloride Call, Br. SO.Cl. Oil. .1 m i da C.H.Br.SO NII. [151°] Bromo-benzene- p sulphonio acid

C.H.Br.SO,H [1:1] [882]. Deliquescent needles.

Formation.—1. By sulphonating bromobenzene with H.SO, CISO,H, or SO₄ (Couper, C. R. 45, 230; Ross Garrick, Z. [2] 5, 549; Armstrong, Z. [2] 7, 321; Nölting, B. 8, 594).— 2. From p-amido-benzene sulphonio acid (Meyer, A. 156, 291; Berndsen, A. 177, 92). Reactions .-- The K salt distilled with K FeCy.

gives terephthalonitrile. Salts. (Goslich, B. 8, 352; A. 180, 93),— NII,A', KA', CaA', 2aq, BaA', FaA', 2aq,— PbA',—ZaA', 6aq,—CaA', 6aq,—AgA', Chloride C_bH_bBr,SO₂Cl. (75°).

Amide [1616] (Bilsstoann, A. 191, 217). Aceinteamide C.H.Br.SO,NHAc [1999]. Anviide C. 11, Br. SO, NHPh [119"]. Bromo-benzeno di-sulphonic acid

C_eH_sBr(SO_sH)_{r [1}:3:4?]. From the corresponding amido-benzene disulphonic acid (Zander, A. 198, 28). Deliquescent needles. - BaA' Baq. -PbA" act.

Chloride Call, Br(SO,Cl), [1013]. Amide C,11,Br(SO,N11), (210 Bromo bonzene di-sulphonic acid

C.H.Br(SO,II), [1:2:4].
Formetion. - I. From the corresponding ancido benzene disulphonic acid (Zander, A. 198, 10).—2. Fran bromo benzene p-sulphonia acid and SO, at 210° (Nölting, B. 7, 1311).

Properties.—Slender deliquescent needles, K,A"aq. - BaA" 4aq. S. (of BaA") 6 at 22°.— $Ag_{x}\Lambda^{\alpha}$

Chloride C.H.Br.(O Cl), [103°]. Amide C. H. Br(SO, N. 1.), [239] Bromo-benzene di-sulphonio acid

CaH,Br(SO,H)2 [1 or 4:5:3]. From the corresponding middo - benzeno disulptionio acid (Heinzelman, A. 188, 177). Deliquescent needles.
Salts. -(NII), A".—BaA"2]aq.—PbA"2{aq.
Chloride C, H,Br(SO,Cl), [99°].
Amide C, H,Br(SO,Cl), [246°]. o-Di-bromo-benzene sniphonic acid

C.H.Br. (SO.H) [1:2:3]. From the corresponding di amido-benzene sulphonio acid (Sachse, A. Saits.—KA'.—BaA', Saq. S. (or
 143 at 10°.—CaA', 2aq.—PhA', Saq.
 Chłoride CH,Br., SO,Cl. [127°].
 Amids O,H,Br., SO,NH, [215°].
 Dil-bremo-bensans sulphonic acid
 H. D. (Orthograms). S. (of BaA'2)

C,H,Br,(SO,H) [1:2:4]. [67']. From eilver bromo-benzene m. or p. sulphonate and bromino (Limpricht a. Goslich, A. 186, 145). From C.H.Br(NH2)SO.H by diazo-reaction (Langfurth, A. 191, 179; Spiegelberg, A. 197, 263). The free

acid orystallises with 3uq [65°].
Salts. — RA'. — NH, A'. — BaA', 2aq. —
BaA', 3aq. S. (of BaA',) 3 at 11°.—CaA'z.—
1°bA', 2uq. S. (of PbA_), 20 at 7°.—AgA'.

Chlorids C,H,Br,SO,Cl. Oil.

Amide C. H. Br. SO, NII. [1750]. m-Di-bromo-banzene anlphoule acld

C.H.Br.(SO.H) [1:3:5]. From dibrominated oor p. amido bouzene sulphonic acid by the diazoreaction (Limpricht, B. 8, 1066; A. 181, 201; Schmitt, A. 120, 158; Lenz, A. 181, 23). It is also a product of the action of conc. II,SO, on bromo-benzone (Herzig, M. 2, 192). Crystalline.

Salts. - NH₄A'. - KA'. - DaA'₂3₂aq. S. (of BaA'₂) ·28 at 18°. - CaA'₂3₂aq. - PbA'₂ 1₂aq. S. (of PbA',) 21 at 18°,

Chloride C. 11, Br., SO, Cl. [57:5°]. Amide C. 11, Br., SO, N11., [203°].

m Dl. bromo-bonzens salpbonic acid C₄H₃Br₂SO₃H [1:3:4]. [80° when hydrated]; [110° anhydrous]. From C₅H₂Br₂ (NH₂)SO₃H [1:3:4:6] by diazo-reaction (Langfurth, A. 191, 184; Bassmann, A. 191, 232). Slender deliquescent needles (containing aq). Sol. alcohol, insol. other. With HBr at 180 gives H2SO, and m-di-bromo-benzeno.

Salts. - N11, A'. - KA'. - BaA', 21, aq (l.). -BaA', 2aq (B.). S. (anhydrous) 2 at 10° (L.); 1 at 20° (B.). - CaA', 3aq. S. (anhydrous) 3'34 at 10°, --1'bA', 3aq. S. (anhydrous) 1'43 at 8', --AgA'.

Chloride Call, Br. SO, Cl. Amide Calla Br. SO, NII, [190 '] (anhydrous). p. Di brome-benzene salphoale acid O.H.Br.(SO.H) [1:4:2]. [128'].

Formation .- 1. By bromination of silver oor m bromo-benzene sulphonate (Bahhnann, A. 181, 206; Limpricht, A. 186, 139),-2. By sulphonation of p-di-bromo-benzene (Douglas Williams, Z. [2] 7, 302; Hilbner a. Williams, A. 167, 117; Wolz, A. 168, 81) .- 3. From nitrated bromo-benzene o. or meulphonic acid (Thomas, A. 186, 129; B.) .- 4. From brominated amido. benzene o-sulphonio acid (B.). Prisus (containing 3aq), [98°]. Couc. HBr at 250° gives H.SO, and p-di-bromb-benzene.

Salts. (cf. Borne, A. 187, 350).—N11,A'.-Natis: (cr. Borne, A. 1897, 350).—Nil₄A;.—NaA' l'huj. K.Y aq. S. (of KA') 5-79 at 2!?.—AgA' l'aq. AgA' 3aq. S. (of AgA') 1-69 at 10°.—BaA', S. 1°01 at 16°.—BaA', aq.—BaA', 5aq.—CaA', 4aq. S. (of CaA',) 5-90 at 22°.—CaA', 5aq.—CaA', 10aq.—PbA', 3aq. S. (of PbA',) 2-56 at 21°.—PbA', 4aq.—CuA', 11aq.—ChA', 25q. (ch.Logida C. Ha, 180, 180, 1792).

S.(01 PDA.) 2700 at 21.—PDA., 2404.—OLA., 2404.—Chloride C.H., Br., (SO, Cl): [72°].

Amids C.H., Br., (SO, NIL.). [193°].

Anhydride (C.H., Br., SO,), O. Obtained by the action of fuming sulpharic acid upon p-distance of the state of the bromo-benzene. Amorphous powder. Nearly insol. alcohol, ethor, and benzene. By boiling alkalie it is converted into salts of the soid; very slowly attacked by boiling water. POl, con-

verts it into the chloride (Rosenberg, B. 19 658).

m-Di-bromo-benzene di-sulphonic acid C₆H₂Br₂(SO₃H)₂. From C₆H₂(NO₂)₂(SO₂H)₂ (Limpricht, B. 8, 290). Deliquescent.

p-Di-bromo-bensene disulphonio acid p.Di-oromo-bensene dissipation acta
C_slt_Br_s(SO_sH)_s. Formed by sulphonating p.dlbromo-benzene (Borne, A. 187, 366). Small
columns.—K_sA".—BaA" 4½aq.
Chloride C_sH_sBr_s(SO_sCl)_s. [161°].
A mide C_sH_sBr_s(SO_sNH_s)_s. [210°].

Tri-bromo benzene sulphonic acid

 $C_6 H_2 Br_3 SO_3 H$ [1:3:5:6]. [145°] (anhydrous). Formed by sulphonating s-tri-bromo-benzene (Reinke, A. 186, 271; Bässmann, A. 191, 206). Also from C₆H(NH.)Br₅SO₄H[1:2:4:6:3] by diazoreaction (Langfurth, A. 191, 191). Six-sided tables (containing aq) [95°]. Wate forms 11,80, and s-tri-brome-benzene. Water at 180°

Salts.—NH, A' aq.—KA' 3aq.—KA' aq. S. (of KA') :81 at 5.5 (B.).—BaA', 9aq.—BaA', 6aq. (or AA) '81 at b'5 (B.).—BaA', 3aq.—BaA', 5aq.—BaA', 2aq.—BaA', 2aq.—BaA', 2aq.—BaA', 2aq.—BaA', 2aq.—BaA', 2aq.—BaA', 3aq.—BaA', 3a

Chloride. C.H.Br.SO.Cl. [65°]. Amide. C.H.Br.SO.NH. Decomposes above 220°.

Tri-brome-benzens sulphonic acid C_bH₂Br₃·SO₃H [1:2:4:5]. [140° anhydrous]. From (1, 2, 4, 5) or (2, 4, 1, 5) di-bromo amidobenzene sulphonic acid by diaze-reaction (Lang-

furth, A. 191, 188; Reinke, A. 186, 283; Knuth, A. 186, 303; Spiegelberg, A. 197, 282). Needles (containing 3aq) [80°]. With IIBr at 200° gives II.SO, and C. II.Br, [1:2:1].

11.50, 4nd c, 1.61, (1.12.1).

Salts, --Ni, A'aq. --KA'aq. S. (of KA') 1.03

at 20° (B.); 77 at 22° (S.). --BaA', 3aq. S. 122

at 12°, --BaA', 2aq. --BaA', 6aq. --CaA', 6aq. -
1'bA', 4aq. --AgA'.

Chloride. --C, H, Br, SO, Cl. [86.5°].

Amide. --C, H, Br, SO, NH. (c. 225°].

Anhydride (C.H.Br., SO.).O. Obtained by the action of funing suphuric acid upon to be a superior of the superi tri-bromo benzene (1:2:1). Amorphous powder. Almost insol. alcohol, other, and benzene. By boiling aqueous alkalie it is converted into salts of the acid; only very slowly attacked by boiling water. PCl, converts it into the chloride (v. supra) (Roscuberg, B. 19, 654).

Tri-bromo-benzen sulphonic acid

C,H2Br,SO,H [1:2:3:5]. From di-brominated C., H., Br., SO., H. (123:3). From di-brominated amido-benzene p-sulphonic acid (Lenz, B. 8, 1067; A. 181, 29). — NH, A'. — KA'. — CaA', 2!aq. S. '39 at 20'. — BaA', 3aq. S. (of BaA', 2) '021 at 18°. — 1'bA', 3!aq. S. '056 at 21°. Chloride C, H.Br., SO., Cl. [127°]. Amids C, H.Br., S. NH, [210°].

Tri-brome-benzene sulphonic acid C H.Br. SO. H. [123:35]. Kron. silver. 1. 2. 41.

C. H. Br. SO. H [17:2:3:5]. From silver (1, 2, 4). di-bromo-benzene sulphonate and Br (Goslich, A. 186, 154). Is perhaps identical with the preceding.—BaA', 3 aq.

Chloride [121]. Amids [152].

Tri brome-banzane sulphonic acid C.H.Br.SO.H [1:3:4:5]. From nitrated (1, 8, 5). di-bromo-benzene sulphonio acid (Lenz, A.

Chloride C.H.Br.SO.Cl. [86°].

Amide C.H.Br.SO.NH. Blackens at 225°.

Tri-brame-bensene sulphonic soid C.H.Br.SO.H. Possibly identical with the preseding. From nitrated (1, 4, 2)-di-hromo benzene sulphonio acid (Borns, A. 187, 364).—
KA' 8]aq.—BaA', 2aq.

Amide [above 220°].

Tri-bromo-benzene enlphonic acid C.H.Br.SO.H. From eilver bromo benzeno osulphonate and Br (Bahlmann, A. 181, 207).

Chloride C.H.Br.SO.Cl. [56°]. Amide C.H.Br.SO.NH. [2020 Tri-brome benzene enlphonio acid C.H.Br.SO.H. Formed at the same time as the

preceding (B.). Chloride C.H.Br,SO,Cl. [72°].

Amide Call_Br.SO.NH. [187°]. Nine tri-bromo-benzeno sulphonic acide are here described, but only six are indicated by theory.

Tri-bromo benzene di-sulphonic acid C₂HBr₃(SO₃II)₂. From benzone m-di-sulphonio acid by nitration, reduction, bromination to C₂HBr₂(NII.)(SO₃H)₂ and diazotisation (Heinzelmann, A. 183, 183). - Salt: K₂A".

Tetra-bromo-bonzene-sulphonic acid

C.HBr.SO, H 5aq [1:2:3:5.6].

Formation. - 1. From C.HBr. (NH.)SO.11 [1:3:5:4:6] by diszo- reaction (Beckurts, A. 181, 216; Langlurth, A. 191, 199; Knuth, A. 186, 229; Reincke, A. 186, 282).—2. From C. H. Br. and H.SO, (Bässmann, A. 191, 223). Needles. Not hygroscopic but v. sol. alcohol and water.

Reaction. - With conc. IIBr at 150° gives H.SO, and (1,2,3,5) tetra-bromo benzene [98-5°].

Salte, - KA', S. 63 at 6°, - BaA', 1. S. 37 at 12° (L.); 16 at 15° (Bä.), - BaA', 1. a. - CaA', 28aq. - S. (ol CaA',) 54 at 3°; 66 at 19° (Bä.), - NH, A', - PhA', PhO 3aq. - PhA', 1. paq. S. (ol PhA',) 89 at 11° (Bä.), - PhA', 4aq. Chloride C. HBr. SO.Cl. [96°].

Amide C. HBr. SO, NII. Minute needles. Not melted at 300°.

Tetra-bromo-benzene eulphonio acid

C.HBr.(SO.H) [1:2:3:4:5]. [169°]. From (1,2,3,5) tri bromo benzene sulphonio acid by nitration, reduction, and diszotisation (Lenz, A. 181, 23). Also in the same way from (1,2,1,5). tri bromo benzene sulphonic acid (Spiegelberg, A. 197, 292). Lamine (containing 2aq).

A, 197, 292). Laminie (containing zaq).
Salts.—KA'aq. S. (ol KA') '194 at 112.—
NH₄A'. S. 95 at 112.—BaA', aq. S. (ol BaA') '0204 at 10.52.—CaA', 3aq. S. (of CaA') '159 at 112.—PbA', 3aq. S. (of PbA') '0181 at 117.—AgA', aq. S. (ol AgA') '146 at 117.—AgA' aq. Chlorida C,HBr,SO.Cl. [1202].

Amide C.HBr SO, NII. Turns brown at 210°. Penta bromo benzene sulphonio acid

C.Br.SO.H. S. 518 at 10°. From either prous cya tetra-bromo-amido-benzene sulphonic acid by chloride (diazo-reaction (Beckurts, A. 181, 226; Hein 18, 1196). zelmann a. Spiegelberg, A. 197, 306; Langfurth,

A. 191, 205). Needles or plutes (containing raq).

Salte.—NH,A'. - KA'nq. S. (of KA') 116 at
10.5°.—CaA',4aq. S. (of CaA_) .78 at 14°.—
BaA',2aq. S. (of BaA',) .0088 at 13°.—BaA', 1½aq. AgA'llaq.

Ohloride C.Br.SO.Cl. [1542].

Amide C.Br.SO.NH. Decomposes at 250°.

BROMO-BENZIDINE v. BROMO-DI-AMIDO.

o-BROMO-BENZOIG ACID C.H.Bro, i.e. C.H.Br.CO.H [1:2]. Mol. w. 201. (Z.; J. a. W.); [150°] (R.). [148°L

Formation .- 1 From o-amido-henzoio acid (Richter, B. 4, 465).—2. By heating C₄H₄Br(NO₂) [1:3] with KCy at 180° (R.).—3. By oxidation of o-bromo-tolueno (Zincke, B. 7, 1502; Rahlis, A. 198, 99), o-bromo-diphenyl (Schultz, A. 207, 353), or o-bromo-benzyl alcohol (Jackeon a. White, Am. 2, 316).

Properties .- Long needles (from water); m. sol. water, v. sol. alcohol, and ether; elightly volatile with steam.

Salts. -- BaA', -- BaA', 211 vEt -- CaA', Saq. -- KA'2nq. -- ZnA', -- PbA', HOEt. -- CuA', aq. [257°], -- IlOCuA'.

Methyl ether Mod. (247°). Ethyl ether EtA'. (255°).
Anilide C.H.Br.CONPhH. [142°].

m Bromo-benzoio acid Call BrCO II [1:8]. [155°]. (280°).

Formation. - 1. From AgOBz and Br (Peligot, A. 28, 246; Angerstein, A. 158, 2).— 2. From benzoic acid and Br in the sunchine (Herzog, N. Br. Arch. 23, 16) or by heating them with water at 100° or 160° (Reinecko, Z. 1865, 116; 1869, 100; Hubner, A. 143, 233; 149, 131). 3. By heating benzamido with Br and water (Reinceke, Z. 1866, 367; Friedburg, .1. 158, 26). - 4. From m-amido-benzoic acid by diazo- reaction (Griess, A. 117, 25; Hübner, A. 222, 100). 5. From C, II, Br(NO) [1:4], KCy, and alcohol at 200° (Richter, B. 4, 464). 6. By oxidation of m-bromo tolueno (Wroblewsky, Z. [2] 5, 332; A. 168, 156) or m-bromobenzyl alcoliol (Jackson, Am. 1, 93). - 7. By the action of a hot solution of cuprous cyanide in petassium cyanide upon m bromo diazobenzene chloride (from m bromaniline), and saponification of the erndo nitrile (Sandineyer, B. 48, 1496). 8. From m-di-bromo-benzeno, CICO by, av I Na (Wurster, A. 176, 119).

Properties - Cry talling; very slightly volatile with steam; sl. sol. water, v. sol. alcohol.

Reactions. 1. Potash-fusion gives m. and a little o- oxy-benzoic acid (Richter, Z. 1860, 157; Barth, A. 159, 236).-2. The K salt fused with sodium formate gives isophthalio acid (V. Meyer a. Ador, A. 159, 15).

Methyl ether MeA'. [32], Ethyl ether MeA'. [32], Ethyl ether PhA'. [65],

Chloride Call, Br.COCl. (2397).

Amide C.H.Br.CONH... [150°]. Nitrile C.H.Br.CN. [28°]. (225°). From the amide (Engler, B. 1, 708). Formed also by the action of a hot solution of potassinm cuprous cyanide upon m. bromo dinzo benzene cirloride (from m brow miline) (Sandmeyer, B.

p. Bromo . benzoic acid C. H.Br.CO.H [1:4]. [251°]. Formation. - 1. By oxidation of p. bromo toluene (Habner, A. 143, 247; Jackson Rolle, Am. 9, 84), p-bromo ethyl-benzeno (Fittig a. König, A. 144, 283), or p-bromo-banzyl bromide (Jackson, Am. 1,93). - 2. From p. bromoaniline vid p-bromo thio carbinuide, the latter when heated with copper at 190° giving p. bromo-benzonitrile (Woith a. Landolt, B. 8, 718).

Properties. Small needles (from ether) or

plates (from water). V. al. sol. water, v. sol.

alcohol and ether. -AgA'.

Ethyl ether Eth' (236° nnoor.) at 713 mm. Formed, together with p-bromo-benzyl-alcohol, by boiling p-bromo-benzyl bromide with alco-holic KOII (Elbs, J. pr. [2] 34, 341). Phenyl ether PhA'. [117]. Scales.

Phenyl ether van. [140°]. Pyrogallyl ether C. H. A'r. [140°]. Chloride. [30°]. (246°). Amide C.H.Br.CONH, [1 [186°]

Anilide C. H.Br.CONPhill. [1977]. Plates. On nitration it gives a dinitro dorivativo. [214°]. (Raveill, A. 222, 178).

Anhydride (C, II, BrCO),O. [213°]. From brome-benzoyl chloride and sodium p-bromebenzoate (J. a. R.). Minuto oblong rectangular plates (from CHCl.). Insol. water. Converted

by hot HOEt into the ethyl ether.

Dl-bromo bonzoic acid CallaBraCOaH [1:2:3] From (a)-bromo-amido-benzoie acid [1:2:3] by diazo-reaction (Hübner, A. 222, 105). From (1, 2, 3) di-bromo-toluene (Nevile a. Winther, B. 13, 965). Silky needles (from water). V. sol. hot water. Heated with H.SO, (3 vols.) and II.O (1 vol.) at 225° it gives a di bromobenzene [218.5°

Salts. BaA', 11aq. S. (of BaA',) 111 at 16°. SrA', 4aq. A'CuOH, - KA', raq.

This acid is perhaps identical with the dibrome-benzoic acid [148°] prepared from o-nitrobenzoic acid, Br. and water at 200° (Claus a. Lnde, B. 14, 1170).

Di-bromo-beazoio acid C.H.Br.CO.H [1:1:3] [153°]. From (β) brome amide benzoic neid [1:4:3] by passing nitrous acid gas into a mixturo of the acid, IIBr, glacial acetic acid and alcohol (Hübner, A. 222, 108). From nitro-pdi-bromo-benzene and alcoholic KCy (Richter, B. 7, 1146). From (1, 4, 3) di-bromo-toluene by exidation (Novile n. Winther, B. 13, 963). Long silky needles (from water). Volatile with steam. Heated with H2SO, (3 vols.) at a water (I vol.) at 225" gives p-di-bromo benzene [89].

Salts. - BaA', 1 aq. - SrA', taq. - ZuA', CuA' 3 3 aq. KA' w1.

This acid is probably identical with the dibromo benzoie neid [153"] from o-nitro benzoie acid, Br. and water at 200 ' (Claus a. Lade, B. 14, 1170).

Di-brome benzoic acid C.H.Br.CO.H [1:3:2]?. [150°-167°]. By oxidising di-bromo-toluene from di-bromo-m-toluidine, [35°], (Sievile a. Winther, C. J. 37, 441).

Di-brome benzoic acid O.H.Br.CO.H [1:3:4]. [169]. By oxidising di bromo toluene from dibromo m toluidine [76°] (Nevile n. Winther, C. J. 37, 111).

Di-bromo-beazoic acid C.H.Br.CO.H [1:3:5]

[207 · 210 ·] (N. a. W.); [209 ·] (R.); [214 ·] (II.).

Fermation. - 1. By oxidation of di-fromotolnene ,39 · from di-fromo-p-tolnidino [73 ·]

(Nevite a. Winther, C. J. 37, 437), -2. From m-di-bromo-benzene by nitration and treatment with alcoholic KCy at 250° (V. v. Richtor, B. 8, 1423).-3. From di-brominated p-amido-benzoic acid (Beilstein a. Geitner, A. 139, 4). -4. From C.H.Br(NH.)CO.H. aqueous HBr (S.G. 148), glacial acetic acid and N.O. at 15° (Hesemann a. Köhler, A. 222, 171).

I'mperties. - l'lates (from alcohol) Noedles (from other solvents).

Salts.—BaA', 4aq.—CaA', 5aq.—CdA'; 4a CaA', 6aq.—NaA' aq.—CdA'; 4aq. Di-bromo benzolo acid C₄H,Br,CO₂H [1:8:x

[223°-227°]. Formed by heating benzoic aci with Br and water at 220° (Angoratein, A 158, 10). Needles.—BaA', 2aq.
Di-brono-benzoic acid C,H,Br,CO,H [1:2:4]

[229°-230°] (B.); [233°] (N. a. W.).

Formation.—1. C.H.Br(NH.)CO.H [4:3:1] i. treated with other and N.O. and the diazo-derivative treated with HBr (Burghard a. Beutnagel A. 222, 181). - 2. By oxidising the corresponding di-bromo-tolueno from brominated p-toluidine (Nevile a. Winther, C. J. 37, 439). -3. A product of the action of Br on p-nitro-benzoic, or (1, 2, 4)di-nitro-benzoic, seid (Halberstadt, B. 14, 908, 2215).

Properties .- Colourless needles (from water),

tables (from alcohol).

Salts.-BaA', 4aq. -SrA', 4aq.-HOCuA',-AgA'.--KA' anq.

Ethyl ether EtA'. [38°

Amide Call Br. CONII... [151°-152°].

Tri-bromo benzolo * acid Call Br. (CO.H). [235°]. From m-bromo-benzoic acid and Br (Reinecke, Z. [2] 5, 110). Tutts of slender silky needles; v, sl. sol, water. -NH, A'.—CaA', 5aq.

Tri-brome-benzoic acid C, II. Br, CO, H.

[1:3:5:6]. [187°]. From tri-brominated m-amidobenzoic acid (Vollbrocht, B. 10, 1708). Needles. BaA', 5 laq.

Tri-bromo-benzoic acid C.H.Br.CO.H. [195°]. From (1, 2, 4)-di-bromo-benzoic acid [229°] by nitration, reduction, and diazotisation (Smith, B. 10, 1706). Needles (from nlcohol).-BaA', 5aq. Tri-brome-benzoic acid C.H.Br. CO.H. [1780] From (1, 4, 5)-bromo-amido benzoic acid (Lawrio,

B. 10, 1705). Needles.—BaA', 3mj. Peata bromo-benzoic acid C_sBr₃CO₂H. (235°) From tri-bromo-benzoic acid and Br at 2000 (Reinecke, Z. [2] 5, 110). Thin plates or flat needles (from alcohol); thick needles (from benz-

ene); v. sl. sol. water.—NH A'.—CaA', 6aq. Nitrile C, Br, CN. [above 300]. Obtained by brominating benzonitrilo (Merz a. Weith, B. 16, 2892)

D-BROMO BENZOIC ALDEHYDE

Call Br.CHO [1:2]. A heavy oil; formed by boiling o-bromo-benzyl bromide with aqueous lead nitrato (Jackson a. White, Am. 3, 33; P. Am. A. 15, 269).

m-Bromo-benzoic aldehyde C. H. Br.CHO [1:3], Oil (J. a. W.).

p-Bromo benzoic aldehyde Call, Br. Cilo [1:4],] (Jackson a. White, B. 11, 57).

BROMO BENZOIC SULPHINIDE v. BROMO SULPHO-BENZOIC ACID.

BROMO - BENZOT TENONE C, II, Bro i.e. Cull .CO.C. 11 Br. [81.53]. 'From benzoic acid, bromo benzene and P.O. at 190 (Kollarits a. Merz₅ B. 6, 547).

BROMO-BENZOYL CHLORIDE v. Chloride of BROMO-BENZOIO ACID.

BROMO-DI-BENZYL v. BROMO-DI-PHENYL-ETHANE.

o-BROMO-BENZYL ALCOHOL C,H,BrO i.a. C.H.Br.CH., OH. [80°]. Prepared by digesting o bromo benzyl acetato with aqueona ammonia at 160°. Crystallises in white needles. Sol. hot water, ligroin, alcohol, other, benzol, and CS.

Volatile with steam (Jackson a. White, Am. 2, 916; R. 13, 1218).

m-Bromo bensyl alcohol C.H.Br.CH.OH. From m.bromo-benzyl bromide and water at 130° (J. a. W.).

p-Bromo-benzylaicohol C. H. BrCH.OH. [77°]. Prepared by bolling p-bromo benzyl bromide with water for 3 days. Long elastic transparent necdles. Sol. alcohol, ether, benzeno, and CS. (Jackson a. Lowery, Am. 3, 246; B. 10, 1209).
o-BROMO-BENZYL-AMINE C. H. Br. CH. NII.

Prepared by acting on o-bromo-benzyl bromide with alcoholic ammonia at 100° for 2 hours (Jackson a. White, Am. 2, 318). Colourless oil; sol. ether. Salts.—B'HCl: [208°]; needles.—B'_4H_2PtCl_6.—B'_HI_CO_3. [95°].

p-Bromo · benzyl-amino Call Br CH NH .. From p-bromo-benzyl bromide and cold alcoholio NH, (Jackson a. Lowery, Am. 3, 251). Oil; volatilo with steam. -B'HCl [160]. B',H,PtCl, -B'2H2CO2 [131°-133°]; small prisms.

Di-o-bromo-di-benzyl-amine

(C_eH₄Br.CH₂),NH. [36²]. From c-bromo-henzyl bromide and alcoholic NH, at 100². Trimetric crystals; insol. water (Jackson a. White, Am. 2, 318; B. 13, 1219). BHCl[166]. - B',H,PtCl,

Di-p-bromo-di-benzyl-amine (C,H,Br.CH₂),NH. [50°],—B'HCl [183°].—B'₂H,PtCl₆ (J. a. W.).

Tri-o-bromo-tri-bonzyl amine(C.H.Br.CII.), N. [122]. From o-bromo-benzyl bromide and alcoholic NH, at 100° (J. a. W.). Small prisms. -B',H,PtCl.

Tri-p-bromo-tri-benzyl-amine(Call, Br.CII.), N. Crystals (from ligroin) [92]; (from ether) [78]. Slender needles; the hydrochloride could not be obtained. -B'HBr. [270]; insol. water.-B'2H.PtCl, (Jackson a. Lowery, Am. 3, 252).

o.BROMO.BENZYL BROMIDE

C₀H₄Br.CH₂Br {1:2}, {30°}, (250°-260°), Dibromo-toluene. Prepared by brominating obromo-toluene (Jackson, Am. 1, 93; 2, 315; B. 13, 1218). Very pungent; volatile with steam. Converted in ethereal solution by Na into anthracene, phenauthrene, s-di-phenyl-ethane, and other products.

m-Bromo-benzyl bromide C.H.Br.CH.Br [1:3]. [41°]. From m-bronio tolucic and Br (Jackson, Am. I, 93; B. 9, 932). Pungent plates; slightly volatile with steam; very volatile with othervapour.

p-Brome-benzyl bromide C_oH₄Br.CH₄Br [1:4]. [62⁵]. Formed by brominating p-brome-toluene or benzyl bromide (Jackson, Am. 1,93; Schramm, B. 17, 2922; 18, 350). Needles (from alcohol); volatile with steam; very pungent. Alcoholic KOH forms p-bromo-benzyl alcohol and pbromo-benzoic ether (Elbs, J. pr. [2] 34, 310).

BROMO-BENZYL SYANIDE v. Nitrile of BROMO-PRENTL-ACETIC ACID.

BROMO-BENZYLIDENE. PHTHALIMIDINE C13H10ONBr. Phthalimidyl-bromo-benzyl. [210°]. Glistening needles. Formed by the action of bromine upon deoxybenzoin-carboxylamide bromine upon deoxybenzoin-carboxylamide C.H. (CO.NH.) CO.CH. C.H. dissolved in chlorolorm (Gabriel, B. 18, 2135).

TETRA-BROMO-BENZYLIDENE-DI-TOLYL-**DIAMINE** (PhCH)₂(NC₀H₂Br₂Me)₂. [160°-165°]. From benzylidene-di-p-tolyl-diamine and Br (Maszara, G. 10, 370).

p.BROWO.BENZYL MERCAPTAN C.H.Br. CH.SH. [25°]? From p.bromo-benzyl bromide and alcoholic KHS (Jackson a. Harts-horn, Am. 5, 208). Crystalline mass; insol. water and glacial HOAo; sol. alcohol, ether, and benzene. - Ilg(SC.H.Br)2: sol. hot alcohol. DI-BROMO BENZYL-PHENOL C. 11, Br., OIL.

[1757]. From henzyl-phenol in CS, and Br (l'aterno a. Fileti, G. 3, 254). Amorphous.

BROMO-0?-BENZYL-PHENOL SULPHONIC ACID C.H., C.11, (O11) BrSO, H. Sa t. KA'. From Br and C.H. C.H.(OH)SO, K (Renne, C.J. 49, 409).

Bromo-p-benzyl-pheuol sulphonic acid

C,H,C,H (OH)Br.SO,H [1:4:x:2].

Salt. - KA' (Rennic, C. J. 41, 35), p-BROMO-BENZYL SULPHIDE

(C,H,BrCH,),S. [59°]. Prepared by boiling p-bromo benzyl bromide with alcoholic Na₂S. Large thin plates. Aromatic edour. Insol. water. Sol. ether, benzene, and CS. (Jackson a. Hartshorn, .1m. 5, 267).

p Bromo - renzyl disulphido (C.H.BrCH.)282 [88]. Prepared by exposing the mercaptan to nir, and also by acting on p-bromo-benzyl bromido with alcoholic Na.S. Needles. Insol. water; sol. ether, benzene, and CS, (Jackson n. Hartshorn, Am. 5, 269).

P-BROMO-BENZYL SULPHOCYANIDE

C,H,Br.SCN, [25°]. From p-bromo-benzyl bromide and potassium sulphocyanide. The o-isomeride is an oil (Jackson a. Lowery, B. 10, 1209; Am. 3, 250).

DI-p-BROMO DI-BENZYL SULPHONE

(C,11,Br.CH_),SO,.. [1897]. From the sulphide and C(O, in 110Ac. Needles (Jackson a. Hartshorn, Am. 5, 269).

BROMO - BENZYL SULPHONIC ACID v.

BROMO-TOLUENE BULPHONIC ACID.

DI.BROMO-BETORGING Br.Me.(OII),[155°]. From tetra-become betorein and III. Also by boiling a mixture of hetorein (3 pts.), bromine (8 pts.) and Ch. (100 vols.). The product is reerystallised trois light petrolemn (Stenhouse a. Groves, C. J. 37, (0)). Long needles.

Totra-bromo-betorcin C.Br.Me.(OBr),. [101]. From bromine (5 pts.), water (100 vols.) and betorein (I pt.). Dissolved in water (50 pts.). Crystallised from light petrolenm (S. a. G.).

Largo colourless prisms. V. sol. ether, benz-

ene and CS₂, less so in petrolenm.

BROMO - BRASILIN C₁₀II₁₃BrO₂. Obtained by brominating acely brasilin and saponifying the product. Glistening red plates. water. Dissolves in KOII with a red colour.

C, H, Br(OAc), O. [201"] (Buchka a. Erck, B. 17, 685; 18, 1110). Tetra-acetyl derivative

Tri-bromo brasilin Ciall, Br.O.

Tetra-acetyl derivative

C₁₈11,Br₂(OAc₁O, [147°], Small white needles; very oxidisable (B.a. F.).

Tetra-bromo-brasilin C₁₈11₁₈Br₄O₂, Slender

red needles. Dissolves in alkalis with a violet colour. Obtained by bromination of brasilin. Tetra-acetyl derivative

C₁₆H₂Br₄(OAc)₁O. [222] (B.a. E.). BROMO-BUTANE v. BUTYL BROMIDE. w.a.Di.bromo.butane C,H,Br, i.e.

CH Br.CHBr.CH, CH, Butylene bronnide. (166"). S.(1. 29 I 820. Formed by the action of Br on nbutylene or n-butyl bromide (Wurtz, A. 152, 23, Linnemann, A. 161, 199; Grabowsky a. Sayizeff,

A. 179, 882). Na forms CH.; CH.OH.CH. ab.Di.brome.butane CH., CHBr.OHBr.CH. (1582). S.G. 21-82. From CH., CH:CH.CH. and Br. Converted by PhO and excess or water at 150° into methyl cthyl ketone (Wurtz, A. I44, Converted by PhO and excess of water at 236; Eltekoff, J. R. 10, 219).

Di-bromo-isobutauo (CH3)2CBr.CH2Br. Isobutylene bromids. (149°). S.G. 14 1-8; 15 1-7434; 25 1-7308. M. M. 11-890 at 14-75 (Perkin). From isobutylene and Br (Linnomann, A. 162, 36). By heating with excess of water at 150° it is converted into isobntyric aldeliyde; if TbO is also present some di-oxyisobutane is also formed (Elitekoff, J. R. 10, 214).

Di - bromo - butano C₄H₈Br₂. (155°-162°). Formed by brominating butane (Carius, A. 126,

Trl-bromo-isobutane C.H.Br. i.e. (CILBr),CBr.CII, (173°-183°) at 235 mm. S.G. 17 2-15. Prone isobutylene and Br (Norton a. Williams, Am. 9, 88).

Trl bromo-isobutane C,II,Br, i.e. (CH₃),CBr,CHBr. (155°-161°) at Frem (CH₃),C:CHBr and Br (N. a. W.). (155°-161°) at 235 min.

Trl - bromo - butane C.H.,Br. (2087-2159). From bromo-butylene (82 - 92') (Caventou, A.

Tetra-bromo hutana

C, II, Br, i.e. CII_Br. CHBr. CHBr. CII_Br. Butineneor pyrrolylene-tetra-bromide. [H9°]. Formed by combination of bromine with the buttinene from di-methyl-pyrrolidino or from crythrite. From orythrite, vinyl-othylene, or gasoils (Caventon, A. 127, 95; B. 6, 70; Henninger, B. 6, 70; Grinnaux a. Cloez, C. R. 104, 1446; Bl. [2] 48, 31). From the gas obtained by passing acetylene mixed with othylene through a red-hot tube (Prunier, El. [2] 20, 72). On distillation it is partially converted into the following body. Colourless needles (from alcohol). Insel. cold petroleum ether (Ciamician a. Magnaghi, B. 19,

Tetra brome butano CH CBr CBr CH, (?) [40°]. Erythrens isobremide. Erythrite tetrabromhydrin. Formed, together with the isomorida [1193], by combining butinene from erythrite with bromine. Large colourless trimetric prious or tables, $a:\hat{b}:c = 9776:1:1:682$ (Chamician a. Magnaghi, B. 19, 569). V. sol. other, alcohol, and petroleum-ether. Alcoholic KOH converts both this and the preceding into the same di-brono-butinene C₄H₄Pr₂, which rapidly polymerises. The latter absoros Br. forming C₁H₂Br₄ [67] and C₄H₄Br₄ 170]

Totra-brome-butano C.H.Br., From Br and di-brome-butylene (140°-180°) from fusel oil nit butylene (Caventou, A. 127, 93). Crystalline; decomposes at 200°.

Totra-bromo-bntane CH, CH, CBr, CHE, From hutineno prepared from methyl chyl Letone by snecessive treatment with PCl, and alcoholio KOH. Sublimes at 105° (Bruylants,

Tetra-bromo-lsobntano $C_{i}\Pi_{a}Br_{i}$ From di-brome-isobutyleno (155°) (Norton a. Williams, Am. 9, 87).

Hexa-bromo-butano C,H,Br,. [109°]. tained by brominating isobutyl bromide at 170°. The yield is 90 p.c. of the theoretical (Merz a. Weith, B. 11, 2245).

Here brome butters C.H.Br. Le. Br. CHBr. CHBr. CH.Br. S.G. 11 2-9. Forme together with the following, by heating crythric tetrabromhydrin with Br at 180° (Coleon, L [2] 48, 52). Liquid; v. sol. ether, sl. sc alcohol. Dilute KOH at 130° converts it in potassium erythrate.

Hexa-bromo-butane C.H.Br. [170°]. S.C 3.4. Formed in small quantity as above (C. Pearly scales, sl. sol. ether and alcohol. Fumin HNO, forms an oil C₁H₃(NO₂)Br₂(NO₃)₂, S.G. 1.81

DI-BROMO BUTYL ALCOHOL C,H,Br,O, i.e Cll, CHBr. CHBr. Cll, OH. Oil. From buteny alcohol CH, CH: CH. CH, OH and Br (Lieben a Zeisel, M. 1, 828). Boiling water forms tri-oxy butane.

ωα.DI-BROMO.n-BUTYL-BENZENE

C .o II .Br 2 i.o. C H3.CII .CH2.CHBr.CH.Br. From plienyl-butylene and Br (Aronheim, A. 171, 229). Reactions .- 1. Red-hot soda-limo forms naphthaloue.—2. HNO, forme a little bromephenyl-propionic acid.

 $\beta\gamma$ -Di-bromo-n-butyl-benzene C.II. CHBr.CHBr.CH, CH. [67°]. From butenyl-benzene and Br (Perkin, C. J. 32, 668). γγ-Di-bromo-n-butyl benzene

C. I. CBr., CH., CH., CH., (?). From butyl-benzene and Br in sunlight. Unstable oil (Schramm, B. 18, 1276).

Di-bromo n-bntyl-benzene C10H12Br4. [70°]. From n-butyl-benzene and Br first in sunlight, thon heated in the dark (Radziszewski, B. 9,

άβ-Dl-bromo-ieobutyl-benzene

C.11, CHBr. CBrMe. From phenyl-isobutylene and Br (Perkin, C. J. 35, 138). Tri-bromo-isobutyl benzene C, H, Br, [64°].

From the preceding by successive treatment with alcoholic KOH and Br (P.). BROMO-BUTYLENE C.H.Br i.e.

(CH3), C:CHBr. (91°). Isocrotyl bromide. From (Butlerow, Z. 1870, 524). Alcoholic KOH at 170° forms (CH₂)C:CH.OEt. Moist Ag₂O at 100° gives isohutyue acid. NH₂ has no action.

Brome-butylene C₄H₄Br. (82° 92°). From

fusel oil butyleno by successive treatment with Br and alcoholic KOII (Caventou, A. 127, 93).

Bromo-butylene C₄II₄Br. (87° i.V.). From

di-bromo-methyl-ethyl-acetic acid and aqueous Na CO, (Jaffé, A. 135, 300; Pagenstechor, A. 195, 126).

Bromo-butylene C.H.Br. (97°). Formed by hoiling the di-bramide of angelic acid with water (Jaffé, A. 135, 300).

Di bromo iso butylene C₁II Br. (155°). From tri bromo iso butane (155° 161°) at 235 mm. (Norton a. Williams, Am. 9, 87).

235 mm. (Norton a. Williams, Am. 9, 87).

Di brome, butyleue (, H. Br. (140?-150°).

From tri-brome-butane (208°-215°). Forms a crystalline di-bromide C.H. Br., which decomposes without melting at 200° (Caventou, A. 127, 93).

Dl -brome -butylene C.H. Br. (148°-158°).

From crotonylene and Br (Caventou, A. 127, 349).

D': brome-butylene C.H. Br. From tatra-

Li - bromo - bntylene C.H.Br... From tetrabrome-butane [1192] (from erythrite) and alooholic KOII. Rapidly polymerises. Combines readily with bromine (Grimaux a. Cloez, Bl. [2] 48, 3I).

Di bromo butylene C.H.Br. i.d. CH,:CH.CHBr.CH,Br. (1903-200°). From the edis deposited by compressed gas, by adding less than the calculated quantity of Br and fractionally distilling. Combines readily with Br, forming C,H,Br, [119°] (G.a. C.).

Hexabromobutylene C,Br,H,... [53°]. Prepared by further bromination of hexabrom-

sobutane (Merz a. Weith, B. 11, 2210).

a.BROMO. n. BUTYRIC ACID C,H.BrO, i.e. OH, CH, CHBr, CO H. (211°-217°). (110°) at 8 mm. S.G. 12 1-54. S. 7.

Formation .- 1. By heating butyric acid with Br at 140°; or by the action of Br on silver butyrate (Friedel a. Machaca, A. 120, 279; Suppl. 2, 70; Gorup-Besanez a. Klinksieck, .t. 118, 218; Naumanu, A. 119, 115; Ley, J. R. 9, 129; Urech, A. 165, 93; Topoleff, A. 171, 249). - 2. From crotonic acid and HBr .-8. From the bromide and water (Kaschirski, C. C. 1881, 278).

Properties .- Oil; mixes with alcohol and NEt, whether dry or in aqueous solution, forms a oxy-butyric acid (Duvillier, Bl. [2]

 Matthy a construction and a construction of the const an other Call (OEt), (252) (Krestownikoff, A. 208, 318).

Bromide C.H. BrO.Br. (173°), From

butyryl bromide and Br.

B-Bromo-butyric acid CII, CHBr.CH., CO.H. Formed in small quantity in the preparation of the a- acid from crotonic acid and HBr (Hemilian, A. 174, 325).

y-Bromo butyric acid CH Br.CH..CH .CO .H. [33°]. From batyro-lactone and HBr (Henry, C. R. 102, 368). Tables or plates; sl. sol. Tables or plates; st. sol. water, v. sol cther.

Methyl ether McA', (187°), S.G. 2 1·45, Ethyl ether EtA', (197°), S.G. 2 1·36,

a - Brome - isobutyrio acid (CII,) CBr.CO.II. [48°]. (199°). S.G. (2015). From isobutyria acid and Br (Markownikoff, A. 153, 229; Hell a. Waldauer, B. 10, 448). Tables. Boiling water converts it into a oxy isolutyric acid, KHS acts similarly (Loven, J. pr. 2/33, 105); boiling baryta water forms also Cll CMc.CO.II (Engelhorn, A. 200, 68).

Ethylether EtA'. (161° cor.). S.G. 21.13. Oil; smelling of raspberries and peppermint (Markewnikoff, A. 182, 336; Hell a. Wittekind, B. 7, 320; Lovén, J. pr. [2] 33, 106).

Bromide (CH,) CBr.COBr. (163°).

8-Bremo-Isobutyric soid CH, Br.CHMe.CO, II, [22°]. From a methyl acrylic acid and cold cone. HBr. Crystals (from CS.). Boiling cone. HBr. Crystals (from CS.). Boiling alkalis convert it into a methyl acrylic acid and a little propyleno (Fittig a. Engelhorn, A.

aa-Di bromo butyrio acid CH2, CH2, CBr2, CO211. (140) at 3 mm. S. 3. S.G. 1.96. From bromo-butyric acid and bromine (Schneider, J. 1861, 458; Michael a. Morton, Am. 2, 15; Otto a. Fromme, A. 239, 275). Thick oil. Water or baryta-water at 275). Thick oil. Water or baryta-water at 120° forms a-bromo-crotonic acid (Erlenmeyer . Müller, B. 15, 49). Converted by reduced silver into di cthyl maleio or zeronio acid CO.H.CEt:CEt.CO.H, butyric acid being formed.

aff Di-bromo-butyric soid CH, CHBr.CHBr.CO,H. [87°].

Preparation .- From Br and crotonic or inc. crotonio scid dissolved in CS, (Körner, A, 187, 234; Michael a. Norton, Am. 2, 12; B. 14, 1202; C. Kolbe, J. pr. 133, 386).

Properties. - Large transparent prisms (from

CS.); sol. alcohol, ether, and hot water.

Reactions. -1. Boiled with water or Na.CO. it gives \$-brome propylene, brome-oxy-butyric acid, di-oxy-batyrie acid, and bromo-crotonie neid. Water gives chiefly bromo-exy-butyris acid; Na CO, gives more bromo propylene, but no propionic aldehyde. 2. Wermed with a solution of NaOII it gives bromo-crotonio acid.

Di-bromo-iso-butyrio acid

CH_Br.CBrMo.CO.H. [48°].

Preparation .- By adding Br to methacrylic acid dissolved in CS, (C. Kolbe, J. pr. (2) 25, 373). Long prisms (from CS₃). Boiling with water or Na₂CO₃ produces CO₃, acctone, some propionic aldehyde, a very little bromo methacrylic neid 63° jand bromo-oxy-iso-butyric acid (q. v.). Warmed with solution of NaOH it gives brome methacrylic acid and HBr.

Tri-breme-butyrio acid C. II Br.O. i.e. CH CBr CHBr CO H ? [114°]. From B-bromocrotonic acid in CS, and Br (Michael a. Norton, Am. 2, 14). Rhombie plates; sol. alcohol and

hot water; sublimes readily.

Tri-brome-butyric acid C. II Br.O. i.e. CH, CHBr.CBr.CO.H? [1110]. From a bromoerotonic acid and Br. V. sol, water and alcohol (M. a. N.).

Tri-bromo-isebutyrle seid C, II, Br, O2. From bromo a methyl-nerglic acid and Br (Cahours, A. Suppl. 2, 349). Prisms,

Tetra-bremo-butyric acid C,II,Br,O2. [1150 From nucobranie acid and Br (Limpricht, A. 165, 293). Monoelinie tables; al. sol. water.

Tetra-bromo isobutyric acid C.H.Br.O From d, brown-a-methyl-acrylic acid and Br(O.)

BROMO-ISO-BUTYRIC para-ALDEHYDE ((CH₃), CBr.CHO)_n, [129°], When the product of the action of NH_a on iso-butyric aldehydo (q.v.) is distilled a product C, II, N is got. This must be combined with bromine, and the compound, (CH2)2CH.CH: N.CHBr.CBr(CH2)2, after keeping for 3 months, is decomposed by water (Lipp, A. 211, 353). Needles (from alcohol). lusol, water, acids or alkalis, v. sol. ether, m. sol. alcohol. Does not reduce ammoniacal AgNO.

BROMO CATECHOL v. BROMO PYROGATEORIN.

BROMO-CAFFEINE C.11, BrN. O. [206°]. Caffeine combines with Br forming the orangered bromide C.H., N.O.Br., which is decomposed at 150° into HBr and bromo cafferne (Fischer, B. 14, 639; Schultzen, Z. 1867, 614; Maly a. An freasch, M. 3, 85). Cry tals, sl. sol. cold water, v. sol. NH, Aq. Reduced by zinc-dust to caffeine. Alcoholio KOH forms ethory-caffeine.

BROMO-CAMPHOR v. CAMPHOR.

BROMO-CAMPHORIO ACID v. CAMPHORIS

BROMO-CAPRIC ACID v. Bnomo-DECO10 ACID. BROMO-CAPROIO ACID v. BROMO-HEXOIS

BROMO-CARBANILIO AOID v. PHENYL-CAM-BAMIC ACID.

Acetyl derivative C, H, AcBrN. [128°]. From acetyl-earbazole and Br. Laminæ, v. sof. alcohol and boiling toluene (Ciamician a. Silber,

TRI. BROMO. CARBOPYRROLIC ACID v. TRI-BROMO-PYRHOL-CARROXYLIC ACID.

y-BROMO-CARBOSTYRIL C. H. NOBr i.e.

C(Ur):Cl1 (Py. 1, 3)-Bromo-oxy-\N: ¢(OH)

quinoline. [265]. Formed by boiling o-amidophenyl-propiolic acid with dilute HBr (Baeyer a. Bluem, B. 15, 2149). Prepared by the action of bromino on carbostyril other and saponification of the product by heating with HCl (Friedländer a. Weinhorg, B. 15, 2682). Needles; may be sublined. The Br is not replaced by boiling with alcoholic KOIf, but requires to be fused with KOII at 200°C.

BROMO - CARVACROL C. 11, BrO i.e. C, H, Mo(OH)Br(C,H,) [1:2:3 or 5:4]. From bromooymidino and INO... Oil (Mazzara, G. 16, 194).
BROMO-CHLORAL v. Di-curoro-Bromo-Alba-

BROMO-CHLORO- v. Chroko-bromo-,

BROMO-CHLOROFORM v. Dientono-bromo-METHANE.

BROMO-DICHROMAZIN v. TRI-AMIDO-PHEмог., р. 172.

BROMO-CHRYSENE v. CHRYSENE. BROMO - CHRYSOQUINONE v. CHRYSO -

BROMO-CINCHONINE v. CINCHONINE. BROMO-CINNAMENE v. BROMO-STYRENE DI . BROMO - CINNAMENYL - THIENYL

KETONE P. THIENYL: DI-HROMO-STYRYL RETONE. a BROMO-CINNAMIC ACID Call, Bro, i.e. C.H. CH: CBr.CO H [131°].

Formation .-- 1. Together with allo-abromocimmunic acid by the action of alcoholic KOH on the di-bromide of cinnamic acid C. H., CHBr. CHBr. CO. H (Glaser, A. 143, 325). -- From exo tri bromo β-phenyl-propianic acid and water at 100" (Kinnicutt a. Palmer, Am. 4, 26; 5, 386).

Preparation. - Dibromide of cinnamic acid (50 grms.) is dissolved in hot alcohol and mixed with the calculated quantity (2 equivalents) of potash dissolved in alcohol. The liquid is neutralised with 11Cl and the alcohol boiled off. The solution of the mixed potassium salts is filtered from bromo styrene [218]. The two acids are separated by partial precipitation with HCl, the a-acid coming down first (Barisch, J. pr. 128, 178).

Properties. Long needles (from water or chloroform). V. c. sol, alcohol.

Salts .- NII, A': actionescent groups of flat neodles, sl. sol. cold water .- AgA'; sl. sol. water. -BaA',: thin rhombio lamine. S. 12 at 6°. Insol. alcohol.

Reactions.—1. Sodium-amalgam reduces it to 8-phenyl-propionio acid .- 2. Alcoholic KOH gives phenyl-propiolic acid, -3. Br gives a tribromo-phenyl-propionio acid [132°] .- 4. Both a and allo-a-bromo-ciunamic acids may be reduced by sino dust and glacial acetic acid to cinnamio

BROMO CARBAZOLE C1:H.BrN. [199°]. (Michael, J. pr. [2] 85, 857).—5. Is not altered from its acetyl derivative and alcoholic KOH. Rhombic plates, v. sol. water.

Methyl ether A'Me: (159°) at 14 mm. Formed by distillation of the isomeric methyl other corresponding to the acid (120°) under

ordinary atmospheric pressure (A. a. S.). Ethyl ether A'Et: (290°); (202°) at 30 mm.; (160°) at 10 mm.; (188°) at 30 mm. V.D. =8.715 (obs.) (Michael a. Browne, B. 20, 551). Formed by distillation of the isomeric ethylether corresponding to the soid [120°] under ordinary atmospheric pressure (Anschütz a. Selden, B. 20, 1384). Prepared by passing HCl into a solution of the acid in alcohol (4 pts.) (Parisch, Inc. cit.). By cold conc. H.SO, it is converted into benzayl-acetic other (Michael a. Browne, B. 19, 1392).

Chloride: (153°) at 12 mm.; clear oily liquid. Formed by the action of PCl, upon salts of the acid [131°] or of its isomeride [120°].

Amide: [1190]; very thin pearly plates; sl. sol. hot water.

Anilide: [80°]; small white needles, which change into six-sided plates (A. a. S.).

E.co-Bromo-einnamic acid C.H. BrO. [134°]. Formed, together with the isomeride [159°] by the addition of 11Br to phenyl-propiolio acid. Long needles (from water). Thick rhombic prisms (from chloroform). V. sul. alcohol and hot benzene, sl. sol. CS₂, hot petroleum-ether and hot water.—A'Nll₄*: flat needles, sol. hot water, sl. sol. cold. A'Ba: glistening rhombic plates; S. '776 at 6°, in. sol. hot water. It is doubtful whether this acid is a distinct isomeride or is identical with the preceding acid; the solubility of the baroun salt appears to point to the former conclusion (Michael a. Browne, B. 20, 550).

Allo-a-Bromo-cinnamie acid

Calla Cli:CBr.COall. [1202]. Formed, together with α-bramo-cinnamic acid (v. Preparation) by boiling the dibromide of cinnamic acid with alcoholic KOII (Glaser, A. 143, 330). Six-sided lamine (from water) or thick prisms (from etbor). V. sol. boiling water.

Salts .- KA': deliquescent needles .- AgA'. The ammonium salt is deliquescent and v. sol. water (difference from a acid).

Reactions .- 1. Reactions 1, 2, and 4 described under a-bromo-ciunamic acid are alse exbibited by the allo-a- acid. Reaction 3 gives, however, a different tri-bromo-phenyl-propionio acid [45° 48°]. 2. On dissolving in 11,50, and pouring into water two products are obtained: (a) A substance C₁₀H₁₂O₁; yellow needles [above 260°], sol. alkalis, alcohol, ether and bonzens. (b) A substance C₁₁H₁₂Br₂O₂; large pearly platos [above 260°], inset alkalis, sol. phonol, nitrobenzeno and aviline, insol. most other solvents, forms a orystalline molecular compound with plienel; on reduction with zinc-dust and acetio acid L gives a substance C₁, H₁₄O₂ which forms colourless crystals, [127°], sol. alcohol, sl. sol. hot water (Leuckart, B. 15, 16).

Methyl ether A'Mo: (146°) at 11 mm.; from A'Ag and Mel; by distillation at ordinary pressure it is converted into the methyl ether of the isomeric acid [131°] (Anschütz a. Solden, B. 20, 1383).

Ethyl other A'Et: (174°) at 30 mm.; (177°) acid, which seems to be the same in both cases at 30 mm.; V.D. = 8.828 (obs.) (Michael

Browne, B. 20, 551). From A'Ag and Etl. By distillation under ordinary pressure it is convorted into the ethyl ether of the isomeric acid [131°] (Anschütz a. Selden, B. 20, 1384).

C.H. CBr:CH.CO.H (?) [159°]. Formed, together with the acid [134°], by the action of aqueous HBr upon phenyl-propiolic acid at 0°. Long flat needles (from water); or rhombic plates (from alcohol). Sl. sol, cold alcohol and benzene. By cold conc. KOII or NII, it is at once converted into an indifferent insoluble nil. Heating with IIBr forms phenyl acetic aldehyde and acctophenone. It combines with Br forming a tri-bromo-hydro cinnamic acul [148].

Salts .- NH, A' : long soluble needles .-BaA', aq: soluble concentric prisms .-- l'bA',: amorphous pp.—CdA'₂: concentrio needles (Michael a. Browne, B. 19, 1378).

Ethyl other A'Et: (151') at 15 mm.; V.D. 8948 (obs.) (M. n. B., B. 20, 551).

Bromo-cinnamic acid [153°]. This was obtained by Erlenmeyer a. Stockmeier by the addition of HBr to phenyl-propiolic acid. Aqueous Na.CO, converted it into phenyl-acetylene, a-bromo-styrene Call Chr.CH, and phenyl propiolic acid. Successive treatment with cone. H2SO, and water gave benzoyl-acetic acid and bromo-neetophenone. Heating with HBr gavo acetophenone and a little phenyl-acetic aldehyde (Erlenmeyer, B. 19, 1936). According to Michael a. Browne this acid is a mixture of the two acids [159] and [134].

o-Bromo-cinnamie acid [1:2]

C.H. (Br) CH:CH.CO.11. Bromo-B-phenyl-acrylic acid. [213-]. Formed by diazotising o-amidooinnamic acid and boiling the diazo-compound with HBr (Gabriel, B. 15, 2294). Flat colour-less needles or scales. Sci. alcohol, ether and acetic acid, sl. sol. chloroform, benzene and CS ...

m-Bromo-cinnamic acid

[1:3] C, II, (Br), CH; CH, CO, H. [179]. Formed like the preceding from mannido-cinnamic acid (Galiriel, B. 15, 2296). Long needles. Sol. alcohol, acetic acid, hot benzene, chloroform, and CS.

p.Bromo-cinnamic acid

[1:4] C_cH₄(Br),C_cH₂,CO_cH₃ [c. 253²]. Fine needles. From p-amido-cinnamic acid (Gabriel,

B. 15, 2300).

Di-brome-cinnamic acids C.H.Br₂O₂ i.e. C.H.,CBr:CBr.CO.H. Bromine unites with phenyl-propiolic acid producing di-brome-cinnamic [139] and allo di-bromo-cinnamic acid The acid [139°] is not converted into an indenaplithene derivative by cone. Il SO, whereas the acid [100] forms CoH. C.Br. [123°] (whence hydroxylamine forms an oxim [195°], and aniline forms an anilide [170°]) (W. Roser, B. 20, 1273, 1576).

a-BROMO-CINNAMIC ALDEHYDE C.H.BrO i.e. C.H., CH. CBr. CHO [73°]. Formed from cinnamic aldehyde di bromide by splitting off HBr (Zincke a. Hagen, B. 17, 1815). Thick tables or large monoclinic prisms. CrO, gives bromo-cinnamic acid [131°]

Phonyl.hydraside O.H.C.HBr.CH:N.HC.H.(130°),glistening yellow

BROMO- CITRACONIC ACID C.H.Bro. LA CH, C(OO, H): CBr.CO, H(?).

Formation .- I. From its anhydride by boiling with water .- 2. From citra di bromo pyrotartario acid, water, and Ag.O in presence of a trace of HCl: C.H.Br.O. = HBr & C.H.Br.O. (Bourgoin, C. R. 88, 312; 89, 418; A. Ch. [5] 19, 285). Properties .- Very unstable, giving off water even over H.SO, and changing to the auhydride.

Reactions. -1. Sodium amalaim forms pyrotartaric acid. - 2. Boiling KOtl forms a syrapy dibasic acid C.H.O. 3. Exeporation of its sulntion produces bromo citraconic . : hydride and, at a low temperature, the acid C.H.D., but at a high temperature 'brome citronic acid' C.H.BrO. (B.). - I. The aqueous solution treated with the equivalent of anitine gives a crystalline pp. of the neid aniline salt. [121]. Standing under water for a few hours, or heating the aqueous solution for a minute, converts it into the

C(CH_s) - CO e | NPh. [145°]. This CBr - CO phenylimide #

forms stellate groups of prismatic needles. sol, hot water; v. sol, hot alcohol; incol, diluto HCl (Michael, Am. 9, 180). An intermediate body is the acid milide C.H.Br(CO.H).CONPhH

[212] (Michael, B. 19, 1373), Salts, — (NH₁), A', — K.A'', — CaA'' 2aq, — CaA'' 14aq. — InA'' aq. — Ag.A''. Decomposed by water at 130° giving off CO₂, and allylene,

 Anhydride C (H₃Bro₂, 100); (220).
 S.G. ⁷ (1935) (Kekulé, A. Suppl. 1, 130; 2, 97;
 Lagermark, Z. 1870, 299; Fittig a. Krusemark, A. 206, 19; Bourgoin, Bl. (2) 31, 252; 32, 388). Formation. -1. From citraconic anhydride and Br at 140?. 2. By distilling citra dibromo-tarturic acid. - 3. From pyrotartaric acid, Br, and water at 120°. Properties. Lamium (from CS); sl. sol. cold water, v. c. sol. alcohol und other; volatile with steam.

Imilio C.H.BrO.NH. [c. 1812]. pyrotart imide or citraconimide and Br at 1500 (Mendini, G. 15, 182). Lamine; may be sublimed. - C. H. BrO NAg.

Di-bromo-citraconie acid. Imide. C.H.Br.O.NH. [144]. From pyrotartrimido or citraconimide and Br (M.). Lamina; may be sublimed. C.H.Br.O.NAg.

BROMO-CODEINE v. CODEINE.

DI - BROMO - COLLIDINE v. DI-BROMO-TRI-MUTHYL PYRADINE.

BROMO- COMPOUNDS. Bromine anites directly with most unsaturated compounds, but there are some cases in which it does not combine with them in the cold, e.g. fumaric, mesaconic, teraconic, terebilenio, aconic, o- and p- coun vie, and (B)-hydropiperio scids (cf. Pittig, A. 227, 29). Aromatic compounds combine with great difficulty with bromine, but when reduced to their di- or tetrahydrides they take up broming as readily as tho ordinary unsaturated compounds. Bromino displaces hydrogen in saturated compounds, the displaced hydrogen being given off as 11Br; this reaction usually requires the aid of heat: the substance is generally heated with bromine and a little water in a scaled tube.

Hydroxyl may be displaced by bromine by treatment with HBr or PBr,; instead of PBr,

of the two being added gradually.

Chlorine may be displaced by bromine by treatment with the bromide of K, Mg, Ca, Sr, Ba, Al, Mn, or Co (Köhnlein, A. 225, 194). CdBr., SbBr, and AsBr, sometimes act similarly; thus AsBr, at 1 to completely converts chloroacetic acid and benzyl chlorido into bromoacctic acid and benzyl bromide respectively.

Indine may be displaced by bromino by means of the bromides of Cu, Ag, Hg, Su, Pb, As, and Sb. Bilir, at 160° only partially converts Etl into EtBr. Bromino itself can dis-place iodine (R. Meyer, J. pr. [2] 34, 104). Amidogen may be displaced by bromine

by using the diazo- reactions (p. 399). In aromatic compounds it is sufficient to add HNO, to a hot solution of the amino in HBr (Losanitsch,

B. 18, 39, v. also Amere).

Carriers. - The displacement of hydrogen by Br is assisted by the presence of carriers which first condine with the bromine, and then pass it on to the organic tody. The most suitable carriers for bromine are: iodin :; Al Bra (Gustayson, B. (0, 971) for benzene an I homologues; nunorphou: P for fatty acids (Hell a. Gautter, B. t4, 891), SbBr., Fe,Br., Felli, and, better still, FeCl. (Scheufelen, A. 231, 151; cf. Willgerodt, J. pr. [2] 34, 264).

Bromination of organic acids. The following is a very convenient method, which depends upon the fact that the acid bromides and anhydrides are much more easily brominated than the acids themselves. The acid is mixed with amorphous phosphorus in quantity sufficient to convert it into the bromide or anhydride, and ! the corresponding quantity of bromine added together with the extra amount required for brominution. The naxture is heated to 100° till decolourised, when the reaction is complete [Hell, B. 14, 891). The brondication of futty acids may also be readily effected by boiling their chlorides with Br and CS, (Michael, J. pr. [2] 35, 92). Alkalis or water acting upon abrome-acids usually produce a oxy-acids, with B. bromo-acids they form unsaturated acids, while y chloro soids become lactones (Fittig, A. 195, 169; Firlenmeyer, B. 14, 1318, 15, 49).

Bromination of aromatic hydrocarbons. Bromine in presence of a carrier enters the henzene nucleus; bromino alone, or in presence of too little iodine, enters the eidt chains of boiling hydrocarbons. Sunlight promotes substitution in the side chains (Schramm, B. 17, 2922; 18, 350, 1272; 19, 212).

Brommation of aromatic amines. Bromino goes first into a position p to the NH and then into the o- positions, but not into a m- position (Nevile a. Winther, B. 13, 962; v. also Aromatic

Stability of brome-compounds. Brome-compounds are less stable than the corresponding chlore- compounds; thus brome acetic and bromo-propionio acids may be converted by reduced ailver into succinic and adipio acids respectively, in this they differ from the corresponding chloro- but resemble the iodo- acids. The relative stability of the alkyl bromides depends upon the reagents attacking them; thus Zn and H.SO, reduces iso propyl bromide more

red phosphorus and bromine may be used, one isopropyl, isoamyl, buiyl, propyl, ethyl; on the two being added gradually. other hand, alcoholic NaOH attacks propy. bromide more vigorously than isopropyl bromide the order being ethyl, propyl, isoamyl, butyl isopropyl (Remsen a. Hillyer, Am. 8, 251).

Other characteristics of bromo- compound may be inferred from the article on CHLORO-COMPOUNDS; the bromo- compounds resemble the chloro- compounds in almost every respect.

BROMO-CONIINE v. CONIINE.

BROMO - COUMARIC ACID O,H,BrO, Bromo-o-oxy-cinnamic acid. Methyl deri vative MeO.C.II, C.IIBr.CO.II. [1719]. S. (CS.) 32 at 14°. From the methyl derivative of α- or β- di-brono-melilotic acid McO.C., Hp. C.H., Br., CO. II and aqueous KOH (Porkin, C. J. 39, 422). Small prisms (from CS₂) sl. sol. boiling water, v. sol. alcohol. Boiling dilute KOH forms the methyl derivative of cumarilic acid MoO.C.H.,C.CO.H. Ethyl derivative Eto.C.H.,C.HBr.CO.H

[161°]. From EtO.C.H.C.H.Br...CO.Et and cold alcoholic KOH (P.). Flat prisms; sl. sol boiling water, m. sol. CS., v. sol. alcohol.

(a)-BROMO-COUMARIN Call Bro. From coumarin dibromide and alcoholio KOII (Perkin, C. J. 23, 368). Prisms; converted by alcoholic KOH into cumarilic acid.

(β) - Bromo · coumarin C, II, BrO, From sodium bromo.o.oxy.benzoic aldehyde and Ac₂O. Bhombic prisms (from alcohol, (Perkin, C. J. 24, 37).

(a) Di-bromo-coumarin C_oH₁Br₂O₂. [183°] From commarin, Br, and I (P.). Needles; alcoholic KOII forms bromo cumarilic acid.

(β) Di-bromo-coumarin C, II, Br, O,. From sodium di-bromo-o-oxy-benzoio aldehyde

and Ac O. Needles (P.). BROMO-o-CRESOL C.H.McBr(OH) [1:3?:6] From bromo-o-toluidine (Wroblewsky A, 168, 165). Golden needles; v. sl. sol. water; the aqueous solution is turned green by Fe₂Cl₂-KC II BrOnq: red scales.

Bromo - m - cresol C. H. MeBr(OH) [1:3:5] [57"]. From s-bromo-toluidiue by diazo- reaction

(Nevile a. Winther, C. J. 41, 421).

Bromo - p - cresol C, H₃(CH₃)Br.OH [1:3:4] (211°). S.G. $^{25}_{23}$ = 1.5468. Liquid. Formed by the action of dry bromine upon sodinm p crssol Methyl ether C, II, MeBr(OMe) : (226°):

S.G. 25 1.418; liquid (Schall a. Dralle, B. 17 2530).

Di bromo o crssol CaH, McBr. (OH). [570] From o-cresol and Br. Volatile with steam (Werner, Bl. [2] 46, 278).

Di-bromo-p-cresol C. H. (CH₃)Br₂-OH [1:3:5:4]. [49°]. Large crystals. V. sol. alcohol, v. sl. sol.

water. Excess of Br forms C.H.Br. (OBr) (W.).

Benzoyl derivative C.H.MeBr. (OBr) [91°], white needles (Schall a. Dralle, B. 17, 253°; Werner, Bl. [2] 46, 278).

Tri bromo m cresol C. HMeBr. (OH). From m-cresol (W.). Needles (from alcohol). Tetra-bromo-p-cresol C.Br.Me(OH). [109°]. From p-cresol (Baumann a. Brieger, B. 12, 804).

Platos; slowly decomposed by cold brominewater into CO, and tri-bromo-phenol. BROMO-CRESOL SULPHONIC AOID

C.H.McBr(OH)(SO.H) [1:2 or 6:57:3]. quickly than propyl bromide, the order being o bromo-toluene by sulphonation, nitration, so -Bah', 4jaq. -Bah', 8aq. -Pbh', 2aq. Brome-cresol sulphonic acid

C.H.MeBr(OH)(SO.H) [1:4:57:3]. From (3,1,2). bromo-toluene sulphonio acid by nitration, reduotion, and diazolisation (S.) .- Ba A', 3mj.

Bromo-cresol sulphonic acid C, H2MeBr(OII)(SO,II) [1:4:7:2]. From (4.1,2)-

bromo-toluene sulphonic acid in a similar way. -BaA', aq.

Di-bromo-cresol sulphoaic acid C.HMeBr.(OH)(SO3H). From (2,1,4) amidotoluene eulphonic acid and Br (llayduck, A. 174, 353). - KA'nq. BaA'. 8laq.

a-BROMO-CROTONIC ACID C.II Bro. i.c.

CH, CH: CBr.CO.H. [106.5°].
Formation.—1. From aa di-bromo n-butyric acid and alcoholic KOH, NH, baryta-water, or Ag₂CO₂ (Michael a. Norton, Am. 2, 15; Erlenmeyer a. Müller, B. 15, 49). 2. From αβ-dibromo-butyric other and alcoholic KOH (Michael a. Browne, Am. 9, 280). - 3. Together with bromopyrotartaric acid by the action of bromine on , propane tri-carboxylic acid (Bischoff a, Guthreit, B. 14, 616).

Properties.-Long needles (from water); needlee (from ligroin); m. sol. cold, v. sol. hot, water .- AgA': white needles, quickly affected by light .- BaA', 2aq.

Allo-a-Bromo-crotonic acid

CH₃,CH:CBr.CO.H. [90'] (K.); [92°] (M. a. N.). From aB-di-bromo-butyrio acid (dibromide of crotonic acid) by treatment with alcoholic KOH (Michael a. Norton, Am. 2, 15) or NaOHAq (C. Kolbe, J. pr. [2] 25, 391).

Properties.—Long needles (from water); needles (from ligroin); m. sol. hot water. - AgA'; needles, little affected by light. BaA', 33aq. -

CaA', 3nq.

B-Bromo-crotonic acid ClI, CBr:CH.CO.H. [95°]. From tetrolic acid and cone. HBrAq at 0° (Michael a. Browne, Am. 9, 277; J. pr. 12, 37, 258). Flat needles (from water); feathery groups of tough needles (from higroin); st. sol, cold water, in. sol. hot water. - AgA': amorphous, readily affected by light. BaA' aq. -- KA'.

aß-Di-bromo-crotonic acid

CH, CBr.CBr.CO.11. [96°]. From tetrolic acid and Br (Pinner, B. 14, 1081). Boiling with

Ag₂O gives (C,11,Br), [116°], BROMO-CUMALIC ACID C,H BrO,(CO,II). [176°]. Formed by the action of bromine on cumalic acid in acetic acid solution (Pechnann) a. Welsh, B. 17, 2396). Colourless glistening needles. Can be distilled in small quantities. V. sol. alcohol, other, acrtic acid, and chloroform, more sparingly in benzene, insol. ligroup.

Methyl ether A'Mc: [134], prismatic needlee, sol. alcohol and benzene, sl. sol. ether, insol. water. Aqueous NII, converts it into the methyl ether of brom-oxy-micothnic anid.

o-BROMO-n-CUMENE C,H,Br i.e. C,H,BrPr [1:2]. Bromo-n-propyl-benzene. (222 i. V.). A mixture of this body with the p-isomeride is formed by the action of Br on propyl-benzene in the dark or on ethyl-benzene mixed with 1 in diffused daylight (Schramm, P. 18, 1274). KMnO, gives o-bromo-benzoio acid.

p-Bromo-n-cumene C.H.BrPr [1:4]. (220° cor.) From C.H.Pr and Br at 0° in presence of

Suction, and disactisation (Schäfer, A. 174, 861), [2] 84, 101). HNO, (S.G. 1-2) forms p-bromebenzoic seid.

> 8.Brome.n.cumene C.H., CHBr.CH., CH., Formed in the cold by the action of bromine (I mol.) upon n-onmeno in direct cunshino. By further bromination in sunlight it gives 8-dibromo-n-cumene C. H. CBr. CH, CH, CH, but in the dark at 100° it yields as-di-bromo-n-cumone C. H., CHBr.CHBr.CH, (Schramm, B. 18, 1275), åß-Di-bromo-x-cumone Call, CHBr. CHBr. CH, From allyl benzene (q.v.) and bromine (Righeimer, A. 172, 131; Radyiszewski, C. R. 78, heimer, A. 172, 131; Indzbazewski, C. R. 78, 1153; Perkin, C. J. 22, 668). From n-canneno and bromine at 160° (Wispel ... Zider, A. 218, 381; cf. S. supra). Silky needles (from alcohol). BB-Di - bromo-n - cumoae C. H. CBr .. CH .. CH .. Phonyl-ethyl-ket-ne-di-bromide. Formed in tho cold by the action of bromine (2 mole.) upon n-camene exposed to direct sunshine (Schramm B. 18, 1275).

waß-Tri-bromo-n-oumcao

C.H., CHBr.CHBr.CH Br. Stycerin tri-bromhydrin. (121°). From cinnamyl bromide and Br. or by the action of HBr on the dibromide of cinnamyl alcohol (Grimanx, Ft. 20, 120).

Tetra-bromo. n. cumene CallaBra. (Fittig, A. 149, 327).

v. Brome-cumene Call Brl'r [1:2]. Bromeisopropul-benzene. (206" cor.) at 710 mm. From isopropyl-phenol and PBr, (Fileti, G. 16, 131).

p. Bromo-cumeno C. H. Bri'r [1:4]. (217). S.G. 15 1:3014. Prepared by brominating cumene at 0 in presence of I (Jacobsen, B. 12, 430). With benzene, Na, and CO, it gives cuminie ucid (R. Meyer, J. pr. (2) 31,93).

Penta-bromo-cumene Call Br., [97]. From cumene and Br (Men el. Z. 1867, 322; Fittig. A. 119, 326). Needles; sl. sol. cold alcohol. Bromo & camene C.H.(CII,) Br (1:2:4:5].

j. (227). White plates.

Formation. 1. By the action of cuprous bremide upon diazo-panado cumens (Haller, B. 18, 93) .- 2. By the action of bromine (1 mol.) in the dar! upon pseudo-cumene; the yield is 60 p.c. (Boi-tem, A. 137, 323; Fittig, A. 139, 188; 145, 138; Schramm, B. 19, 216; Süssenguth, A. 215, 243).
3. By the action of bromine upon an aqueous solution of pseudo-cumencsulphonic acid (1:2:4:5) 76 p.c. is converted into bromo v-cumene, the remainder forming bromobromo y-emmene, the remanance forming monacy cumene sulphonic acid C_ellMe_sBr(SO_sH) [1:2:13:5] (Kelbe a Pathe, B. 19, 1547).

Bromo - y - cumene C_eH Me_sBr [1:2:4:3), (226 '-229'). Colourless oil. Formed by the

bydrolysizof the aulphonic acid C. HMe, Br (SO 11) (1.2: £3:5) by means of experimental strain at 200° 210°. By Mol and Na it yields externmethyl-benzene (Keibe a. Pathe, B. 19, 1551).

Bromo-4-cumeae C. H. Me, Br. (1):24:63 (2372), Liquid; still fluid at 15°. Obtained by hydro-lysis of the suphoric acel (1):34:5(2). By sulphonation it is recovered into the same sul-

phonic acid (Jacobsen, B. 19, 1223).

ω-Bromo-ψ-cumoae C.11 (CII,), CI1, Cl. Pscudocumul bromide. Liquid. Decomposes on distillation. Formed by the action in direct sunshine of 1 mol. of bromine upon 1 mol, psoudocumene (Schrainm, B. 19, 217).

Di-bromo-\u03c4-cumene CaliMe,Br. [1:2:4:5:6] [64°]. (294°). Formed by the action of 1 mol 1 (Meyer a. Müller, B. 15, 698; R. Meyer, J. pr. of bromine upon 1 mol. mono-bromo-pseudosumene by gaslight, or in the dark (Schramm, B. 19, 216; Jacobsen, B. 19, 1220). Long flat needles, v. sol. alcohol.

ω1-ω2-Di-bromo.ψ-oumene C4H2(CH2)(CH2Br)2 [1:2:4]. Pseudo-cumylene bromide. [97°]. Flat glistening needles (from petroleum-ether). V. sol. alcohol and benzene. Formed by the action

of 2 mols, of bromine upon 1 mol. pseudo-cumene in direct sunshine (Schramm, B. 19, 218; Hielt a. Gadd, B. 19, 867).

Tri-bromo-4-cumene C, Me, Br, [1:2:4:3:5:6].

[226"] or [233" cor.]. V. sol. hot teluenc, sl. sol. alcohol. Formed by the action of Br (3 mols.) in the dark con ψ-cumene (1 mol.) (Fittig a. Laubinger, A. 151, 264; Schramm, B. 19, 217; Jacobsen, B. 19, 1222).

BROMO-4-CUMENE-SULPHONIC ACID

C_ollMo_oBr(SO_oll)[1:2:4:5:6]. [c.121°]. Formed by dissolving bronno-4-cumenc [73°] in warm slightly fuming 11₂SO₄. Needles (containing slightly furning 11.SO₄. Needles (containing 2aq). By treating the Na salt with zinc-dust and aqueous NII, it is debrominated with production of (1,2,4,6) pseudo-cumene sulphonic acid.

Sults. - A'Na aq. - A', Ca 3aq. - CuA', 4aq. --BuA', !, aq. -- KA' aq. Amide C, IIMc, Br(SO, NH.): [185°]; small

needles, v. sol. alcohol, nearly insol. water (Jacob-Ben, B. 19, 1218; Kelbe a. Patho, B. 19, 1553).

Bromo. 4. cumone-sulphonio acid C.HMe.Br(SO.H) (1:2:4:6:3). Formed, together with the di-bramo- acid, by the action of CISO, H upon di bromo-\psi-cumene. NaA' \aq

Amide C, HMe Br(SO, NH2): [158°]; thin

ncedles (Jacobsen, B. 19, 1223).

Brome-\psi-cumenc-sulphonic acid $C_bHMe_3Br(SO_3H)$ [1:2, f:3:5]. [116°]. Colourless needles. Formed by the action of bromine upon an aqueous solution of pseudo-cumenesulphonic acid [1:2:4:5]; 76 p.c. of the pseudocumene-sulphonic acid is converted into bromopseuda-cumena (73°), the remainder yielding the brome-sulphonic acid. It is also corned by sulphonation of bronto-pseudo-cumene [1:2;4:3]. The latter body is formed by the action of super-

healed steam upon it at 200°, 210°, Sults. - NaA'aq. - KA'aq. - AgA'aq. -

BaA', uq. | PbA', 3uq. | Amide | C,HMe,Br(SO,N11) : {188°]; thin needles (Hellie a. Pathe, $B.\ 19,\ 1547$).

Di-bremo-4-cumene-sulphonic acid CaMe, Br. (SO, 11) (1:2:1:5:6:3). Obtained by sul phonation of distronto pseudo-comene with sulphuric chlorhydriu.

Salts. -- NuA'. · NuA' uq. -- BaA'. Amide C.Mc.Br (SO,NH.): [above 250°];

plates (Jacobsen, 7, 19, 1222). BROMO . 4 . CUMENOL C. IIMe.Br.OII

[1:2:4:3:5], [35°], Formed by bromination of pseudo-cumenol in cold acetic acid. Slender yellow needles. Insid. water, v. sol. other solvents (Reuler, B. 1f. 29; Auwers, B. 18, 2657).

Bromo - iso - cumonol C.H.PrBr(O11) [2:4:1]. [49°]. Bromo isopropyl-phenol. From o-isopropyl-phenol (o-isocumonol) and Br (Fileti, G. 16, 117). Decomposes ut 250

Methyl other CallaPrBr(OMe). (250°).

Di . brome . 4 . onmenol

C.Me.Br. OH [1:2:1:3:6:5]. [149°]. Formed by bromination of pseudo-currenol dissolved in a

prisms or needles. Insol. water, m. sol. alcohol.

scetic scid, and benzene, v. sol. ether.

Methyl ether C.Me.Br.OMe: Formed by methylation of the above or by bromination of peeudo-cumenol-methyl ether. White needlee. Incol. water, v. col. other solvents (Auwers, B. 18, 2657).

Di-bromo-\psi-oumenol

CaMeaBr.OH [1:2:4:3:5:6], [152°]. From C. HMe, Br. [1:3:4:2:6] (Edler, B. 18, 630; Jacobsen, B. 19, 1220).

Di-bromo-iso-cumenol

C,H,PrBr,(OH) [2:4:6:1]. From o-iso-cumenel and Br (Fileti). Liquid.

Methyl ether Call I'rBr (OMe). (279°). HNO, forms a nitro-derivative

benzoic acid. From Br and cuminio acid or silver cuminato (Naquet a. Luginin, Z. 1866, 333; Gerichten, B. 11, 1719). From bromooymene and HNO₃ (Fileti a. Crosa, G. 16, 296). Exo-bromo-cuminio acid C₃H₆Br.C₄H₄.CO₂H₄.

From Br and cuminic acid at 120° (Czumpelik,

13. 3. 478

BROMO-CUMYL-BUTYRIC ACID

Cull, BrO, i.e. Call, Pr.C. H, CHBr.CO. H. [150]. From cumenyl-crotonic acid and HBr. Prisms. Decomposed by Na.CO, giving allyl-isopropyl-benzenc (Perkin, C. J. 32, 662). BROMO-CUMYL-PROPIONIC ACID

C₁₂H₁₃BrO, i.e. C₆H₄Pr.CH, CHBr.CO₂H. [87°]. From cumyl-acrylic acid and HBr (Perkin, C.J.32, 661). Resolved by beiling water into HUr and cumyl-acrylic acid. Na CO Aq forms vinylisopropyl-benzene.

Di-bromo-cumyl-prepionic acid

C. H. Pr.CHBr.CHBr.CO.H. [190°]. From cumylnerylic acid and Br (Widman, B. 19, 258). BROMO-CUMYL-VALERIC ACID

C.4H.5BrO. i.e. C.H.Pr.C.H. CHBr.CO.11. enmenyl-angelic acid and IIIr. 1 Prisms. Na,CO,Aq gives isopropyl-butenyl-benzene (Perkm, C. J. 32, 663). BROMO - CYANO - BENZENE v. Nitrils of

BROMO-BENZOIC ACID.

BROMO p-CYMENE C10II 13 Br i.e. C₄H₃(CH₃)(C₄H₂)Br [1:4:2]. Bromo p methyl-n-propyl - benzene. (234° i.V.). S.G. 17:1-27.

Formation .- I. From cynicne and Br (Landolph, B. 5, 267).—2. Together with brono-p-cymene - sulphonic acid C_aH₂MePrBr(SO₂H) [1:4:5:2] by the action of bromino upon an aqueous selution of p-cymene-sulphonie acid (Kelbe a. Koschilzsky, B. 19, 1730). - 3. Tho same or the following brome-cymene is fermed by hydrolysis of bromo-p-cymenc-sulphonic acid

C.H.MePrBr(SO₃H) [1:4: 5or6 :2! (K. a. K.). Bromo-Poymeno C.H.(CH₃)(C₃H₃)Br [1:4:3]. (232°). From thymol and PBr, (Fileti a. Crosa, (i. 16, 287). Formed also by the action of bromine upon an aqueous solution of p-cymenesulphonic acid (Me:Pr:SO.II = 1:4:3) (Claus ...

Christ, B. 19, 2165, v. supra).

Di.bromo-p-oymene C₈H₁MeBr₄Pr. (27
S.G. 12 1-596 (Claus a. Wimmel, B. 18, 903).

(a).Bromo-m-isocymene C_s11_sMePrBr [1:3:6]. (225° cor.). Formed, together with bromo-isocymene sulphenic acid, by the action of bromine small quantity of acctic acid. Long colourless ou m isocymeno sulphonio scid (Kelbe, A. 210, 6; 285, 281). Oxidised by dilute HNO, to

bromo-toluie acid [210°].
(β)-Bromo-isocymene O_cH_sMe(Pr)Br [1:3:4]. (221°). Got by distilling C.H.Me(1'r)Br(SO.K) [1:3:4:6] with superheated steam (Kelbe a. Czarnomski, A. 235, 293). Dilute HNO, oxidises it to bromo toluic acid C.H. MeBrCO.H [1:4:3] [153°].

Di-bromo-m-Isocymeno C1411:2Br... (273)uncor.). Obtained by brominating (a)-bromom-isocymene sulphonie acid (Kelbe a. Czarnomski, A. 235, 281). Converted by furning HNO. intonitro-bromo-iso-cymene $C_{16}H_{17}(NO_s)Br$ [83].

Bromo.p.cymono-sulphonic acid $C_uH_2MePrBr(SO_uH) = [1;1;5or6;2],$ methyl-n-propyl benzene sulph m.cacid. Formed, together with bromo p cymene C.H.Mel'rBr [1:4:2], by the action of bromine upon p-cymenesulphonio acid C, 11, McPr(SO, 11) [1:1:27 in aqueous solution at 40 1502 (Kelben, Koschitzky, B. 19, 1730). Formed also from cymidine by sulphonation and diazotisation (Widman, B. 19, 248). Sodium amalgam reduces it to ncymene sulphonio acid.

Salts. KA'aq. BuA'223aq. S.1 at 173,-BaA', 11, aq. —CuA', 12aq.

Amide C,H,MePrBr(SO,NH,): [152]. Bromo-p cymene sulphonic acid

 $C_sH_2\mathrm{MePrFr}(\mathrm{SO}_sH)$ [1:1:2:5]. Formed by sulphonation of bromo-p-cymene (1:1:2) with H SO, (Kelbe a. Koschitzsky, B. 19, 1732; Claus a. Christ, B. 19, 2163; Remsen a. Day, Am. 5, 151), or CISO, II (Paterno n. Canzoneri, G. 11, 126). Long needles containing 3aq [c. 60°]. By zine-dust and NII, it is easily debrominated to p-cymene-sulphonic acid (1: t:a)?.

Salls, KA' 3aq.— PbA₂, S, 2.1 at 30'.— AgA'.—CaA'₂ 8aq.—NaA' 1!aq. (R. a. b.).— ZnA'₂ 8aq. (R. a. b.).—Ma'A'₂ 9!aq (R. a. b.).— CaA', 9!aq. (R. a. b.).—CaA', 5aq.—BaA', 9!aq (R. a. b.).—EaA'₂5aq.—8. 1:37 at 27'.—CaA'₂8aq.—Amide.—C,H.McPrB(SO,NH₂): [188']

(K. a. K.) [195°] (C. n. C.).

Chtoride CH McFrBrSO Cl :82 %

Bromo-p-cymene-sulphonic acid $\mathbf{C}_{\mathbf{s}}\mathbf{H}_{\mathbf{s}}(\mathbf{C}\mathbf{H}_{\mathbf{s}})(\mathbf{C},\mathbf{H}_{\mathbf{s}})\mathbf{Br}(\mathbf{S}\mathbf{O}_{\mathbf{s}}\mathbf{H})=\{1;1:5:3\}$ nucor.]. Formed by bromination of an aqueous solution of p - cymene - sulphonic C.H.M.Pr(SO,H) (14.3). Glistening colourless plates (Claus a. Christ, B. 19, 2166).

(a) Bromo iso cymene sulphonic acid

C,H,Me(Fr)Br(SO,H) [1:3:6:1]. [657] and [1267]. From (a) broanc iso cymene by sniphoration, or from (B)-is ocymene sulphonic acid by bromination (Kelbe a. Czarnomski, A. 235, 277). Ne dles containing 3aq (from water). After a week's exposure over H₂SO₂ it melts at 126% Salts.—BaA', 7aq.—CuA', 7aq.—KA' mq.—NaA' 2aq.—Ant de C₀H₁₂Br.SO₂NH₂ [1712].

(B)-bromo-m-iso-cymene sulphonic acid m isocymene sulphonic acid (Kelbe a. Czarnomski, A. 235, 272). Salts. - PbA', aq. --BaA 2. — CuA' 4aq. — KA' aq.

Am (de.— C. H. Br.SO NII, [162?]

BROMO-CYMENOL C. II Metalir(OII) [1:4:3:20r6]. From amido-cymenol by the diazo-reaction. Oil (Mazzara, G. 16, 191).

Tri - bromo - oymenol C.Me(C,H,)Br,(OH) 2:4:1]. [222°]. Glistening golden plates. formed by shaking an aqueous solution of 210). Pearly plates (from alcohol).

cymenol with a solution of bromine in HBr

(Jesnrun, B. 19, 1414).
BROMO-CYMIDINE C.H.MeBr(C.H.).NH. From bromo-nitro-oymene. Liquid (Mazzara G. 16, 193)

BROMO-DECANE v. DECYL BROMEDE.

Di-brome decane Coll roller. Decylene bromide. Oil. From petroleum decyleno (Reboul a. Truchot, A. 144, 248).

*Di-bromo - decane C_b11 aBr., Diamylens bromide. From diamylene and Br (Baner, A. 135, 311). Liquid. Alcola lie KOH gives rutyl-

ene C_{le}H_m (150°).

Tri-bromo-docane C_{le}H R°, Oil. From diamylene and Br (Valz, Z. 1868, 315). Decomposes at 100".

Tetra bromo decane Call, Br. Menthens tetrabromale. From menthene and Br (Beckett n. Wright, Report of British Ass. 1875, ii. 88). Oil, split up by distillation into HBr and

Tetra-bromo decano Ciallin Br. Decenyleno tetra bromide. From decinene (165') and Br. Oil (Reboul a. Truchot, A. 111, 219).

Tetra bromo-decane CallaBr. From allylpropylidenc propyl methane (158°) and Br (Ro-

formatsky, J. pr. [2] 27, 389]. DI-BROMO-DECINENES C 10 II ,. Br. Described as hydrobromides of terpenes. V. also BornyL

BECOMED .. Di-bromo docinene C10II 16 Br. From diamylene and Br. p. 211.

Tetra bromo decineno C₁₀H₁₄Br₄. Di camphiludene dibromide [160°-164°]. Di-bromo-From camphor and PCl,Br, (De la Royère, Bl. [2] 38, 579). Unctuous tables.

7-BROMO-DECOIC ACID C10 II 11 II IV. i.e. C.H., CHBr.CH.CH.CO.H. From decencion acid (q. v.) and HBr (Schneegans, A. 227, 92). Heavy oil. Na CO, removes 11Br forming the lactor a of oxydecoic acid.

Di-bromo-decoic acid Ciallin Br. O. Di bromocapriz acid. [135]. From decenoic ('umydecylenic') acid and Br (Hall a. Schoop, B. 12, 194). Monoclinic prisme (from benzene).

DI-BROMO-DECYL ALCOHOL G, II, Br.O. Oil. From allyl-di-isopropyl-carbinol and Br (Lebedinsky, J. pr. [2] 23, 22).

BROMO DECYLENE Cigili, Br. (215), S.G. 15 1 109. Oil. From decylene bromide (v. sup.) and alcoholic KOH (Reboul a. Truchot, A. 144, 218). Alcoholic FOII forms C, II, (165°) and CETEORIC

Bromo-decylene v. MENTHYL BROMIDE.

Di bromo-decylene $C_{10}\Pi_{18}Br_{1r}$. Decinene bromide. Oil From $C_{1r}\Pi_{18}$ and Br (R. a. T.). Di-bromo-decylene $C_{1r}\Pi_{18}Br_{1r}$. From ter-

pilene hydride and Br (Montgolfier, A. Ch. [5] 19, 158).

Di-bromo-decylene C, ll, Br. From rutylene und Br (Bauer, .). 105, 344).

DI.BROMO DODECANE C. H. Br. [-15] Dodecylene bromide. Formed by the addition of Br. to dodecylene (Krafft, B. 17, 1371).

BROMO-ISO-DURENE C.HBrMe, 11:3:4:5:6]. (253°). Liquid; solidified by cold (Bielefeldt, A. 198, 338).

Bromo.s.durene C. IIMe. Br [1:2:4:5:3]. [610 By bromination of durene (Gissmann, A. 216, Bi brome a darens C. h.; Br. t.s. C.Me.Br. [1:2:8:4:5:6]. Di bromo pribnitons. [210°]. From e-durene, Br, and I. Prisms (Jacobsen, B. 19, 1213).

Di-bromo-iso-durene C10H12Br2 [1:3:4:5:2:6]. [209°]. Long fine needles. Sl. sol. hot, v. sl. eol. cold alcohol. Prepared by bromination of isodurene (Jacobsen, B. 15, 1853; cf. Jannasch, B. 8, 356).

Di bromo-s durene C.Me.Br., [1:2:4:5:3:6]. [203°]. Needles (from alcohol) (Fittig a. Jannasch, Z. 1870, 161; Friedel a. Crafts, A. Ch. [6] 1,515).

BROMO DOREMOL Ci, H. Br.OH [1180]. Formed by bromination of durenol in acetic acid. Long prisms. V. sol. alcohol and ether, insol. water (Jacobsen a. Schnapauff, B. 18, 2814).

DI-BROMO-ENNANE Collin Bry. Nonylene bromide. From Br and ennylene (from paraflin). Alcoholic KOII forme bromo-emylene Call Br

(c. 210°) (Thorpe a. Young, A. 165, 18).
BROMO-ENNOIC ACID Call, C. 11, Br. CO. 11. From ennenoic acid C, II, O, and HBr. Decomposed by warm aqueous Na₂CO₃ forming an oil, probably C.H., C.H. (Schneegans, A. 227, 83). BROMO-ENNYLENE v. Di-BROMO-ENNANE.

BROMO-ERUCIC ACID C. II. Bro. From di bromo behenic acid and nlcoholic KOII. Converted into behenolic acid by alcoholic KOII (Haussknecht, A. 143, 50).

Di-bromo-erucio acid C. Il Br.O. From behenolie acid and bramino (II.).

BROMO-ETHANE v. ETHYL BROMIDE.

Di-bromo-ethane v. ETHYLENE EROMIDE and ETHYLIDENK BROMIDE.

u-Tri-bromo-ethane CliBr, CliBr. Bromoethylene bromide. (188) nt 752 mm. S.G. 17-5 2.6189; 2.6107 (Anschülz, A. 221, 138). M. M. 12:897 at 11:7°. From CHBr:CH, water, and Br (Wurtz, A. Ch. [3] 51, 84). Also formed by the action of Br on ethyl bromide, ethyleno bromide, or ioda ethylene (M. Simpson, P. M. [4] 14, 544; Caventou, A. 120, 323; Tawildaroff, A. 176, 22; Staedel, B. 11, 1711)

Reactions .- 1. Abroholic KOH gives u-dibromo-ethylene, bromo acetylene, and acetylene. 2. u-Di-bromo ethylene is also formed by tho action of alcoholic KOAc, water and PbO, and NaOEt (Michael, Am. 5, 192). 3. SbCl, gives CHCl, CH, P. (Henry, Bl. [2] 42, 262). - 4. Benz. ene in presence of Al,Cl, produces biomo benzenc, s di-phenyl chane, and u-di-phenyl-ethane (Anschütz, A. 235, 333).

u-Tetra-bromo-ethano CBr. CH.Br. Acetyl idenetetra bromide. (103 5°) at 13 5 mm. S.G. 2.9216. From CBr.; CH, and Br. (Anechutz, A. 221, 140; Lennex, C. J. 13, 206; Reboul, A. 124, 270). Also from tri-brono-chane and Br (Donzel, B. 12, 220%. Decomposed by heat, giving off Br, and HBr. Converted by benzeno and Al,Cl, into u-di-phenyl-ethane, and s-tetraphenyl-ethane CHPh CHPh, (210); hromobonzene and acetylene dibromide being also formed (Auschütz, A. 235, 196).

s-Tetra-bromo-ethane CHBr, CHBr, Acetylone tetra-bromide. (114°) at 12 mm. S.G. 215 2-9629. Acetylene, from C.H.Br. and alcoholio KOH is passed directly into bromine. The pro-

alcohol and sine dust and CHBr: OHBr is some. rated from CH2:CHBr by fractional distillation, and ie then mixed with bromine (Anschütz, A. 221, 138; cf. Reboul, C. R. 54, 1229; Sabanejeff. B. 9, 1441; A. 178, 112).

Properties .- Smells of camphor and chloroform. Takes up moisture from air, becoming cloudy. At 185° it begine to decompose, giving off Br, and HBr, and leaving C,HBr,. water and bromine at 185° it gives C.Br. and C2Bre. Boiling alcoholic KOII forms acetylene and bromo-acetyleno. Zinc added to its alcoholio solution forms neetylene dibromide in the cold, but on warming it forme acetylene. With benz. ene and Al, Cl, it forme bromo - benzene, u-diphenyl-ethane, nuthracene, and anthraquinone (Anschütz, A. 235, 161). Al₂Cl₆ alone forms CHBr₂.CH₂Br and C₂Br₆. Tolueno and Al₂Cl₆ give di-methyl-anthracene [225°]. o. m. and p. xylene appear to give tetra-methyl-anthra.

Penta bromo ethane CBr., CBr., II. [49°] (S.); [54°] (D.); [57°] (B.). (210°) at 300 mm.

Formation. -1. From CHBr: CBr, and Br (Lennox; Sabanejeff, A. 216, 281).-2. From bromo-ethylene and Br (Denzel, B. 12, 2208) .-3. From bromo-acetylene and Br (Reboul, A. 124, 268). -4. By spontaneous decomposition of tri-bromo-ethyleno exposed to air (Demole, Bl. [2] 34, 204).-5. From acetylene tetrabromide and Br (Bourgoin, Bl. [2] 23, 173) .- 6. From euccinic acid, water, and Br (Orloveky, J. R. 9, 280).

Hexa-bromo-ethane CyBr. Carbon hexabromide. Formed by brominating C.Br.H (Reboul). Formed also by the action of Br and Al on CCl₄, C₂Cl₄, or C₂Cl₄ (Gustavson, J. R. 13, 287). Also from uncobromic acid, water, and Br at 130° (Delbrück, A. 165, 253). Prisms (from CS.); insol. alcohol. At 200"-210° it decomposes, without previous fusion, into Br and C.Br., Unlike the latter, it is not volatile with steam.

BROMO-ETHENYL-NAPHTHYLENE-DIA-MINE

[229°]. Formed by reduction of the acetyl derivative of (1:3:4)-brome nitro-(α)-naphthylnmine with SnCl. Small white needles. V. sol. alcohol and ether, insol. water. The ethenvl group is very stable. Salts.—B'ICL.—B'(H.SO.*: sparingly soluble needlee.—B'HNO.*: fine white sparingly soluble needlee (Prager, B. 18, 2160; cf. Meldola, C. J. 47, 505). BROMO-ETHOXY- v. Bromo-oxy-.

BROMO-ETHYL-ACETO-ACETIC ETHER v. BROMO-ACETO-ACETIC ETHER.

d. Bromo-ethyl-aceto-acetic-ether C, H, BrO, i.e. CH .CO.CH(CO.Et).CH .CH.Br. Heavy yellowish oil of comphor-like odour. Not distillable. Formed by dissolving trimethylene methyl-CH. CO.CH,

three times its weight of strongly cooled cone. IIBr (1.85 S.G.), allowing to stand 10 mins. at duct, containing CHBr, CH,Br, is treated with 15° and pouring into iced water. By boiling with HOI is is converted into scetyl propyl alco-hol OH_OO.CH_CH_CH_OH (v. p. 46) with formation of alcohol, CO, and HBr (Perkin, jun., a. Freer, C. J. 51, 833, B. 19, 2565).

BROMO-ETHYL ALCOHOL v. GLYCOL BROM-

Dibromo ethyl alcohol CHBr, CH,OH. (180°). S.G. 2 2.35. From bromo-ethylene and dilute HBrO (Demole, B. 9, 49). Reduces auunonincal AgNO. Alcoholio KOII gives bromo ethyleno Acetyl derivative CHBr. CH. OAc. S.G. 2 198.

BROMO . TRI . ETHYL . AMINE. bromide C.H.Br.NEt,Br. From NEt, and ethylene bromide. Alcoholio KOH fornus C2H2.NEt3Br (Hofmann, C. R. 49, 880).

p-BROMO-ETHYL-ANILINE C.H.Br.NHEt. From p-bromo-amiline and EtBr (Hofmann, A.

74, 145).

p-Bromo-di-ethyl-aniline C. H. Br. NEtz. [33°]. (270°). Needles or prieme. Formed by bromination of diethylanilino (Claue a. Howitz, B. 17,

p-BROMO-ETHYL-BENZENE C.H.BrEt [1:4]. (201°). S.G. 32 1'34. From p-ethylbenzene, Br, and 1 (Kekulé a. Thorpe, C. J. 22, 866). From C. II, Br., EtI, and Na (Aschenbrandt, A. 216, 222). Does not solidify at -20°. A mixture of o- and p- ethyl-benzenes is formed by the action of bromine in the dark upon ethyl-benzene, or by the action of bromine in presence of 3 p. c. of iodine upon ethyl-benzene in diffused daylight (Schramm, B. 18, 1272).

ω-Bromo-othyl-benzone C_aH, CH_aCH_bBr (c. 148°) nt 30 mm. S.G. ²² 1 311. Formed by direct combination of styrene with IIBr (Bernthsen a. Bender, B. 15, 1983). Formed also by treating a mixture of benzene and bromo-othylene with Al₂Cl₆ (Hanriot a. Guilbert, C. R. 98,

525).

a.Bromo.ethyl.benzene C.H., CIIBr.CII, (37°) at 17 mm. (c. 150') at 500 mm. From Br and ethyl-benzene at 140° (Radziszewski, B. 6, 492; Berthelot, C. R. 67, 328; Anschütz, A. 235, 328). Formed also by passing HBr into cooled phenyl-methyl curbinol (Engler a. Bethge, B. 7, 1125). Partially decomposed by distillation into etyrene and HBr.

w.a.Di.bromo.ethyl.benzens

Ph.CHBr.CH,Br. [74°]. (140°) at 15 mm. Styrene di-bromide.

Preparation .- 1. From etyrene (10 g.), other (25 g.) and bromine (17 g.) (Blyth a. Hofmann, A, 53, 306; Zineke, A. 216, 288). -2. From hot ethyl-benzene und Br (Radziszewski, B. 6, 493; Friedel a. Balsohn, Bl. (2) 35, 55).

Properties. - White plates or needles (from 80 p.c. alcohol). V. e. sol. other, benzene and glacial HOAc, v. col. alcohol or benzoline.

Reactions .- 1. KOAc and alcohol at 160' gives chicily \$\textit{B}\$-brono styred (150^2-160') at \$\text{C}_1\text{B}\$rOs (40'-45) (40 \$\phi\$ (Dernols, B. 9, 49),—75 mm.).—2. KOAc and glacial 110Ac gives 7. Dry oxygen at 23' has no action.—8. ICI chicily the diacetate of phenyl-plycol, forms CliBrl.CH_Cl and a less quantity of Ph.CH_(OAc)CH_(OAc).—3. Alcoholic KOH or Cll l.CliBrC (tlerry, C. B. 98, 680).—9. With water at 190° give a-bromo-styrene (Glaser, A. 154, 154).—4. Givee PhCH(OH).CH₂(OH) by treatment with AgOAc, alcohol and AgOBz, or aqueous K₂CO₃. AgOBz in presence of toluenc forms Ph.CH(OBz). CH₂(OBz).—5. Benzone and Al.Ol. give s-di-phenyl-ethane (Anschütz, A.

Tri-brome-ethyl-benzene O.H. CHBr.CHBr. [38°]. From a-bromo-styrene and Br (Fittig a. Binder, A. 195, 142). Acted upon by bensene and Al₂Cl₄ in presence of CS₃, it is converted into s-tetra-phenyl-ethane [209°] (A.).

Tetra-bromo-ethyl-bensene C₃H₃Br₄. From

di bromo ethyl benzene and Br (Kinnioutt a.

Palmer, Am. 5, 387).

Penta-bromo-ethyl-benzene Call Br. From ethyl-benzene, Br, and Al Br, Gustavsen, Bl. [2] 30, 22).

Di-bremo-di ethyl-bonzen C,II,(C,H,Br), (200°-230°) at 30 mm. From bromo ethyleno. benzene, and Al. Cl. (Hauriet a. Guilbert, C. R. 98, 525).

Bromo-tetra-ethyl-benzene Call(Call,) Br (281 nncor.). Heavy liquid (Gulle, B. 16, 1745).

Di-bromo-tetra ethyl bonzeno C. (C.II.) Br., [75°]. (above 330°). Prisms. V. sol. alcohol (Galle, B. 16, 1745).

BROMO.ETHYL.BROMIDE v. Dr. Bnomo.

BROMO - ETHYL BROMO - ACETATE v. BRONO-ACETIC ACID.

BROMO-ETHYLENE C.II.Br i.e. CHBr:CII. Vinul bromide. (16"). S.G. 14 1-5167 (Anschütz). Formed by the action of alcoholic KOII upon either di-bromo-ethuno (Regnault, A. Ch. [2] 358; Beilstein, J. 1861, 509; Glöckner, A. Suppl. 7, 109; Semenoff, J. 1864, 480). Formed also from acctylene and HBr (Rehoul, C. R. 74, 947). Gas or liquid; when kept in a sealed tube and exposed to snalight it changes to an amorphous polymeride, insel. water, alcohol, and ether; S.G. 2075. This substance carbonises when heated (Hofmann, C. J. 13, 68; Baumann, A. 163, 308); it is not affected by boiling alcoholic KOH. Polymerisation is arrested by the presence of MeI or Etl, but not by chlorinated or brominated hydrocarbons; a little I steps polymerisation of the liquid, but not of the gas. An line arrests, but SO, promotes, tho change (Lwow, Bl. [2] 35, 169).

Reactions .-- 1. Split up into HBr and acotylene by alcoholic KOH, NaOEt, NaOC, H111 or ammoniacal AgNO₃ (Sawitsch, Bl. 1861, 7; A.119, 185; Miasnikoff, Bl. 1861, 12; A. 118, 330). --2. Conc. HBr at 6° forms s di-bromo-ethane; a more dilute acid gives u-di-brome ethane (Rebool, A. 155, 29, 212) .- 3. Funning HCl at 100° forms CH₂CHBrCl.—4. Conc. HI at 4° glvos CH₂CHPrI; at 100° it forms also Origi.CH₂Br. (R.). - 5. Aqueons solutions of metallio salts either have no action or split it up into acetylene and HBr (Kutscheroff, B. 14, 1532; Linnemann, A. 1(3, 317; Saytzeff a. Glinsky, Z. [2] 3, 675). 6. duc ressive treatment with cone. H2SO, and water forms crotonio aldohyde (Zincke, A. 191, 379). 7. Aqueoue BrOH at 0° gives CilBr. Cif.Oll, CllBr. CH.Br. and benzene and Al Cl, it produces styrene, ethylbenzene, u-di-phenyl ethane, and di-methylanthracene dihydride (Anschotz, A. 235, 331). If elevation of temperature be avoided and the Al_Cl. be clowly added the products are C.H. C.H.Br and C.H. (C.H.Br). (Hanriot a. Guilbert, C. R. 98, 525).

s-Di.brome-ethylene OHBr:OHBr. Acetylene dibromide. (110°-111°). S.G. 175 2.2714 (Ansoliütz, A. 221, 141); 2 2.268 (Sabanéeff, B. 9, 1441; Plimpton, C. J. 39, 536); 12 2.223 (S.). V.D. 6.47 (calc. 6.44).—Formed by mixing acetylenc tetrabrosaide (100 g.) with alcohol (20 g.) and adding zinc dust slowly, with cooling (A.; cf. Sabanéeff, A. 216, 252).

Properties.—Oil; does not polymerise.
Reactions.—1. Heated for soveral days with 50 pts. of water at 135°, it is not affected .-2. Heated with diluto K.CO3, bromo-acctylene is formed, which is spontaneously inflammable. Alcoholic KQII and aqueous KCy also form brome-acetylene.—3. Heated with dry KOAo at 160° for two days it forms CHBr:CH(OAc), the acetate of bromo-vinyl alcohol (Sabanéeff, Bl. [2] 41, 253).-4. Heated with AgOAc and a little 110Ac at 100° it forms a compound C2H2Br2AgOAo. This is decomposed by 11Cl with evolution of acetylene, -5. Combines with AgNO .- 6. With alcoholic KCN it forms a compound which, on saponification, gives an acid C₁H₂O₄ (163°-168°). Its silver salt is C₄H₄Ag₅O₄ (S.).—7. Alcoholic KOPh gives CHBr.CHOPh. (c. 223°); S.G. 15 1-185, 8. With alcoholio NMe, at 120° it forms NMe, Br, NMe, HBr, NMe_II_Br, and carbonaceous hodies:

2NMc₃ + C₂ll₁Br₂ = 2NMc₃llBr + C₂ (Plimpton, C. J. 39, 536). - 9. With NEt₄ it nets similarly. — 10. Acts upon beuzene in presence of Al₂Cl₄ forming CHBr₂-CH₂Br₄ unthracene, and Cli₂Ph.CH₂Ph (Anschütz, A.

u-Di-bromo othyleno CBr.: Cll., Acetylidene dibromide. (92°) at 751 mm. in CO. S.G. 2:1780 (Anschütz). From CIBr., CH.Br by treatment with alooholic KOH, NaOEt, sodium, or solid KOH (Cahours, C. R. 31, 293; Fontaine, C. R. 70, 1361; Sawitsch, A. 119, 183; Reboul, A. 124, 270; Tawildaroff, A. 176, 22; Michael, Am. 5, 192). From Chr. CH.Br by bolling with alcohol and KOAc (Demole, Bl. [2] 29, 205), or by treatment with zine and alcohol (Sabanceff, A. 216, 255). Also from CH Cl.CHBr, and alcoholic KOII (Henry, Bl. [2] 42, 262).

Properties .- Pungent liquid, attacking the eyes. Readily absorbs oxygen, changing to broma-acetyl bromide. Polymerises with groat

ease, becoming solid.

React 1 : ... 11BrO forms CBr, CO.CH, CBr, [90°]. S.G. ² 2°88 (Demore, B. 31, 1710).—2. Benzeno and Al₄Cl₄ give u-di-phenyl-ethyleno (Auschütz, A. 235, 158).

Tri-bromo-ethylene CHBr:CBr. (163°-164°). S.G. 205 2.708 (S. a. D.); 2 2.69 (Demole, B. 11, 318). From di-bromo-ethylene bromide and

alcoholio KOH (Lennox, A. 122, 125). Preparation. -- Acetyleno tetra-bromido (1

mol.) is dissolved in twice its weight of alcohole and (somewhat more than 2 mols, of) KOAe and Na,CO, (1 mol.) are added; the mixture is heatod on a water bath 21 hours with inverted condenser (Sabandeff a. Dworkowitsch, A. 216, 280; cf. Sabanécff, A. 178, 122; Bl. (2) 29, 207).

Reactions.—1. Alcoholic KOH or Zn and

110Et give a mixture of C.H., and C.HBr.— 2. Alcoholio KOPh at 100° forms phenyl di-bromo-vinyl exide.—3. Alcoholic KOPh at 170° forms the phenyl derivative of glyoxylic seid, Pho. CH, CO, H. 4. Reacts upon bensene in presence of Al₂Cl, producing u-di-phenyl-ethylone and tri-phenyl-methane (Anschütz, A. 285, 336).—5. Absorbs oxygen, becoming CHBr. CO.Br (Demole a. Dün, B. 11, 1302).

Tri-bromo-ethylene (C.HBr.). [174°]. A by-product in formation of C.H.Br. from acetyl-

ono (Sabanéeff, A. 178, 114).

Tetra-hrome-ethylene C.Br., [542]. (2159). Formation.—1. By the action of Br on alcohol or ether (Löwig, A. 3, 292).—2. From C.HBr, and alcoholio KOII (Lennex, A. 122, 126).—3. From CH.Cl. and 1Br, (Höland, A. 210, 234).—4. From CBr, by heating at 220° (11.) .- 5. From di-bromo-succinic acid, Br, and water at 190° (Bourgoin, B. 7, 1644).

Properties .- Plates; volatile with steam; no. affected by oxygen even at 216° (D.).

BROMO-ETHYL-ETHER v. BROMO-ETHYL

BROMO-ETHYL-KAIRINE v. Ethyl ether of Bromo-(B. 4)-oxy-(B. 4)-ETHYL-QUINGLINE TE-TRATIYUNIDE,

γ-BROMO-ETHYLºMALONIC ACID

C.H.BrO, i.e. Br.CH, CH, CH(CO, H), [1160]. From vinaconio (tri-mcthylene di carboxylio) acid and HBr (Röder, A. 227, 19; Perkin, jun., C. J. 47, 814). Sl. sol. benzene, CS, or light potroleum, sol. chloroform. Boiled with water it gives the lactone of y-oxy-ethyl-malonic acid (q. v.).

Bromo-ethyl-malonic acid

CII, CHBr.CH(CO.H), (?). [141°]. From orotaconio acid C,H,O, and HBr (Claus, A. 191, 79).
TRIBROMO. (a) - ETHYL: NAPHTHALENE

C.H.Br.C.H. [127°]. Slender needles. Prepared by the action of bromine on (a) ethylnaphthaleno (Carnelutti, B. 13, 1672).

BROMO ETHYL (B) NAPHTHYL ETHER v. (B)-NAPHTHOL.

€wa.BROMO.DI.ETHYL OXIDE

CH.Br.CH.OEt. (128). S.G. 2 1:371. V.D. 5:29 (calc. 5:28). From the iodo-compound and Br (Henry, C. R. 100, 1007).

Di-ω-bromo-di-ethyl oxido CH.Br.CHBr.OEt. From Br and vinyl ethyl oxide. Vory unstable liquid. NaOEt gives CH_Br.CH(OEt)₂ (Wislicenus, A. 192, 111).

Tetra hromo di ethyl exide C4H,Br.O. fuming liquid obtained by saturating ethylidene oxychloride with Brat 120° (Kessel, B. 10, 1667). Octo-bromo-di-ethyl oxide C, H, Br, O. (c. 1346) at 160 mm. By heating ethylideno oxychlorido with Br for 10 hours from 100°-210° (Kessel, B. 10, 1667). Furning oil.

Eso-cro. DI. BROMO o.ETHYL . PHENOL C. H. Br(C. 11, Br)OH. From o ethyl-phenol and Br in the cold. Decomposed by distillation into HBr and C.H.Br(C.11.)OH (Suida a. Plohn, M.

Tri-hromo (a) ethyl phenol C. HBr. Et. OH. [55°]. Formed, togother with the following hody, by treating (a)-cthyl phonol with excess of Br in the cold (Fittig a. Kiesow, A. 156, 251).

Eso-exo- Tri-bromo-ethyl phenol C.11,Br(OH).C11Br.CH.Br. [108]. Long white needles. Obtained by the action of alcoholic KOII upon Call, Br(OH). CHBr. CliBr. CO.H, the product of the action of bromine upon p-cou-

Acetyl derivative C.H.Br(OAc).C.H.Br.: | one. [B4°]; needles (Eigel, B. 20, 2535). Tetra-bromo-ethyl phenol C_sBr_sEt.OH [106°]

BROMO- TETRA - ETHYL - PHOSPHONIUM BROMIDE CH.Br.CH., PEt, Br. [235°]. From PEt, and ethylene bromide in the cold (Hofmann, Pr. 9, 287; A. Suppl. 1, 154). Rhombic dodeca-

Reactions.-1. Moist silver oxide gives CH. (OH).CH.PEt.OH (difference from corresponding derivatives of AsEt, and NEt, which give vinyl base) .- 2. With silver acetate and water at 100° it gives acctute of the vinyl base C₂H₃PEt₃.OAc.-3. Zine and H₂SO₄ give tetra-ethyl-phosphonium bromide. - 1. Alcoholic potash has no effect. -5. Combines with PEt, AsEt, and NH, forming di acid bases.

Salts .- (C.H.Br.PEt,Cl),PtCl, Palo orango monoclinic prisms, sl. sol. cold, v. sol. hot, water. $-(C_2H_4Br.PEt_3Cl)AuCl_3.-C_2H_4Br.PEt_4I.$

Hydroxide. - C.H. Br.PEt.Off. From the sulphate by the action of baryta. Unstable, rapidly changing to C.H.OH.PEt,OH. BROMO ETHYL QUINOLINE

C.H. (C.H.Br)N. The hydrobromide is formed hy heating quinoline with ethylene bromide. --B'HBr: thick needles .- (B',H,Cl2)PtCl4: orangeyellow needles (Berend, B. 14, 1349).

DI-BROMO DI ETHYL SULPHATE

(CH,Br.CH,)2SO4. An oil formed by warming Ag₂SO₄ with benzene and ethylene bromide (Beilstein a. Wiegand, B. 15, 1369).

Brome-ethyl-sulphurie acid

CH_Br.CH_O.SO_H. From ethylene bromide and SO_1 (Wroblewsky, Z. 1868, 563). Ba.A.,... PbA'2 3aq, scales. An isomeric acid appears to be formed by heating ethylene bromide with Ag₂SO₄ and water (B. a. W.).

BROMO ETHYL THIOPHENE

C,SH,(C,H,)Br. (195° uncor.). Oil. Formed by shaking ethyl-thiophene with bromine water (Demuth, B. 19, 684).

Di-brome (B) ethyl-thiophene C.SIIBr. (C.II.). Oil. Formed by adding 2 mols, of bromine to 1 mol. of (B)-ethyl-thiophene dissolved in acctic

acid (Bonz, B. 18, 550). Tri-bromo (β)-ethyl-thiopheno C, SBr, (C, II,). [108°]. White plates. SI. sol. cold alcohol and ethor. Formed by complete bromination of

(B)-ethyl-thiopheno (Bonz, B. 19, 519). BROMO-o-ETHYL TOLUENE C. II, McEtBr

[1:2:4]. (221°). Formed by bromination of oethyl-toluene in presence of Fe.Br_a. Liquid. By HNO₃ (1·1) at 200° it is oxidised to p-bromoo-toluie neid [118] (Claus a. Pieszcek, B. 19, 3088).

Bromo-p-othyl-tolueno Call MeBrEt [1:2:4]. From p-ethyl toluene and Br. Oxidised to bromo p-toluic acid (Morse a. Remsen, B. 11, 224).

wa. Di. bromo.m.ethyl-teluene C.H., CHBr. Cil.Br. 15°]. Formed by the combination of m-tolyl-ethylens (m ni-thylstyrene) with bromine. Colourless crystals (Müller, B. 20, 1216).

Tri.brome di ethyl teluene C.Br.MeEt .. [206°]. From (1, 3, 5) di ethyl-to-uene (Jacobsen, B. 7, 1435).

TRI.BROMO ETHYL-XYLENE C.Br. EtMe. [8:5:6:1:2:4]. Tri-bromo di-methyl sthyl benzens. [91°]. From (Anschütz, A. 235, 824). From ethyl-m-zyleno (187°)

BROMO EUGENOL C.H.Br(CMe)(CH)C.H.,
Methyl ether C.H.Br(OMe), (190°) at
20 mm. S.G. 2 1396. From the dibromide, hot alcohol, and Zn (Wassermann, C. R. 88, 1206), Di-bromide C.H.Br.(OMc)... Dimethyl ether of di-bromo-di-oxy-propyl-benzene. (78°). From Br and a well-cooled solution of methyl-ougenol in other. Silky needles.

Ethyl ether Call Br(OMc)(OEt). [48°]. Prepared by the action of alcohol and Zn on its dibromide (Wassermann, A. 179, 385). Dibromide C, H.Br. (OMe) (OEt). - [80]. From

ethyl-eugenol and Br. Di-brome ougonel CoHBr. (OMc) (OH) (C1H2).

Collistening hexagonal prisms. V. sol. [59°]. Glistening hexagonal prisms. alcohol. Formed by boiling an alcoholic solution of the di-bromide with zine dust (Chasanowitz a. Hell, B. 18, 824).

Di-bromide C. HBr. (OMe) (OH) (C.H Br.). [119°]. Glistening quadratic or trimetric tables. Sparingly soluble in other and cold alcohol. Formed by bromination of eugenol.

BROMO-FLUORENE

 $C_{12}\Pi_{z}Br$ i.e. $< C_{z}\Pi_{z}Br$ $> CH_{z}$. [102°]. fluorene, CHCl₁ and Br in the cold (Hodgkiuson a. Matthews, C. J. 43, 165). Needles (from alcohol). V. sol. CHCl₃. Oxidises to bromedi-phenylene ketone.

(a)-Di-breme-fluerene

CaH Br i.e. CaH Br CH. Di-bromo-diphenylene-methane. [165"] (Barbier, A. Ch. [5] 7, 479; Hodgkinson a. Matthews, C. J. 43, 161). Got by adding bromine to a solution of fluoreno in CHCl. Monoclinic crystals, $a:b:c=1.167(1(1.065); \beta \approx 77^{\circ}.52')$ (Arzumi, Z. Eryst. 1, 624). Sol. boiling alcohol. Turned yellowish by light. CrO, gives di-bromo diplanylene ketone.

Supphonic acid Call, Br. SO. H. [1420]. Formed by sulphonation with CISO II in CHCl.

-BaA

(β)-Di-breme-fluerene $C_{13}H_{s}Br_{2}$. Formed together with the preceding (Fittig a. Schmitz, A. 193, 134). Monoclinic crystals; a:b:c=563:1:697. $\beta=78^{\circ}$ 21' (A.). Readily changes into two isomerio modifications (7) and (8) (Lehmann, Z. Kryst. 1, 626). Tri-brome-finorene

 $C_{12}H_{2}Br_{2}$ i.e. $< \frac{C_{01}I_{3}Br}{C_{0}H_{3}Br} > CHBr$. [1622]. From fluorene (1 mol.) in CS₂ and Br (3 mols.). Oxidised by CrO₂ to (β)-di-bromo-diphenylene ketone (B.).

DI-BROMO-FLUORESCEIN C20 H10 Br2O1 [260° 270°]. From fluorescen (1 mol.) and Br (2 mols.) in HOAe (Uneyer, A. 183, 1). Reddishbrown needles with g.con reflex. Dyes wool and silk salmon-pink.

Di-acetyl derivative Call AcaBraO. [210°].

Tetra-bromo fluoresceïn

 $\mathbf{C}_{zo}\mathbf{H}_{a}\mathbf{Br}_{4}\mathbf{O}_{\bullet}\text{ i.e. }\mathbf{O} < \begin{matrix} \mathbf{C}_{a}\mathbf{H}\mathbf{Br}_{v}(\mathbf{OH}) \\ \mathbf{C}_{o}\mathbf{H}\mathbf{Br}_{4}(\mathbf{OH}) \end{matrix} > \mathbf{C} < \begin{matrix} \mathbf{C}_{a}\mathbf{H}_{4} \\ \mathbf{O} \end{matrix} > \mathbf{CO}_{\bullet}$ Eosin. Formed by adding Br to a solution of fluorescein in HOAc. It is purified by conversion into the K salt (Baeyer, A. 183, 38). Prepared by dissolving fluorescein (1 mol.) in NaOHAq, adding a solution of Br (4 mols.) in NaOIIAq, and acidifying. Orange needles (containing HOEt) (from alcohol), or fissh-coloured crystals C20H, Br.O. (from dilute alcohol containing HCl). Tetra bromo fluorescein is v. sl. sol. ing HCl). water and benzene; its alkaline solutions are pink and show strong yellow fluorescence, they dye wool and silk pink. Zinc-dust and NaOll reduce it to a lenco-compound, which is reoxidised by air. Potash fusion forms di-bromoresorcin and di-bromo resorcin-phthalein. Conc. forms C H Br O ... I'Cl, forms C, Il, Cl, Br, O, Sodium maalgam forms fluorescein. Warming with cone. KOll gives a deep blue solition sheare HCl pps. unstable

 $O < \stackrel{C_u H \operatorname{Br}_u(OH)}{C_u H \operatorname{Br}_u(OH)} > C(OH) \cdot C_u H_u \cdot CO \cdot H.$

Salts. - K.(C., H.Br.O.) 5aq. S. 50. - K.A"HOEL. - (NII.) A". - InA" 2aq. - CaA" kaq. -Ag.A".—(IIÒPb).A".

Methyl other Coll, MeBr.O. Methyl

erythrin.

Ethyl ether Czall, EtBr.O. Erythin. Spirit-soluble cosin. From K.A", KEtSO., and alcohol at 150°. From thoreseein, boiling alcohol and Br. Red crystals (from alcohol). Formed, together with a colourless ethyl-cosin, by heating silver cosin with EtI and alcohol at 100°. - KC, II, EtBr.O, aq: dyes a more violet shade than cosin.

Disethyl other Czoll Et Br.Os.

Ag A" and Etl.

Acetyl derivative $C_{20}H_6Ac_2Br_4O_4$ (?).

| BROMOFORM CHBr., Tri-bromo-methane. | Mol. w. 253. [8°]. (151°). | S.G. 4° 2°8341 (T.); | \(\) 2°9045 (Perkin, C. J. 45, 553); \(\) 2°8842 (P.). | C.E. (0° 10) \(\) 000944 ; \(\) (0° 100) \(\) 0010116. | S.V. 103°53 (Thorpe, C. J. 37, 203). | M.M. 11·626 at 17·9° (P.). | V.D. 8·63 (calc. 8·75) 11 626 at 17 9 (P.). (Cahours, A. Ch. [3] 19, 484).

Occurrence.- In erude bromine (Hermann,

A. 95, 211; Dyson, C. J. 43, 36).

Formation .- 1. By the simultaneous action of Br and KOH, or of 'bromide of lime,' on alcohol or acctone, or by decomposing bromal with alkalis (Löwig, A. 3, 295; Dumas, A. Ch. [2] 56, 120; Günther, Ar. Ih. [3] 25, 373).

 From CH Cl₂ mul IBr, (Höhmd, A. 210, 236).
 Reactions. — 1. Alcoholic KOH forms CO (Avols.) and ell-plene (I vol.) but no formate (Long, 37 J. 3. 23). -2. Br in presence of dilute KOII in sunlight forms CBr, (Illabermann, B. 6, 549). -3. Reduced to CH, by KI, water, and Zn or Cn (Berthelot, A. Ch. [3] 51, 48) or by the copper zine couple (Gladstone B. Tribe, C. J. 28, 510).

BROMO - FUMARIC ACID C.HBr(CO.H).

[1785].

Formation .- 1. From iso-di-bromo succinic acid by heating at 180° or by boiling with water (Kekulé, A. Suppl. 2, 91; A. 130, 1).—2. From di-bromo-succinic acid and water at 140 (Bandrowski, B. 12, 315) .- 3. By distolving acetylane-di-earboxylic acid in strong aqueous HBr (Bandrewski, B. 15, 2697). -4. From (βδ)dibromopyromucic acid and from (8) bromopyromucic acid by dilute IINO, (Hill a. Sanger, A. 232, 82, 64). 5. From bromo-maleic acid and cold fuming HBr (Fittig a. Petri, A. 195, 67)

Properties.-Lamine; v. e. sol. water, v. sol. alcohol and ether. At 200 it changes to brome maleic acid or its anhydride. Sodiumamalgam forms fumaric acid. Br gives the samo tri-bromo-succinic acid as with bromomaleïc acid. Furning HBr combines slowly in the cold; at 100° it quickly forms iso di-bromosuccinic acid. With its equivalent of aniline it unites immediately to form the acid aniline salt. This does not give an anilide on standing for weeks in contact with cold water. On boiling its aqueous solution the same substance is obtained as on heating aniline brome-maleate, viz. C_{1e}H₁₂N₂O₂. [230°] (Michael, Am. 9, 180). Salts. - Ag₂A". - PbA" 2aq. - A"H(NH₂Ph):

[154] (Michael, B. 19, 1373).

Dimethyl ether A"Meg. [30] (Anschütz, B. 12, 2284).

Di bromo fumarie acid

CO.H.CBr:CBr.CO.H. [220]. Colonricsservstals. Prepared by the addition of bromine to acetylenedicarboxylic acid. On distillation it is converted into dibromomaleïe acid [108°].

Salts. -AgrA" ang - PbA" (Bandrowski, B. I2, 2213).

(β)-BROMO-FURFURANEC, H, BrO. (103°). From the corresponding bromo-pyromucio acid by distilling with lime (Canzoneri a. Olivori, G.

17, 42). Ileavy oil. (a) Di-bromo-furfurane Call Br.O i.c.

CBr. O. [10°]. (63 ') at 15 mm.; (165°) 110 - CBr

at 761 mm. Formed by adding bromine to an alkaline solution of (δ) -bromo-pyromucic acid [184]. On oxidation it gives fumaric and maleic acids.

Tetra-bromide G,H,Br,O: [1110]; by long boiling with water it yields bromo-fumuric and bromo-maleje acids (Ilill a. Hartshorn, B. 16, 1132; B. 18, 448; A. 232, 53).

(β)-Di-bromo-furfuranc CBc:CH >0. (166°).

Formed by distilling di-bromo-pyromucic acid (1 pt.) with Ca(OII)2 (2 pts.) (Canzoneri a. Oliveri, G. 15, 116).

Tetra-bromo-furfurans C.Br.O [65]. From (βγ)-di-bronio-pyromucio acid, or from tribrome-pyromucic acid, water and bromine va-pour. Formed also by the action of alcoholic KOII on di-bromo-furfurane tetrabromide (Hill a. Sanger, A. 232, 86, 96; B. 16, 1132; 17, I760).

Di. bromide CaBraO. [1237]. Six-sided plates. V. sol. ether, m. sol. alcohol and benzene. By boiling with water it yields di-brouto-maleio acid (Hill a. Hartshern, B. 18, 450).

BROMO-FURIL P. FURIL

BROMO-GALLIC ACID v. BROMO-TRI-OXY. BENZOIC ACID.

TRI-BROMO-GUAIACOL C.H. Br.O. i.e. C.H.Fr. (OMc) (OH). [102°]. From guaiacol and Br (Tieuiann a. Koppe, B. 14, 2017).

BROMO.GUANIDINE CH, BrN. guanidino carbonats and Br (Kamenski, B. 11, 1600). Needles; detonates just above 100°.

BROMO GUANINE C, H, N, OBr. From guanine and bromins, crystallised from water. White powder. Sl. sol. boiling water, insol. cold water, alcohol or ether .- B'HOL Prisms. Converted by NaNO, into bromo-xanthine (Fischer a. Reese, A. 221, 341).

TRI-BROMO-HEMIMELLITHOL TRI-BROMO-TRI-METHYL-BENZENE-(1:2:3:4:5:6).

BROMO-HEPTANE v. HEFTYL BROMIDE.

Dl-bromo-heptane $C_rH_{11}Br_{2r}$ Heptylene bromide. S.G. 153 I-515. From lieptane of paraffin oil. Decomposes at 150° (Thorpo a. Young, A. 165, 12).

Di-bromo-heptane C.H., Br. (211°). heptane in the oil of Pinus subiniana (Venable,

A. C. J. 4, 22).

Di-bromo heptane CMeg. CMeBr. CH.Br. From penta-methyl-cthyl alcohol and PBr. Easily fusible solid (Kaschirski, C. C. 1881, 278).

Di-bromo-heptano CH, CH, CH, CH, CH, CH .. CHBr.. Heptylidene bromide. From quanthol and PCl, Br. (Bruylants, B. 8, 409).

Hexa-bromo-heptane C.H. Br. From heptonene and Br. Oil (Saytzell, .f. 185, 144).

a-BROMO-HEPTOIC ACID CallerBrO. CH, CH, CH, CH, CH, CHBr.CO.H. (2507), From Br and heptoic acid (Cahours, A. Suppl. 2, 83; Helms, B. 8, 1168; Hell a. Schüle, B. 18, 625). Ethyl ether EtA', (c. 225). S.G. 201211.

TETRA-BROMO-HEPTYL ALCOHOL C,H, Br,O i.e. (CH Br,CHBr,CH) CH.OH. From di-allyl-carbinol and Br (M. Saytzeff, A. 185, 135).

Acelyl derivative C. H. Br. OAc. verted by AgOAc into C, H₁₁O₁OAc)₃, S.G. ^a₆ 1480, where the state of the whence baryta forms a syrup C/H₀O(OH)₃ (Dieff, J. pr. [2] 35, 17).

BROMO-HEPTYLENE C.H₁₃Br. (158°). From the heptylene bromide of Venable.

Bromo-heptylene C.H., Dr. (1654). From the heptylene bromide of Bruylants.

BROMO-HEXADECANE P. CETTL BROWNER. Di-brome-bexadecane C_mH_nBr₂. Cetene bromide.

[13]). Colourless crystalline solid. Formed by addition of Br, to cetene (Kraff, B. 17, 1373).

BROME TEXANE C. HEXYL BROMHOR.

Di-bromo-hexane C.H. Hr. i.e. CH₂CH₂CH₂CHBr.CHBr.CHBr.CH₂ (196) at 710 mm. S.G. P 15809. From the corresponding hexylene (Erlenuwyer a, Wanklyn, A. 135, 141; cf. Hecht a. Sto. 14, A. 172, 69; Hecht, B. 11, 1423).

Di-bromo-hexane Mc CBr CBr Me., (170 ± (K.); [1402] (E.). From Me,C:CMe, and Br. Needles (from ether). Converted by heating with water and PbO at 150' into pinacolin (Pawlow, Ar 196, 121; Effekoff, J. R. 10, 220; Kaschursky, J. R. 13, 81).

Di bromo hexanc Me C.CHBr.CH Br. Prom Me, C.CH: CH, Crystadine (Friedd a Silva, Bl. [2] 19, 289).

Di-bromo-hexane C, H₁₂Br₂, (211°). From hexane of petroleum (P-louze a. Cahours, A. 121,

DistlytTetra-bromo hoxane C.H.Br. tetrabromide. [63°]. From diallyl and Br (Wagner a. Toilens, B. 6, 588).

Tetra bromo hexano C. H. Br. -1429]. From iodo-hexylene and Br (Bouchardat, Z. 1871,

Tetra-brome hexane C.H., Br., From hoxis sorbic acid to pone derived from mannito (Hecht, B. 11, 1054). B. 15, 619).

Tetra-bromo-hexane C.H.Br. From hexinene from di-methyl-allyl-carbinyl chloride (Saytzeff, B. 11, 2152).

Tetra-bromo-hexane O.H., Br., [112], (318° cor.). From hexineno derived from coal-tar (Schorlemmer, A. 139, 250).

Hexa-bromo-hexane Call, Br. From diallylene (Henry, Bn. 1, 199).

Hexa-bromo hexane C. H. Br. [77°]. From di-fromo-diallyl (Henry, $\hat{B}, \hat{7}, \hat{23}$).

Hexa-bromo-hexane ('11,13. 152]. From see hexyl iodide and Br at 130 (Merz a. Weith, B. 11, 2250).

Hexa-bromo hexane C.H.Br., Tand Br at 125' (Wahl, B 70, 1254). Trem bexana

Octo brome hoxane Call Br. From hexane and Br (W.).

Octo-bromo-hexano C.D.Br., [135"]. From sec-hexyl iodide and Br at 130 (M. a. W.).

Octo bromo hexano Call Br. Dipropargyl octobromide (111 'j (Hemy, B. 7, 21).

BROMO-HEXENOIC ACID C. H. Br.O. Dibromo-hydrosorbic acid. [95°]. From sorbic acid and Br. Lamine (Fittig a. Kachel, A. 168, 287).

DI-BROMO HEXINENE C. H. Br. Di bromodiallyl, (210). S.G. 1 1 656. From diallyl-tetrabromide and solid KOH (Henry, J. pr. [2] 8, 57).

Totra-bromo-hexinene C.H.Br., Proparayll tetralremide, S.G. 2 2:161, Liquid (Henry, B. Propargyl 6, 959).

a-BROMO-n HEXOIC ACID

CII. CH .CH .CH ..CHBr.CO.H. Bromo-caproic acid. (210). From caproic acid and Br (Calionra, A. Suppl. 2, 78).

Ethyl other EtA' (205 ' 210°) (Hell, B. 17,

γ-Bromo n hexoic acid

CH_CH_CHBr.CH_CH_CO_H. From hydrosorbie, or iso hydrosorbie, need and HBr. Oil. Sodium-amalgam reduces it to n-hexoic wad. Boiling water converts it into hydrosorbic and oxy-hexpic fields (Fittig, A, 200, 42; Hielt, B_{\bullet} 15, 618).

y-Bromo-iso hexoic a. id

Mc CBr.CH,,CH,,CO,H.

Ethyl ether A'Et. Formed by saturating an absolute alcoholic solution of isocaprolactone with HBr. By distillation it is decomposed into the original Inctone and EtBr (Bredt, B. 19, 514).

Bromo-nexoic acid C.H., Brog. [86"]. From iso-pyroterebic acid and IIBv., Prisma. AgA' (Lagermack a. Eltekoff, J. R. 11, R. 9)

Brome hexoic acid Call BrO. [25"]. From ethyl-crotonic acid and conc. HBrAq. Sodiumanalgam forms hydro-ethyl-crotonic neid. Aqueous Na_sCO_sAq even sat 0 forios ainylene, NaBr, and CO...

oß.Di-bromo-iso-hexoic acid

Pr.CHBr.CHBr.CO.H [99]; From pyrotoro-bic acid and Br.(W. 11, Williams, B. 6, 1095; Geisler, A. 208, 46). Ladge crystals (from CS₂).

Di-brome-hexoio acid $C_x\Pi_{10}{\rm Br}_xO_x$. [68°]. From sorbic acid and funning HBr. Boiling water or alkalis produce sorbic acid, and other bodies (cf. Hjelt, B. 15, 620).

Di-bromo-hexoic acid Callin Br. Og. hydrosorbic acid and Br in CS, hiquid; decomposed by boiling water, giving oxy-hydrosorble acid (Fittig, A. 161, 314; 200, 46; Hjelt, Di-brome-hexoic acid C.H.Br.O. From iso-sorbic acid and HBr. Oil (L. a. E.).

Di-bromo-hexoio acid CHEtBr.CMeBr.CO.H. From methyl-ethyl-acrylio acid and [98°]. From methyl-ethyl-acrylio acid and Br. Monoelinic crystals, a:b:c=96:1:1:53.8=94° 36′. Water at 100° forms brome amylene CHEt:CMeBr, methyl othyl-acrylic acid, di-oxyhexoic neid, and methyl ethyl ketone (Lieben, a. Zeisel, M. 4, 78).

Di-bromo-hexoie acid C.II, Br.O. From ethyl-crotonic acid and Br. Decomposed 1 by cold Na,CO,Aq into bromo-amylene, NaBr,

and CO, (Fittig, A. 200, 35).

Tetra-bronio-hoxoie acid Callabr.O. [183]. From sorbic acid and Br (Fittig, A. 161, 323; 168, 277; 200, 58). Monoclinic crystals (from alcohol). More stable than the preceding acids, not being attacked by water at 100 '. - NuA' 2aq. -CaA', 7aq. BaA', Paq.

DI BROMO-HEXOIC ALDEHYDE C. H. Br.O. GH, CH, CHBr, CMeBr, CTIO. Di-bromomethyl-propyl-acetic aldehyde. From Br and cold methyl-ethyl-acrolem. Pungent oil. -(C_sH_{to}Br_sO)NnHSO_s 3aq (Lieben n. Zeisel, M. 4, 19).

BROMO-HEXONENE C.H.Br. Bromo diallylene. (150°). From di-bromo di-allyl and KOII. Pps. ummediacal AgNO, and enprous chloride (Henry, B. 14, 400).

Octo-bromo-hexoneno CoBrs. From sec-hexyl iodide and Br at 140°. Prisms. At 200° it splits up iido Br and hexu-bromo-benzene (Merz a. Weith, B. 11, 2217).

BROMO HEXYL ALCOHOL Caller BrO i.e. CH, CHBr.CH(OH), CH, Et. (189°). S.G. 1-3. Liquid. From bexyleno oxide and Br (Henry, C. R. 97, 260; Bl. [2] 41, 363).

Di-brome-hoxyl alcohols v. Dibromides of HEXENYL ALCOHOLS

DI BROMO-HEXYL-BENZENE C, II, Br. i.e. Ph.CHEr.CHBr.CH_x.CH_{(CH_x), [80]).} From

hexyl-benzeno (q. v.). Needles or pletes.

BROMO HEXYLENE C. II., Br. (138° 111) at
739 mm. S.G. V. 1:2025. From (β) hexylene
bromide and alcoholic KOH (Caventou, A. 135, 126; Reboul a. Truchot, A. 111, 247; Hecht, B. 11, 1421; A. 172, 70). See also HEXENYL BROMIDE.

Di bromo hoxylono Calladar, S.G. 9 1:598. From hexylene (derived from immuite) and Br (Henry, B. 11, 1054).

Tetra was of hexylene C.H.Br., 1 allylene and Br (Henry, C. E. 87, 171).

Ooto-bromo hexylene C.H.Br., (181°), From sec-hexyl iodide and Br at 130 (Merz a, Weith, B. 11, 2219; Hecht, B. 11, 1420).

Octo bromo hexylene Call Br., From hexane

and Br at 125° (Wahl, B. 10, 402).

BROMO-HIPPURIC ACID C. II. BrNO. i.e. C.H.Br.CO.NH.CH..CO.H. From hipparic acid, alcohol, and Br. Slayder needles. Possibly identical with the following acid.— CaA'₂ (Mayer, Z. 1865, 415).

p-Bromo-hippurio acid
[1:4] C.H.Br.CO.NH.CII. CO.H. Excreted when p-bromo toluene is taken with food. Flat needles (from water). Boiling HClAq forms glycocoll and p-oxy-benzoic acid (Pronsec, H. 5, 63).

BROMO. HYDRATROPIC ACID v. BROMO-

PMENTL-PROPIONIO ACID.

BROMHYDRIN C. GLYCEBIN.

Tri-bromhydrin v. Tri-bromo-profane. p-BROMO-HYDROCARBOSTYRIL C.H.BrNO

i.e. C_eH_z(Br) < NH, CO 4:2 [178°]. Long

V. sol. alcohol, ether, benzene flat needles. and acetic acid. Prepared by reduction of p. bromo-o-nitro-hydro-einnamie acid with and HCl (Gabriel a. Zimmermann, B. 13, 1683).

BROMO-HYDROCINNAMIC ACID v. BROMO-PHENYL-PROPIONIC ACID.

DI-BROMO-HYDRO-CŒRULIGNON v. CORRE-LIGNON.

BROMO HYDROQUINONE C.H.BrO. i.c. C_sH_eBr(OII)_c. [111"]. Formation, -1. Together with di-bromo-hy-

droquinone, by the action of conc. HBrAq on quinone (Wichelliaus, B. 12, 1501).-2. From hydroquinone (1 mol.) and Br (1 mol.) in etherchloroform (Sarmuw, A. 209, 99).

Properties. - Leaflets; may be sublimed. V. e. sol. water, alcohol, and benzene. FogCla forms bromoquinone.

Di-acetyl dericative C.H.Br(OAc) .. [73]. Formed, together with di acetyl di bromohydroquinone by heating quinono with AeBr (Schulz, B. 15, 655). Needles, sol. alcohol and benzene.

Di-bromo-hydroquinone CaH.Br. (OH) .. [1879]. Formation. -1. As above (W.). -- 2. From hydroquinone and Br in HOAc (Benedikt, M. 1, 345).-3. From quinone and Br (Sarauw). Properties. - Long needles (from water).

Converted by Fo2Cld or bromine water into dibromoquinone.

Di-acetyl derivative C. H.Br. (OAe), [161°]. Formed as above (Schulz). Needles, sol. chloroform and ether,

Methyl ether C.H.Br.(OH)(OMe). From methyl-hydrogoinone and Br (B.).

Bi-methyl other Call Br (OMe) ... [142°]. From di-methyl-hydroquinone and Br in HOAc (Habermann, B. 11, 1036). Methyl-ethyl ether C_aH Br₂(OMe)(OEt). [88]. Prepared like tho preceding (Fiala, M. 6, 913).

Di-isobutyl ether C.H.Br.(OC.H.) From C, 11 (OC, IL), and Br (Schubert, M. 3, 684).

Plates (from HOAc).

Di. bromo-hydroquinoneC. H. Br(OH)(OBr)(?). Bromoxy-bromo-phenol. [87]. From quinous (Uniol.) and Br (Uniol.) in chloroform (Sarauw). Golden tables, sl. sol. other and CHCl, decomposed by water into HBr and bromoquinone; changes slowly to C, H, Br, (OH),..

Tri - bromo - hydroquinone CallBra(OH)2. [136°]. Formed together with tetra bromohydroquinoue, by treating di-bromo-quinone with cone. IIBs, or by the action of Br (6 mols.) on hydroquinone (2 mols.) or quinone (3 mols.) (Saranw, A. 209, 116). Silky needles, sol. alcohol and benzene, v. sol. boiling water. Fe,Cl. gives tri-bromo quinone.

Totra - bromo - hydroquinone C_oBr_{*}(OH)_r. [244]. Prepared as above (Sarauw). Prepared also by reducing tetra-bromo-quinone (bromanil) with SO₂ or 11I and 1' (Stenhouse, A. 91, 310; Suppl. 8, 20) or by warming it with cond. IlBrAq. Slender needles (from HOAc); insol. boiling water, v. sol. alcohol. Fe Cl. forms tetra bromo-quinone.

BROMO-HYDROQUINONE-PHTHALKIN v. ! HIDBOQUINONE-PHIHALEIN.

DI-BROMO-HYDROSORBIC ACID v. BROMO-

HEXENOIC ACID.

BROMO-HYDRO-THYMOQUINONE

C1.H1.BrO, i.s. C4H(C1H.)(CH1)Br(OH). [58]. From thymoquinone and HBr (Schniter, B. 20, 1318). Oxidises to bromethymoquinouc [45"]. The di-acetyl derivative is formed by the action of acetyl-bromide on thymoquinone.

Di-acetyl derivative [91°]. Rhombohedral crystals (Schulz, B. 15, 657).

Di-bromo-bydro-thymequinone

C.(C.H.)(CII,)Br (OII)...

Di-acetyl derivative [1220]. Colourless tables (Schulz, B. 15, 658),

BROMO-HYDRO-TOLUQUINON E

C,H2MeBr(OII), [0. 160]. Formed by the action of cold cone. HBr npon toluquinoue. Glistening plates (Schniter, B. 20, 2286).

Tri-breme bydre teluquinene C₂H₃Br₂O₂ i.e. C₂Br₃Me(OH)₂. [202°], tri-bromo toluquinone and SO₂. Need From Needles, sol. water (Canzoneri a. Spica. O. 12, 471).

BROMO - HYPOGÆIC ACID v. Hypogrie

DI - BROMO - ICOSYLENE C. H. Br. From icosinene and Br (Lippmann a, Hawliczek, B. 12, 69).

DI-BROMO-INDIGO CioH.Br.N.O. i.c.

$$C_{\mathbf{4}}H_{\mathbf{4}}Br < \begin{matrix} O \\ C \\ N \end{matrix} > C = C < \begin{matrix} O \\ C \\ N \end{matrix} > C_{\mathbf{0}}H_{\mathbf{4}}Br. \qquad \text{Formed by } \\ H \qquad \qquad H$$

boiling w-di-bromo-m-bromo-o-amido-acetoplienone [5:2:1] C₆H₂Br(NH₂).CO.CHBr, or w-dichloro · m · bromo · o · amido · acetophenone [5:2:1] C.H.Br(NH)CO CHCl or their acetyl derivatives with dilute NaOH and exposure to the air (Baeyer a. Bloem, B. 17, 968). Prepared by heating bromo isatin with PCl, and treating the product with a 10 p.c. solution of III in acetic acid and then with aqueous SO. Some isomeric di-bromo-indipurpurin is formed at the same time (Baeyer, B. 12, 1315). Small black needles, may be sublimed; v. rl. sol. most menstrua. May be reduced to a 'vat' like indigo.

BROMO-INDIRUBIN Cinfl Br N.O. ncedles. Formed by the action of Na CO, on an alcoholic solution containing indoxyl and bronio isatia (Baeyer, B. 14, 1745)

BB-BROMO-IODO-ACRYLIC ACID

C,H,BrO, i.e. CBrI;CH.CO,H. [110], S. 1.7 at 20°. From brome-propiolic seid and 111 (Hill, Am. 3, 175). Scales. - BaA', 3nq. BaA', 16 at 20°. - CaA', 3¹, aq. - AgA'. S. (of

as Brome iede-acrylic acid Clil.CBr.CO.11. [96°]. Formed by the addition of IlBr to B-iodo-propiolic acid (Homolka a. Stolz, B. 18. 2284). Needles, sl. sol. cold water. By heating in alcoholic solution with AgBr it yields αβ-αίbromo-acrylic acid [85] (Stolz, B. 19, 537).

Ba-Bromo-iodo-acrylic acid CliBr:Cl.CO.11. V. sol. water. [71°]. Colourless crystals. Formed by boiling propiolic acid with an ethereal solution of BrI (Stolz, B. 19, 536).

BBa-Bromo-di-iede acrylic acid CBrl:CI.CO₂II. [160°]. S. 2 at 20°. From bromo-propiolie acid and lodine in ethor. Plat being treated with Na.S.O. and distilled with six-sided plates.—Bah', 4aq. S. 16'26 at 20'.——steam (Plimpton, C. J. 41, 805; Sabanejeff, A. bromo-prepiolie acid and lodine in ethor. Flat

CaA'r-KA'.-AgA' (Mabery a. Lloyd, Am. &

αββ-Bromo-di-iodo-acrylic acid

Cl. CBr.CO.H. [182']. Glistening colourless plates. Formed by the action of iodine bromide upon iodo-propiolio acid in ethercal solution (Homolka a. Stolz, B. 18, 2286).

Bia Di-brome-iede-acrylic acid

CBr.:CLCO.H. [110°], S. 3:5 at 20°. From bromb propolic acid and IBr (Mabery a. Lloyd, Am. 4, 91; N. .1m. 4, 17, 94). Monoclinic prisms (from water); ark, ~617;1: 581, — BaA' Baq. S. 16 7 at 29 . CaA' - AgA'.

BaB-Di-brome-iedo-acrylic acid

CIBr.CBr.CO.B. [147]. Long salky needles. Sl. sol. cold water. Formed by the netion of a solution of bromine in chloroform upon iodopropiolic acid IC:C.CO.11 (Homolka a. Stolz, B, 18, 2285).

DI-BROMO-IODO-ALLYL ALCOHOL.

Ethyl other C. II, Br, IO i.e.

CIBr:CBr.CII_OEt. Oil. From iodo-propargyl ethyl ether and Br (Liebermann, A. 135, 286).

o-BROMO-IODO-BENZENE C.11, Br1 [1:2]. (2572). From o-bromo-aniline or o-iodo-aniline by the diazo- reaction (Körner, (l. 4, 339),

m Bromo iedo benzene C.H.Bil [1:3]. (252°). Similarly prepared (K.).

p Bromo iodo benzone C. H. Brl [1:4]. [92°]. (252). Similarly prepared (Griess, J. 1866. 452 ; K.).

Dichloride Call Brl.Cl. (115° 120°, with decomposition). Ppd. as yellow needles when chloring is passed into a solution of brome-iodobenzene in chloroform (C. Willgerodt, J. pr. [2] 33, 158). With alcohol it forms aldehydo and C.H.Brl. It converts metallic and alcoholic iodides into chlorides, ioding being separated.

Tri-bremo-iode-benzene C_e11 Br., [1:3:5:6]. [1047]. Obtained by adding cone, 111 to a solu tion of Call Bos N N NO . Colonrless needles, sel, het algohol (Silberstein, J. pr. (2) 27, 120).

Tri breme iode benzeno C.H. Br. [1:2:4:5]. [165]3.

Dichlorodo Call HaroCle | Formed by dissolving C.H. HBr, in CHCl, and passing in Cl, (Willgerodt, J. pr. [2] 33, 159).

s.BROMO-IODO-ETHANE CallaBri i.e. C11 Br.C11 1, Ethylene bromwdide, [234], (1632) S.G. 22 2 516. From bromo-eth; lene and conc IIIAq at 100 (Reboul, A. 155, 2 " aly 1 hour ethylene are BrI (Marwell Simper for 22, 51). Needles; sl. vol. cold alcohel. Alcohelic KOH gives iodo-ethylene and acetylene (Lagermark, J. R. 5, 334).

u-Brome iode-ethane Cl., CHBrl. idene bromiodide, (142.). S.G. 30 2 452. From bromo-ethylene and cone. InlAq at 4' (R.), or from u-di-iodo-ethane and IBr (Maxwell Simpson, Pr. 27, 421). Alcoholic KOII forms bromoclhylene.

Di-bromo-iodo-ethane C,H, IBr2. (170°-180°). S.G. 20 2:86. From bromo-ethylene and IBr (M. Simpsen, Pr. 22, 51). Ag.O forms C.H.Br.

s - BROMO - IODO - ETHYLENE CallalBr. Acctylene bromiodide. [c. 8']. (150° cor.). S.G. (solid) ? 2.750; ¹⁷⁵ 2.627. Got by passing acctylene into aqueous solution of Brl, the product

With alcoholic NaOH it appears to 216, 266). give off C,HBr.

24-Bromo iodo-ethylene CH,: CBrI. (129°). S.G. 2 2:565. From chloro-bromo-iodo-othane and alcoholic KOH (Henry, C. R. 98, 741). Slowly absorbs oxygen from air.

Di-bromo-iodo-ethylene ClBr:CllBr. [66°]. Small prisms. Formed by the action of an aqueous solution of bromine upon iodo-propiolic acid IC:C.CO.H., CO. being evelved (Homolka a. Stolz, B. 18, 2285).

BROMO-IODO-METHANE CILBrI. Methylene bromiodide. (139). S.G. $\frac{165}{192}$ 2·926. V.D. 9.65. From methylere iodide and Br or 1Br (Henry, C. It. 101, 509).

Di-bromo-iodo-mothano CIIBr. I. Bromiodoform. [6]. From iodoform and Br (Scrublas, A. Ch. [2] 34, 225; 39, 97; Bouchardat, J. Ph. 23, 10).

BROMO-IODO-NAPHTHALENES Cight Brt. The three following are obtained from the corresponding brome-naphthylamines by the diazereaction (McIdola, C. J. 47, 523):

(a)-Brome iodo-naplithalene C, Il Brl [1:4]. Large flat needles; insol. water; sol. alcohol and glacial HOAc, v. c. sol. benzene and other.

(αβ)-Bromo-iodo-naphthaleno C₁₀11 Br1 [1:3]. [68°]. Needles.

(άβ)-Bromo-iodo-naphthalone C₁₀H_aBrI [1:2].
[1]. Thick needles.

BROMO IODO NITRO BENZENE

C, H, Br1(NO) [1:2:5]. [106°]. From C, H, BrI [I:2] and IINO, or from C.H.Br(NIL)(NO.) [1:2:5] (Körner, C. 4, 383). Needles or prisms.

Alcoholic N11, displaces I by N11... Bromo - iodo - nitro - Senzono - C.H.Brl(NO.) [1:4:3].(90]. From C₆H₂Br(NH₂)(NO₂)[1:4:3](K.). ⁴

Bromo - iodo - nitro - benzono CaH.Brl(NO2) [1:3:4]. [84°]. From C.H.Br(NIL)(SO.) [1:3:4] (K.). Alcoholic NII, displaces I by NII,

Bromo - iodo - nitro - benzeno | CallaBrl(NO2) [1:3:6?]. [127]. Formed, together with the following body, by dissolving m-bromo-iodobonzene in funning HNO3 (K.). Alcoholic NII3 displaces Br by NIL

Brome - iodo - nitre - benzeno CallaBrl(NO2)

[1:3:22]. Needles.

BROMO-IODO-NITRO-PHENOL

C₆H₂(OH)(NO₂)BrI[4:2:4:6], [404°], From (4,3.1)brome-attre-phenol, KOH, HIO,, and I (Korner, 5. (-67, -615). Monoclinic tables a:b:c:=-520:11. 7; $\beta=65^{\circ}$ 32' (Groth, 2. Kryst. 1, 437); volatile with steam. – KA'. – NaA'.

Bromo-iodo-nitro-phonol C.H.(O11)(NO.)Brl [1:1:2:6]. From (1,3,6) bromo-nitro-phenol as above (K.). Prisms (from ether). - KA': yellow

BROMO IODO NITRO TOLUENE

C_all_aMcBr1(NO₂) [1:3:4:x], [118°]. Formed by nitrating (1,3,6)-bromo-iodo-toluene. Needles (Wroblewsky, A. 168, 160).

Bromo-iodo nitro toluone Call McBrl(NO.) [1:3:2 or 6:x]. From the corresponding bromoiodo-tolueno (W.).

Dl-bromo-iodo-nitro-toluene

C.HMeBr.1(NO.) [1:3:5:4:2]. [69°]. From C.H.MeBr.I by nitration (Wroblewsky, A. 192, 210). Large needles. Volatile with steam.

Dl.bromo.dl.lodo.nltro.toluene [129°]. From | C.MeBr, 1, (NO2) [1:3:5:4:2:6].

C.HMeBr. I, and fuming HNO. Yields on reduction by Sn and HCl an amido- compound which is converted by further treatment by Sn and HCl to o-toluidine (W.).

 $\Sigma_{i}(f, \mathcal{T}) = \bigcup_{i \in \mathcal{T}} (-1)^{i} = \sum_{i \in \mathcal{T}} (-1)^{i} \mathcal{T}$

BROMO-DI-IODO-PHLOROGLUCIN

C_s(OII)₃BrI₂. From tri-bromo-phloroglucin and aqueous KI. Decomposed by heat (Bonedikt a. Schmidt, M. 4, 605).

BROMO-IODO-PROPANE C.H. BrI i.e. CH, CBrl.CH, (118). S.G. 112.20. Formed by union of III with allylene hydrobromide (Reboul, C. R. 74, 669, 944).

Bromo-iodo-propane CII, CHI.CH, Br or CII, CilBr.CH, i. (160°-168°). From propyleno, water, and IBr (M. Simpson, Pr. 22, 51).

DI-BROMO-IODO-PROPYLENE Calla Br. (?). From iodo-altyleno and Br. Doos not combine with Br (Liebermann, A. 135, 275).

DI-BROMO-IODO-STEARIC ACID

C18 1133 Br. 10 ... From ricinoleic acid C1811 110, vid C₁₅11₂₁1O₂ (Claus, B. 9, 1917). BROMO 10DO-TOLUENE

C.H.MeBrl [1:2:3or5]. (260). S.G. 18 2:139. From C, H, MeBr (NH f (Wroblewsky, A. 168, 164). Bromo-iodo-toluene C_11,McBr1 [1:3:4].

(265°). S.G. " 2.011. From the corresponding brome totaldine (W.).

Di-bromo.iodo-tolueno C.H.MeBr.I [1:3:5:4]. [86°]. (270°). From C.11, McBr(NO.)(NH.) vid C, H, MeBr(NO.)1, and C, H, MeBr(NII,)1 (Wroblewsky, A. 192, 209). Also from di-bromo-ptolnidine, CaH McBr (NIL) by diazo- reaction.

Di-bromo-di-iodo-toluono C. HMeBr. I [1:3:5:4:2], [68°], From C, HMeBr I(NH2) by diazo- reaction (Wroblewsky, 1, 192, 212).

DI-BROMO-ÎODO-TOLUÎDINE

C_11MeBr_1(N11_) [1:3:5:4:2]. [64°]. By reduction of the corresponding uitra- compound (Wroblewsky, A. 192, 210). Converted by sodium amalgam into o-toluidine.

Acetyl derivative CallMeBr I(NHAo) [121"]. Small white needles.

BROMO-ISATIC ACID v. ISATIC ACID.

BROMO-ISATIN v. ISATIN.

BROMO-ISATOIC ACID v. ISATOIC ACID.

BROMO-ISO- v. Buomo-

BROMO-ITACONIC ACID C,II,BrO. [164°]. Formed by the dry distillation of ita-di-bromopyrotartarie acid (Swarts, J. 1873, 584). Its anhydride is formed similarly from ita-di-bromopyrotartario anhydride (Petri, B. 14, 1637). Alkalis form aconic acid; Su reduces it to itaconic acid.

BROMO-LACTIC ACID v. BROMO-OXY-PRO-PIONIC ACID.

DI-BROMO-LAURENE CallaBr. (?) [210° From laureno and Br (Montgolfier, A. Ch. [5] 14, 93).

Tri-brono-laurene C₁₁H₁₂Br₃? [125°]. From lanrene and Br in the cold (Fittig, Köbrich a. Jilke, A. 145, 149). Cf. Laurene.
**CROMO-LEVULIC ACID v. BROMO-ACETYL-

PROPIONIO ACID. BROMO-LUTIDINE v. BROMO-DI-METHYL-

BROMO-MALEIC ACID C.HBr(CO.H), i.e. CO,H.CH.CBr.CO,H, or CO,il.C.CHBr.CO,H,

CBr.C(OH) CH.C(OH) 0. or ij. ĊН ČBr .

Formed by boiling di-bromo-succinic acid or its Ba salt with water (Kekulé, A. Suppl. 1, 367; Petri, A. 195, 62). Formed also, together with bromo-fumuric acid (q. v.) by the action of Br and water on succinic acid at 180° (Kirkulé, A. 130, 1), or fumarie acid at 100° (Carins, 4, 149, 264). Deliquescent prisms or needles; v. e. sal. water, alcohol, and ether, splits up into water and its anhydride on distillation.

For discussion of formula see Man ic acm.

Reactions, -1. Sodium-amalgam gives succinic acid. 2. Furning HBr unites in the cold, forming di bromo succinic acid. 3. Electrolysis of its Na salt gives CO and HBr. A. Boiling cone, birgha-water forms oxalle and areticacids. -- 5. When it is discolved in water and an equivalent of aniline is added there reparates a crystalline pp. C.HBr.3CO/Hi4CO/H.NH/C.H.). [128]. This acid aniline saft dissolved in water and allowed to stand deposits the acidanilide CO H.C HBr.CO, NHC, H. which crystallises in prisms; in sol. in didute BCl. If instead of allowing the sub-tance to react in the cold the solution is heated, the compounds $\mathbf{C}_{l_0}\Pi_{12}\mathbf{N}_1\mathbf{O}_2$ and $\mathbf{C}_{l_0}\Pi_{l_1}\mathbf{N}_1\mathbf{O}_3$ are obtained. $\mathbf{C}_1\mathbf{N}\Pi_1\mathbf{P}_1\mathbf{h}_2\cdot\mathbf{C}\mathbf{O}_2$

$$C_{16}H_{12}N_2O_{27}$$
 probably β

$$CH = -CO$$
NPh,

[230°], forms ochre - coloured microscopie needles. Insel, het, A. rol, cold aq; rol, hot, alcohol, less in cold; Collas O, pro-C(NHPh) CO_H

, (176), fram syellowich bably CILCO.NIII'h

indistinct cry tabs. Sol, bot, m. sol, cold ag. Sol, nikalis : acide pp. the end thance unchanged (Michael, Am. 9, 1891; B. 19, 1873).

Salts. As A" it ery taking pp. CaA" tag.

-CaNa A" 4 ap. PbA" ap.

2008 i.V.). Con-Dimethal other A'Me. verted by iodine rate dimethyl brong from rate. Distingly other A'Vit. (256 i.V.). (140° 150°) at 25 mm. (Anachitz, B. 12, 2254)

Schacherl, A. 229, 91). Anhas 'r de C 115rO., (215 'i.V.), Tormed ! as above, and also by heatner distremos-accinic

geid with Ac O at 139 (An shotz, B. 10, 1884). Water form cloomo-maker acid.

Amide C.H.ErN.O. (1968) 175 []. From the ! imide an LML,

Imide (CAB)(O)NH. [151]; Formed, together with the graide of distronounching acid, by locating saccinimide with Br at 160 [Ciamici in a. Silber, B. 17, 557; Ki ichm ki, Saz, D. 74, 560).

Iso brome - maleic acid is Bhotheri mane ACID (q. r.).

Bromo mulcic acid (2) C.H L.O. From mucobromic acid and buyla (Hal, E.

239a - K A" aq. - GaA" 2.aq. - V₂ A".

Di bromo-maleic acid Cilir II O. Formed, together with bromo-scale acid, by brominating succinic as I (Keladé, A. 130, 2). From (3)-di-brome-pyromacic acid and from tri-bromo-pyremacie acid by Let dilute $11 \times O_{\chi}^{-1}$ (Hill a. Samper, A. 232, 89). Formed also by heating muco comic and with Br at 140' (Hill, Am. 3, 48; B. 13, 731). Sander felt dure dies, v. sol. water, alcohol, and other, v. sl.; of, benzene and ligroin. An equivalent quantity of aniling added to a solution of the sold in water disciders " Vol. I.

and then deposits the acid aniline salt. When this is allowed to stand under water it gradually forms an anilide. The neutral salt in the same way gives the di-anilido C. Br (CO.NHPh), (110 % By heating the acid (3 pts.), with water (60 pts.) and aniline (22 pts.) for 30 minutes C(XIIFh) = CO,

NPh [183] is obtained in the ·co/ Chr. ... torn; of that, yellowish red pri men book mp.; m. sal. hot alcohol (Michael, Ang. 9-1-9).

Salts. BaA' 2aq. 8.6% of 20 . PhA' aq

—Ag A": explodes when hear h. Annyarian CBr G. attack Prepared by

heating the acad. Needles (by sublimation) yelsol, cold water,

heating succinimide with broming. ervstals, athe = (13)2.1; (9649); \$\beta\$ (14) (50). By boiling with aqueous KOH it is converted into di bromo mateie, acid (Ciamic), te n. Softer, B. 17,556; 6,14,35; cf. Kalicharki, Sit. R 74,561).

Brome na cie-acid-di-bromide c. The promo-SUCCENIC ACTO.

DI BROMO MALEIC ALDEHY DE C.H.Br.O. 190 : Got in small quantity from (6) - delicano Lycomucie acid and aquaous, biomire (Tonnics, 7), 12, 1203 ; Hill a. Sanger, A. 23 5, 87). Long thin prisms (from water). V. sed. alcohol, ether, Chleroform, and benzene, m. rol. light petrolearn. In a carrent of CO, it may be subfurned. On oxidation it gives mucchronne acul-

BROMO-MALEYL BROMIDEC THE O . 15621 Permed by treating contain brome pyromanic acid with cold bromine (Hill a. Sarger, A. 232, 80). Long prietos (from light petroleam).

BROMO MALIC ACID Call late. The Bodinm salt NallA" is formed by the action of NaOF t sodium di bromo ruccinate. Reilung limeswater convert at into calcium racena te. Sodiumamal/mo fori action suci sta Tb(OAc), $\mathbf{p}_{\mathbf{l}}(\mathbf{s},\mathbf{P})\mathbf{A}$.

Ethstderivative. Sodium salt CO Na.CH, CHIOEi).CO Na. Hygro oper mass; Famed by adding alcoholic buttlet to codimic disbroma succinate (Malder & Handsteger,

R. T. C. 1, 151).

BROMD-MALONIC ACIP 6 I have i.e. CHBr(CO II). Oldained by ritter or the dibrominated and with sudiam and the related of B. 40, Co. B. 41, Co. Delay brown tyrinist Ar. O form battonic and the ArthA". Ar Ar.

Di bisman (anionie nerd Clia (1) H) ste. 126 J. From malonic head CHCl, and Ri (Febrieff, \vec{B} 7, 400; J. R. 10, 65; Vat 't Holl, B. 8, 3.5). Sociles, v. e. sol. water. Bolling baryta water form one, oxahe held.

Acride CBr (CONIL). (206 % Located by adding Br to make made in aquest. soluwoon (Ureund, B. 17, . 2). Former at a by the action of alcoholic NH, on the amide of pentabeome-acete acid CBr, CC, Cla CO, MI, (Stokes a. Pechasam, Am. 8, 380). Nordles, prions, or large netaledra. Sl. sol. hot water, alcohol, and acetic acid. - CBr2 CO.NH Hg: white amorphous powder, insoluble in water and

alcohol. Methylamide CBr. (CO.NHMe) : [1620];

From tetra-hydro-phthalic acid and bromine-water (Bueyer, A. 166, 353). Prisms or tables (containing laq). Baryta-water converts it into C,II,,O,

BROMO MELILOTIC ACID v. BROMO-OXY-PHISAL-PROPIONIC ACID.

BROMO MESITENE LACTONE

 $C_aH_aBr<\frac{O}{CO}>$, [1052]. From mosite ne lactone, CS, and Br (v. Aceto-acetic ether). Sl. sol. cold alcohol and water (Hantzsch, A. 222, 18).

BROMO . MESITOL Call Bro t.r. C,HMe,Br(OII), [802]. Needles (from alcohol) (Biedermann a. Ledonx, B. 8, 59).

Di bromo mesitol C. Me Br. (OII). [150]. From mesitol, Br. 110Ac, and I (Jacobsen, A. 195, 265).

BROMO MISITYL ALCOHOL v. BROMO- w-OXY-MESITYLENE p-BROMO - MESITYL BROMIDE v. p-w-1)1.

RROMO-MEST LYERNE

BROMO-MESITYLENE Call, Mea Br [1:3:5:1]. Bromo-s-tri-methyl-ben enc. [-12]. (2272 i. V.). S.G. 49 1:32. Formed by the action of I mol. Br. upon cold mesitylene in the dark (Fittig a. Storer, A. 117, 6; Schramm, B. 19, 212). (5:3:1) . Bromo-mesitylene

JH,(CH) (CH Br) [1:3:5]. Mesital bromide. [38]] (231). Prisms. Formed by bromination of meatylene at 130° (Wispek, B. 16, 1577; Colson, A. Ch. [6] 6, 89; C. R. 96, 713).

es Di-bromo mesitylono CallMe Br... (278'). Long needles. Formed by the action of 2 mole, of bromine upon mesitylene in the dark (F. a. S.; Süssenguth, A. 215, 248; Sehramm, B. 19, 212). Funning HNO₄ gives bromo-di-nitro mesityleno [194°].

n-w.Di-bromo mesitylene

C.H. (CH.) Br(CH.Br) [5:3:4:1], p - Bromo - mesityl browide. Oil, fluid at 2 19". Decomposes on distillation. Formed by the action of bromine (1 mol.) in sunskine upon esosbremomesitylene (Schramm, B. 19, 213).

ww.Di-bromo mesitylene Call Mc(CH Br). [66"]. Formed by passing CO, charged with bromine vapour into boiling mesitylene (Colson, 1. Ch. [6] 6, 92; C. R. 96, 713; Robinet, C. R. 96, 500). Formed also by treature ωω-di-oxymeshly been with HHr (Rebinet a. Colson, Bl. [2] 40, 111). Then prisms; decomposed by alcohol.

Tri-cso-bromo-mesitylene C.Mc.Br., (221). Formed by the action of 3 mole, of bromine ayon mesitylene in the dark (Schraum, B. 19, 213). Triclinic crystals, v. sl. sol, alcohol.

: ω.ω..Tri.bromo mesity lene C.H.(CH_)Br(CH_Br), [5:153:1], [122]. Obtained by the action of bromine (1 mol.) in sunshine upon hot p-e-di-bromo mesitylene (p-bromomesityl-bromide) GML(CH,),Br(CH,Br). Very slender needles (from alcohol) Schramm, B. 19, 215).

c.ω.ω..Tri-bromo mesityleno C,11,MeBr(CH,Br), [5:2:1:3]. 1817. bromo was-di-oxy mesitylene and cone. HBr. Can be formed by brominating mesitylene (Colson, A. Ch. [6] 6, 101; Id. [2] 41, 362).

large white needles or trimotric crystals (Freund, B. 17, 782).

BROMO MALOPHTHALIC ACID C. II., BrO. BROMO MASITYLENE, GLYCOL v. BROMO BROMO MESITYLENE, GLYCOL v. BROMO DI-OXY-MESITYLENE.

BROMO-MESITYLENE SULPHONIC ACID C.II, BrSO, i.e. C. 11Me, Br.SO.H. Formed by the action of bromine-water on a very dilute solution of mesitylene sulphonic acid or its Ba salt; formed also from bromo-mesitylene and furning H.SO, (Rose, A. 164, 56). Deliquescent trimetrie needles (from ether), - BaA', aq. -PbA', I laq. KA'aq. CuA', 1aq. NaA'. (α)-BROMO-MESITYLENIC ACID C., H₀BrO.

i.e. C. H. Mc Br(CO.H) [1:3:1:5; '117']. Formed, together with some of its isomerides, by the slow action of Br on mesitylenic acid in the cold Formed also from the corresponding amideme itylenie acid (Schmitz, A. 193, 172). Trimetric prisms (from alcohol), a;b;c = 927:1: 470 - BaA' laq: monoclinie, a:b:c = 3:068:1: 801; β=63' 21'. CaA'. 2aq.
(β) Bromo mositylenic acid

G.H. Me.Br(CO/H) [1:3:2:5]. [215°] (Sch.); [212] (8.). Formed by oxidising bromo-mesitytone (Fittig a. Storer, A. 147, I), or from the corresponding amido-acid (Sch.). Monoclinic $\begin{array}{lll} & \text{crystal} s_i \, a(b) c & \text{PB3}(1) 760 ; \; \theta = 70 \, ^{\circ} \, 35', + \, \text{BaA'} \; , \\ & - \, \text{CaA'}_{,*}, & \text{CaA'}_{,*}, \; \text{Suq.} & \text{KA'}, \end{array}$

Di-bromo-mesitylenic acid CallBr.Me. (CO.H). [195]. Formed by oxidising di-bromo-mesitylene (Süssenguth, A. 215, 250). Needles (by sublimation). - CaA', 7aq. - BaA', 34aq.

BROMO-METHACRYLIC ACID C. II Bro., i.e. CHBr:CMe.CO.II. Bromo-crotonic acid. [63°]. (229). From citra- or mesa- di-bromo-pyrotartaric acid by frontment with water, Na,CO,Aq, or KOHAq (Kekulé, A. Suppl. 2, 97; Cahours, A. Suppl. 2, 317; Fittig a. Krusemark, A. 206, 7; Friedrich, A. 203, 354). Also from aß-di-bromoisobutyric acid and NaOHAq (C. Kolbe, J. 1r. (21.25, 382). Flat needles, sl. ad. cold water. Reduced by sodium amalgam to isobutyric acid. Decomposed by heating with alkalis into methane, allylone, and sectic acid (F.) .--CaA', Saq. S. (of CaA',) 5 75 at 11". - AgA', -HO,CaA', NH, HA', (Morawski, Sitz, B, 74, 39).

Ethyl ether EtA'. (193) (C.).

Bromo-methaerylic acid CH :C(CH.Dr).CO H. [66]. Formed, together with the preceding, by boiling most-di-bromo-pyrolartaric acid with water or Na.CO₂Aq (Krasemark, A. 206, 12). Lamino (from water), v. sol. water; volatile with steam. Reduced with difficulty by sodium amalgam to isobutyric acid. - CaA', 2aq. S. (of CaA',) 80 at 5

Di-bromo-methacrylic acid C,H,Br,O, Needles. From tri-bromo-butyric acid (dibromido of bromo-methacrylic acid). At 120 'it takes up Br forming tetra-bromo-butyric acid, whence boiling alkalis form

Tri-brome methacrylic acid C.H.Br.O. Needles (C.)

BROMO-METHANE v. Magnyt promide. Di-brome methane p. METHYLINE BROMDE. Tri-bromo methane v. BROMOFORM.

Tetra-bromo-methane CBr., Curbon tetrabromile. [92°]. (189°). Occurs in commercial bromine (Hamilton, C. J. 39, 18).

Formation .- 1. By heating Br with CS, in ω.ω.σ. Til-brome-mesitylene (C.H. (CH Br), presence of I or SbBr, (Bolas a. Groves, C. J. [94°], (21°) at 10 mm. From boiling mesityle 23, 161; 24, 773; A. 156, 60; 160, 160}.—2. From CH2Cl, and IBr, (Höland, A. 240, 236). -8. From alcohol and Bc (Schäffer, B. 4, 356) .-4. By exposing a mixture of dilute KOH, bromoform, and Br to sunlight (Habermann, A. 167, 174).-5. By heating bromoform or bromopicrin with SbBr, or BrI at 1502, 6, From CI, and Br (Gustayson, A. 172, 176), -7, From CCl, and Al Brout 100 (Gustavson, J. R. 13, 286). 8. From CH,Br and Br in presence of animal charcoal (Damoiscan, C. R. 92, 42).

Preparation, CS, (2 pts.) is heated with iodine (3 pts.) and Er (11 pts.) for 96 hours at

1502 (Höland, A. 240, 238).

Properties. Tables with frint complor like smell; extremely prese to sublimation. At 220° it splits up into C Br, and bremine. Boiling alcohol gives bround, HBr, and ablehyde. Alcoholic KOH gives K CO, and KIr. Sedium analgam forms CHEr, and CHEEr

BROMO METHANE DI SULPHONIC ACID CHBr(SO(H)). Patassium salt KA". From bromo di sulpho abdeley le CBr(SO, He, CHO, ley boiling with aquacous K CO, (Rathke, A. 161, 161).

Di bromo methane sulphonic acid CHBr. SO, H. Barium sait BaA'. Thin uncloses plates, formed by the action of Brancon terriori sulpho actate at 130 (Ambiensch, 56, 7, 157).

BROMO PENTA METHYL TRI AMIDO TRI-PHENYL CARBINOL Contraction Co. Hydrobromide C, II, N, HBr. Formed by heating di methylamiline with Br at 120 (Branch abar, Brunner, H. 10, 1845, 11, 697a

DI-BROME-METHYLAMINE McNBr. r. METHYLAMINE.

p BROMO METHYL ANILINE

Calla Br; NHMe. (11 % (260)). Prepare I from the nitro anime.

Acetyt desirative '99.

Nitrosamine C.H.(Br)NMc(NO), 471%. Long needles. Formed by the action of BNO, on polaromo-di-methyl-aniline (Wurster n. Selo de, B. 12, 1818).

m-Brome di methyl-aniline CJI (Br) NMe₂ 11(3), [112], (264) corr.). Prepared by the methylation of actor manifum. By the action of HNO, it gives a nitroso-compound which forms baht green needles and melts at labout 11s.

Methylociode is Cat Br. NMc L. Leaflets (Wurster b. Scheibe, B. 12, 1818).

p-Bromo di-metavl aniline C.H. dligNMe.

[14], [55] l. (261 covr.).

Preparation, 1, by boundation of dimethylanilimedieserved in acetic acid (Weber, 8, 711; 10, 763).
 19 methylation of ρ. brota miline. By the action of HNO, it gives a mixture of printro-demethyl anifine and pe bromo-plicayl-methyl-mitrosamene

Methylo-iodide C.H.Bo.: Me.L. [185].

(Wurster a. Schelbe, B. 12, 1816).

Ferrocyanide B H. he(CN) (#6): barter, Ferricannide Ballication, bug: very soluble yellow crystala (Wurzter a. Beser, B. 12) (DI-BROMO-MET. Fileme-DI-PHENYLENE

BROMO . DI . METHYL. ANILINE - PIITHA . LEIN C., H., Br., N.O., i.e.

 $C_{i}H < \frac{C(C_{i}H_{i}BrNMc_{i})_{2}}{COO}$. The hydrockleride, CO.0 formed by heating p-bromo-di-me by amiline with phthalyl chloride, crystallises in 1001 blue (1130), by adding lar to o-methylene needles. Conc. HClAq pps. dings given li 2HCl oxide supended in water. Black os at c. 225° (O. Fischer, B. 10, 1623) B 'H P(C).

DI-BROMO-METHYL-ANTHRACENE

C₁₃H₁₀Br₂. [138°-140°]. From methyl-anthracene by Br in CS₂ (Liebermann, A. 212, 85). Yellow needles (from glacial HOAo).

Tetra bromo methyl anthracene C. H.Br. Needles (from toluene). Oxidises to dishromo-

methyl-authraquinoue (b.).

DI BROMO - DI - METHYL - ANTHRACENE. DIHYDRIDEC, Haller. From dimethyl anthrarens dihydrale and Br in HOAc (An.eliatz, A. 235, 309). Oxidises to author home.

DI-BROMO METHYL-ATROLACTIC ACID v. Di Blomoloxy, folky i planetonie se in

TETRA BROMO METHYL AUR'NE

C. II. Br.O. Formed by loominating methylnature B'HBr 2aq (Zulkowsky, M. 3, 471).

BROMO . METHYL - BENZENE v. BROMO-TOUTTSIL.

Bromo-di-methyl-benzene v. Bureto xylena. Tri-bromo tri methyl benzene Ca(CH,)Br. [1:2:3:1:5:6). Tri bromo hemimeliithene. [245"]. Needles. St. sol. alcohol. Formed by bromination of (1.2;3)-tri-methyl-benzene (Jacobsen, B. 15, 1858). Other bromo-tri-methyl-benzenos are described as Bromo-y-cumenes and Bromo MUSITYLENES.

Bromo - tetra - methyl - benzene r. Browe

Bromo penta methyl-benzene C. BrMe #161 . (289). From C.Me.H. Br, and I (Friedd a. Craft , A. Ch. [6] 1, 473).

Hexa & brome hexa methyl benzene

C (Cit Er), (255°) (F. n. C.); (227°) (H.). From hexa methyl-benzene, water, and Br at 10) (Hofmann, B. 13, 1732; Friedel a. Crafts, A. Ch. [6] 1, 468).

BROMO METHYL BENZOIC ACID e. Bromo TODAYAC ACID

Bromo-di-methyl benzoic acid

C.H.BrMe CO.H. [1734]. Bromo-pseudo-cumeric a id. Bromosylylic acid. From C. H. BrMc. (4:2:1:5) by C O, in HOAr (Sue enouth, A. 215, 241). And from C.H.Me. (CO.H) [188,4] and Br (Ganter, B. 57, 1608). Needles (from y V. e. sol. ale Nol. - CaA', 2nq. - Ba V 6nq. Needles (from water).

Bromo-di-methyl-benzoic acid C.H.BrMe,CO.H. [189]. Bromo-p vydytic acid. I ion G H, Me (CO H) [1:2:4] and Re (Gorder, B.

17, 1609). Needles (from dilute alconed). Other isomerides are described at Bromo-

ero-BROMO-DI METHYL-COUMARIN

destruitment series (g. r.).

,C(CH_a;.CBr $e'n'(cn') < 0 \longrightarrow c0$ Formed by bromina-

tion of diamethylocommunicalis olyed in CS. Crystalline rolid. Sl. rol. redad. Converted by het alcoholee KOH into di noshy) commarilie ner l eli-methy i-comourence carboxylic acid) (Hantzsch a. Lang, B + c9, 1209).

C.[11,Br.; 1962]. From methylene-di-pheny'ene (q. c.). Nocides or actahedra (from ether) (Car-nebey, C. J. 37, 719).

HEXA-BROMO-METHYLENE-DI-PHENYL-ENE OXIDE C., H.Rr.O. Formed, together with the heats branisated compound Cislishr,O (Salzmann a. Wichelbaus, B. 19, 1401).

BROMO-METHYLENE-PHTHALIDE

C=CHBr >0 [133°]. Long C,H,BrO, i.e. C,H, c=0

colourless needles. Formed by heating phthalylbromo-acetic acid in vacuo; or by bromination of acetophenone-carboxylic acid. It combines with Br., forming C_bH₁ < CBr(CBr.H) O. [118°]

(Gabriel, B. 17, 2525).

p-BROMO-METHYL-ETHYL-ANILINE

C,H,BrN i.e. C,H,Br.NMeEt. (265°). From methyl-cthyl-aniline and Br. Solidifies below 0° (Claus a. Howitz, B. 17, 1327).

TRI-BROMO-DI-METHYL-ETHYL-BENZ-ENE C₁₀H₁₁Br₃ i.e. C₆Br₃Me₂Et. [218°]. (Jacobsen, B. 7, 1434).

DI-BROMO - METHYL - ETHYL - GLYOXA -LINE C₃Br₂(CH₃)(C₂H₃)N₂. Di-bromo-oxalethyline. [38]. Colourless crystals. Sol. acids. Formed by bromination of methyl-cthyl-glyoxaline (oxal-ethyline) (Wallach, B. 16, 537).

HEXA-BROMO-METHYL-ETHYL-KETONE C₄H₂Br₆O i.e. CBr₃.CO.CH₂.CBr₃. [90°]. From u-di-bromo-ethylene and HBrO (Demole, B. 11, Reduced by sodium-amalgam to methyl ethyl ketone. Fuming IINO, gives malonic acid.

DI-BROMO-(B, 2-Py, 2)-DI-METHYL-(Py, 3)-ETHYL-QUINOLINE $C_{13}H_{13}Br_2N$. [144°]. White needles (Harz, B. 18, 3389)

TRI-BROMO-METHYL-GLYOXALINE

C₃Br₃(CH₃)N₂. Tri-bromo-oxalmethylin. [89°]. White crystals. Insoluble in cold water.

Formation .- 1. By the action of McI on tribromo-glyoxaline-silver .-- 2. By bromination of methyl-glyoxaline (oxal-methyline) dissolved in dilute H.SO, (Wallach, B. 16, 537).

BROMO-METHYL-INDONAPHTHENE-CAR-BOXYLIC ACID $C_6H_4 < \stackrel{\mathrm{CBrMe}}{\underset{\mathrm{C11}}{\mathrm{C1}}} < C.CO_2H_1$ [215°].

Formed by bromination of methyl-indonaphthene in chloroform. Needles. Sl. sol. alcohol (Roser, B. 20, 1575).

HEXA - BROMO-DI-METHYLENE DIKETONE C₃H₂Br₆O₂i.e. CBr₃.CO.CH₂.CO.CBr₃. Hexa-bromo-acelyl-acetone. [108°]. From the diketono and Br (Combes, A. Ch. [6] 12, 210). Needles; decomposed by alkalis into tri-bromomesty. and tri-promo-acetic acid.

eso BRoz (298° corr.). Colourless fluid. C₁₀II₀Br(CH₃). action of bromine on a cold

Formed by the "I-naphthalene in CS... solution of (a)-methy round

Picric acid compa5°]. Yellow needles C₁₁H₉Br,C₆H₂(NO₂)₃OH. [1. (Schulze, B. 17, 1528).

eso-Bromo-B-methyl-naphthale.;d. Formed C₁₀H_aBr(CH₂). (296°). Colourless fluctuon of by the action of became on a cold so (β).methyl-naphthalenc in CS

Pieric acid compound C₁₁H₂Br,C₂H₂(NO₂)₃OH [113°], yellow need.] (Schulze, B. 17, 1528).

ω-Bromo-(β)-methyl-naphthalene C₁₀H, CH₂Br. [56°]. (213° at 100 mm.). White glistoning plates. Formed by passing gaseons bromino into (B)-mothyl-naplithalene heated to 240° (Schulze, B. 17, 1529).

Tri-bromo-di-methyl-naphthalene C, H, Br. [228°] (Cannizzaro a. Carnelutti, G. 12, 410; cf. Giovanozzi, G. 12, 147).

TRI-BROMO-METHYL-DI-PHENYL-AMINE $C_{13}H_{10}Br_3N$ i.e. $NMe(C_8H_3Br_2)(C_6H_4Br)$. [98°]. From methyl-di-phenyl-amine and Br (Gnohm, B. 8, 926). HNO₃ forms {C₃H₂Br(NO₂)₂}₂NH.

Totra-bromo-methyl-di-phenyl-amine (C, H, Br.) NMe. [129°]. Formed at the same time as the preceding (G.).

DI-BROMO-DI-MÉTHÝL-QUINOL v. Dimethyl- Bromo-Rydroquinone.

BROMO METHYL PIPERIDINE

CH₂ CH₂ CH₂ NMe. The methylo-bromids (B'MeBr) is formed very readily by isomeric change of di-methyl-εδ-di-bromo-n-amyl-amino (so-called 'di-methyl-piperidiuc-di-bromine'), CH Br.CHBr.CH .. CH .. CH .. NMe., by warming its alcoholic solution for a short time (Merling. B. 19, 2630).

DI-BROMO-METHYL-PYRIDINE C.H.Br.N i.c. C, NH, MeBr. [109°]. Formed, together with ethylene bromide, from tropidine (q. v.) hydrobromide and bromine at 165° (Ladenburg, A.

217, 145).

s-Di-bromo-di-mothyl-pyridine C,NHMc,Br2 [1:5:2:4]. Di-bromo-lutidine. [65]. Formed by the action of bromine upon an aqueous solution of the potassium salt of s-di-methyl-pyridine-di-carboxylie acid. - B'2H2Cl2PtCl42aq: needles (Pfeiffer, B. 20, 1350).

Di-bromo s-tri-methyl-pyridine C,NMc,Br, [1:3:5:2:4]. Di-bromo-collidine. [81°]. (262°) at 726 mm. Obtained by the action of bromino upon an aqueous solution of the potassium salt of s-tri-methyl-pyridine-di-carboxylic acid; the yield is 50 p.c. of the theoretical. White pearly plates. Very volatile with steam. Weak base.

Salts .- B'HCl; easily soluble small glistening crystals. B'_ll_Cl_PtCl, 2aq: orangeyellow needles. B'.H.Cr.O.: [146°]; needles. — ^B'C,H.(NO.),OH: [160°] dark-yellow tlat prisms, v. sol. hot alcohol, insol. water (Pfeiffer, B. 20, 1345).

DI BROMO TRI METHYL PYRIDINE DI. CARBOXYLIC ETHER. Dibromide

NC_sH₁Br₂(CO₂Et)₂Br₂. [102°]. From the following body and funning HNO, (Hantzsch, A.215, 17). Di-bromo-tri-mothyl-pyridine di-carboxylic

ether. Di-bromhydride NC₈H₂Br₂(CO₂Et)₂H₂Br₂. [SS°]. From the di-hydride of (1,3,5,2,4) - tri - methyl - pyridino - di earboxylic ether by Br in CS. (Hantzsch, A. 215, 14). Yellow twin crystals.

DI-BROMO - DI - METHYL - PYROCATECHIN v. Di-methyl ether of Di-Beomo-Pyrocatechin.

HEXA-BROMO-D1-METHYL TRISULPHIDE C₂Br₄S₃ i.e. (CBr₃)₂S₃. Carbotrilhiohexabromide. [125°]. S. (alcohol) 5·5 at 78° ; S. (ether) 2·35 at 0° . From CS₂ and Br (Hell a. Urech, B. 15, 275, 987; 16, 1147). Prisms or tables, insol. water. Hot cone. NaOH gives NaBr, Na CO. and Na2S3. Decomposed by heat into CBr4, CS2Br, SBr2, and a blue substance CaBr4S4 2aq.

DI-BROMO-METHYL-THIOPHENE HBr. (CHa)S. (228°). Oil (Moyer a. Kreis, B.

787). 11 i-bromo-(a) methyl-thiophene C₁Br₂(CH₃)S. 186°l. Formed by bro-Tri-b. (β)-thiotolene. [86°]. Formed by brominatio. (β)-methyl-thiophene (Egli, B. 18,

545). Long colourless silky needles. V. sol. ether and hot alcohol.

Tri-bromo-methyl-thiophene C,Br,(CH,)S. [39°]. Formed by bromination of the methylthiopheno from pyrotartaric acid. Large colourlese needlos (Volhard a. Erdmann, B. 18, 455). Forms a molecular compound [74°] with the preceding (Gattermann, \vec{B} , 18, 3005)

Bromo-di-methyl-thiophene C,H(CH3),BrS. Bromo-thioxenc. (194° uncor.). Formed by bromination of thioxene dissolved in CS. Volatile with eteam. Colourless fluid. Heavier than water (Messinger, B. 18, 1637).

Di-bromo-di-methyl-thiophene C4(CH3), Br.S. Di-bromo-thioxene. [46°]. (217° nncor.). Long colourless needles. Formed by adding 2 mols. of bromine to cooled thioxene (from coal-tar)

(Messinger, B. 18, 563).

Di-bromo-di-methyl-thiophene C,(CII3), Br.S. Di-bromo-thioxenc. [47° 50]. Needles. Volatile with steam. Formed by bromination (with 2Br.) of thioxeno (from acetonyl-acetone) (l'aal, B. 18, 2253).

Tri-bromo-di-methyl-thiopheno CaH BraS i.c. C.Br.,(CH.,)(CH.,Br)S. Tri-oromo-thioxene. [141°]. Crystallises and sublimes in needles. Formed by the action of an execss of bromine upon the di-bromo- derivative of the thioxene obtained from acetonyl-acetone (Paal, B. 18, 2253).

Ooto-bromo-di-mcthyl-thiopheno C₄Br₂(CBr₃)₂S. Octo-bromo-thiovene. [114°]. Small needles. Formed by the action of an excess of bromine upon thioxene (from coal-tar) (Meseinger, B. 18, 565).

BROMO-DI-METHYL-O-TOLUIDINE

C₉H₁₂BrN i.e. C₈H₃(CH₃)(Br)NMe₂. (245°). Prepared by bromination of di-methyl-o-toluidme, or by methylation of bromo-o-tolnidine (Michler a. Sampaio, B. 11, 2172). Liquid; sol. alcohol and ether, volatile with steam.

Bromo-di-mothyl-m-toluidine

 $C_1H_2(Br)(CH_2)NMe_2$ [1:2:4], [98°], (276°), Prepared by bromination of di-niethyl-m-toluidine. White leaflets. Insol. water, sol. alcohol, ligroin and C.H. (Wurster a. Riedel, B. 12, 1800).

Ferrocyanide B'.H. Fe(CN), 4aq: crystals. Ferricyanide B'H.Fe. (CN)12 9aq. Very soluble yellow crystals (Wurster a. Roser, B. 12, 1826).

TETRA BROMO-MYRISTIC ACID

C₁₄H₂₁Br₄O₂. From myristolio acid and Br (Masino, A. 202, 176). DI. BROMO-MYRISTOLIC ACID CHILL Br.O.

Obtained by gently warming the preceding (M.). α.BROMO-NAPHTHALENE C_{10} II Br [1], [5°], (277°); (280° cor.) (Ramsay a. Young, G. J. 47, 650). S.G. $\frac{24}{3}$ 1·1750; $\frac{12}{3}$ 1·503. R_{22} 81·9 (Nasini, G. 15, 93).

Formation.-1. From naphthalene in CS2 and Br (Laurent, A. Ch. [2] 59, 196; Glaser, A. 135, 40; Wahlforss, Z. 1865, 3; Guehm, B. maphthalene sulph mic acid or nitro (a) bromo-15, 2721).—2. From diazo-bromo-naphthalene amphthalene [85] with PBr (John, Bl. [2] 28, salts by boiling with alcohol (Rother, B. 4, 851; Stallard, C. J. 49, 188).-3. From Hg(C1011,)2 and Br (Otto, A. 117, 175).

Properties.—Liquid, insol. water, miscible with alcohol, ether, and benzene.

Reactions.—1. CrO, gives phthalic acid (Beiletein a. Kurbatow, C. C. 1881, 359).—2. A colution in CS, gently heated with Al,Cl, gives (A) heams noulithalene as the chief product, to-

gether with di-bromo-naphthalenes, and naphthalene (Roux, Bl. [2] 45, 510) .- 3. Toluene in presence of Al₂Cl₄ forme bromo-toluene and naphthalene (Roux).—4. Reduced by sodiumamalgam to naphthalene.-5. Cl.CO.Et and Na give naphthoic acid.

Picric acid compound C₁₀H₂BrC₆H₂(NO₂)₃OH, [135]. Yellow needles (Wichelhaus, B. 2, 305; R.).

Dichtoride $C_{10}H$ BrCl₂: [165°]; tables. (β)-Bromo-naphthalono $C_{10}H$ Br [2]. [59°]. (282 cor.). S. (92 p. c. alcohol) 6 at 20 .

Formation. -1. By heating (β) -diazo-naphthalene with a large excess of HBr (Gasiorowski Wayss, B. 18, 1911; J. Liebermann, A. 183, 268). 2. By running a volution of (β)-diazonaphthalene bromide into a hot solution of cuprous bromide; the yield is 30 p.c. of theoretical (Lellmann a. Remy, B. 19, 811). - 3. From (\hat{B}) -naphthol and PBr₅ (Brunel, \hat{B} , 17, 1179).—I. From (α)-brome-naphthalenc and Al₂Cl₆ (Roux, Bl. [2] 45, 513).

Properties. - Trimetric scales, v. sol. CS, CHCl3, benzenc, and ether.

Picric acid compound $C_{10}H_2BrC_6H_2(NO_2)JOH.^{-1}$ [79"] (R.); [86°] (B.); S. (alcohol of 92 p.c.) 6 at 20° (R.).

Di-bromo-naphthalono C₁₀H_aBr₂ Formed in small quantity by brominating naphthalene (Jolin, Bl. [2] 28, 514; not observed by others).

o Di-bronto-naphthalene $C_{10}H_{o}Br_{2}\{1;2\}$. [63°]. From (1, 2)-broma- (β) -naphthylamine by the diazo-perbromide reaction (Meldola, C. J. 43, 5). Oblique rhombic prisms (from alcohol, acetono or petroleum).

m-Di-bromo-naphthalene $C_{10}H_8Br_2$ [1:3]. [61°]. From di-bromo-(a)-naplithylamine, [119°] by removal of N11, (Meldola, C. J. 43, 2).

Needles. Di-bromo-naphthalene Cull, Br2 [c. 683]. Formed, together with two isomerides, [81°] ar I [15d'] by the action of Br (2 mols.) on naphthalene (I mol.) (Guareschi, G. 7, 24). Also from bromo-(β)-naphthol and PBr₃(Canzoneri, G. 12, 425). Prisms (from alcohol).

 $(\alpha\beta')$ -Di-bromo-naphthalene $C_{i0}H_aBr+1:2'or3'$]. [740]. From (1, 2'or 3', 2)-di-bromo-naphthylamine by diazo- reaction (Meldola, C. J. 17, 513). Silvery scales (from dilute alcohol).

 η -Di-bromo-naphthalene C_mH₁Br₂[1:x], [77°]. Formed, together with the isomeride (1)^{20°} , by brominating napla halene (a) sulphonic acid (Darmseidter a. Wichelhaus, A. 152, 301).

(β)-Di-bromo-naphthaiene C1011, Br2 [1:4]. [82]. (310). S. 93.5 per cent. alcohol) 1.33
 at 114°; 6 at 56° (Guarczeli, A. 222, 269).
 Formation.—1. The chief product of the

action of bromine (2 mols.) on naphthalene (Glaser, A. 135, 40).—2. By distilling (a)-bromo-511). - 3. From acctyl-(a) naplethylamine by brominating, saponifying, and treating the resulting C, 11 Br(N112) by the diazo-reaction (Meldola C. J. 43, 4).

Properties.-Long needles. Oxidised by HNO, to di-bromo-phthalic and bromo-nitro phthalic acids and bromo-nitro-naphthalene CrO, in acetic acid gives di-bromo-naphtho quinone and di-bromo-phthalide. Reacts with Br forming C₁₀H₄Br₂ [173°] (Guareschi, G. 16, 141).

Constitution.—This follows from the oxidation to di-bromo phthalic acid, coupled with the observation that the bromo-(α)-naphthylamine from which it may be formed (v. supra) gives (α)-bromo-naphthalone by the diazo-reaction.

(γ) Di-brome-naphthaleno $C_{10}H_0Br_2$ [1:1'or4']. [131° cor.]. (326°). S. (93.5 p.c. alcohol) 2 at 56°.

Formation.—1. By brominating naphthaleno (G.; Magatti, G. 11, 357).—2. From diazobromo-naphthaleno (from bromo-naphthylamino [64°]) by adding hromine-water and warning the pp. with HOAe.—3. The chief product of the action of Br on naphthalene (a)-sulphonic acid (Darmstädter a. Wichelhaus, A. 152, 303).—4. By the action of PBr, on (a)-di-nitro-naphthalene or hromo-naphthalene (a)-sulphonic sicid (1).

or hromo-naphthalene (a) sulphonic acid (J.).

Properties.—Tables. 11NO, gives bromonitro-phthalic acid. CrO₂ in HOAc gives bromophthalic acid [176]. Does not form a tetrabromide with Br.

 δ -Di-bromo-naphthalene $C_{to}\Pi_{\delta}Br_{2}$. [141°]. From naphthaleno (α)-sulphonic acid and PBr_{δ} (J.). Thin platos.

ε-Di-bromo-naphthalene C₁₀H_cBr₄. [160°]. From (α)-bromo-naphthalene sulphonic acid and PBr₄ (J.).

Di-brome-naphthalene tetra-chloride $C_{io}\Pi_aBr_zCl_*$. [156°]. From di-brominated naphthalene (? [82°]) and Cl (Laurent).

Di-bromo-naphthalene tetra-bromide $C_{10}H_aBr_a$. A mixture of three bodies of this composition, [c.100°][190°] and [173°] is formed from naphthalene and Br (G.).

Tri-brome-naphthalene C₁₀H₃Br₃. [75°]. Formed by brominating naphthalene, or by heating di-brome-naphthalene tetra-bromide with alcoholic KOH (Laurent, A. Ch. [2] 59, 196; Glaser, A. 135, 43). Needles (from alcohol).

Tri-bromo-naphthalene C₁₀H₃Br₃ [1:4:1']. [85°]. From di-bromo-nitro-naphthalene [117°] and PBr₃ (Joliu). Needles.

Tri-bromo-naphthalenc C₁₀H₅Br₃ [87°]. From di-bromo-naphthalene (\$\textit{\beta}\$)-sulphonic acid and PBr, (J.). Needles.

Tri-bromo-naphthalone C₁₀H₃Br₃ [1:3:1'or4']. [105°]. From (3, 1' or 4', 1)-di-bromo-naphthylamine [102°] by d. zo-reaction (Meldola, C. J. 47, 516).

Tri-bromo-akahtbalene C_{1e}II.Bf₃ [1:3:2'or3']. [110°]. From (1, 2' or 3', 3)-di-bromo-naphthylamine by the diazo-reaction. Needles (from alcohol) (Meldola, C. J. 47, 513).

Tri-bromo-naphthalone C₁₆H₂Br₃ [1:2:4].

[111]. I'rom C₁₆H₄(NH) 'Br₂ [1:2:4] by diazoreaction (Meldola, C. J. 43, 4). Formed also by heating C₁₆H₂(NH₂)(NO₂)Br [1:2:4] with cone. HBrAq and glacial 190Ac at 130° (Prager, B. 18, 2163). White needles (from dilnte C₂H₄O₂). Dilute IINO₃ at 180° gives phthalic acid.

Tetra-bromo-naphthalene C₁₀H₄Br₄ [1:4:2':3']. [175°]. S. (95 p.c. alcohol) '5 at 78°. From di-bromo-naphthalene tetrabromide [173°] and NaOEt (Guareschi, G. 16, 141). Needles (from alcohol) or plates (by sublimation). Cr0, in HOAc gives di-bromo-phthalide [188°] and tetra-bromo-(a)-naphthoquinone [224°].

Tetra-brome-naphthalene $C_{10}H_1Br_4$. [120°] From di-brome-naphthalene tetrabromide [100°] and NaOEt (Gn.). Needles (from alcohol).

Tetra-bromo-naphthalene tetra-bromide $C_{i0}H_4Br_b$. [173°]. From (1, 4)-di-bromo-naphthalene and Br (Gu.).

Ponta-bromo-naphthalene C₁₀II₃Br₃. From C₁₀II₄Br₄ and Br at 150° (Glaser). Granules, insol. alcohol.

Hexa-bromo-naphthalene C₁₀IL_BBr_e. [252°]. From naphthalene, Br, and I at 400° (Gessner, B. 9, 1505). Also from naphthalene (20 g.), Al.Cl₀ (15 g.) and Br (300 g.) (Roux, Bl. [2] 45, 515). Noedles; easily sublimed. Does not combine with pieric acid.

BROMO-NĀPHTHALENE DI-CARBOXYLIC ACID C₁₂H,BrO₄ i.e. C₁₀H₅Br(CO₂H)₂. [210°]. From bromo-acenaphthene and CrO₅ (Blumenthal, B. 7, 1095). Needles (from benzene). Converted by NH₃ into the imide C₁₀H₅Br(CO)₂NH [above 255°].

BROMO-NAPHTHALENE-(β)-SULPHINIC ACID C₁₀H₀BrSO₂Π. From naphthalene (β)-sulphinic acid and Br (Gessner, B. 9, 1503).

(a)-BROMO-NAPHTHALENE SULPHONIC ACID C₁₀II_Br(SO_III) [1:4]. [139°]. Formed by sulphonating (a)-bromo-naphthalone (haurent, Compt. chim. 1849, 392; Darmstädter a. Wichelhaus, A. 152, 303; Otto, A. 147, 184). Flat needles. Oxidised by KMnO₁ to phthalic acid (Meldola, B. 12, 1964). Potash-fusion gives no bromo-naphthol (M.). Br gives chiefly C₁₀II_Br₂ [82°]. — CaA'₂ 3aq. — BaA'₂ 2aq. — PhA'. 1 ag.

PbA'₂ 1'₂aq. Chloride $C_{10}H_0Br(SO_2Cl)$. [87°]. (Jolin, Bl. 28, 516). In its preparation there is also formed $C_{10}H_0Cl(SO_2Br)$ [116°] (Gessner, B. 9, 1504).

Bromide C₁₀H₀Br(SO₂Br) [115°] (J.).

Amide C₁₀H₆Br(SO₂NII₂) [190°] (J.); [195°]

(O.).

Bromo-naphthalene (a)-sulphonic acid

C_{1a} ¹⁷ Br(SO₃II). [104°]. Formed by brominating naphthalene (a)-sulphonic acid (D. a. W.).

PBr_s gives di-bromo unphthalene [131°].—KA'.

Chloride C₁₀H Br(SO Cl) [90'] (J.). Amide C₁₀H Br(SO NH.) [205'] (J.).

Bromo-naphthalene (β) -sulphonic acid $C_{1a}H$, $Br(SO_a\Pi)$. $[62^{\circ}]$. Formed by brominating naphthalene (β) -sulphonic acid (D, a, W). Crystalline mass, sol. ether (difference from the two preceding acids).—KA'.

Bromo-naphthalene sulphonic acid C₁₀11,Br(SO₂11). Formed in small quantity in preparing its isomerido [139°] by sulphonating (a)-bromo-naphthalene with H₂SO₄ or ClSO₃II (Armstrong a. Williamson, C. J. Proc. I, 234).

Chloride C₁₀H_aBrSO Cl [151].

Di-bromo-naphthalene (β)-sulphonic acid
C₁₀H_aBr_a(SO₄H). Formed by brominating naphthalene (β)-sulphonic acid (J.). Crystalline.
PBr, gives tri-brono-naphthalene [87°].
Chloride C₁₀H_aBr_a(SO₂Cl) [109°].
Achide C₁₀H_aBr_a(SO₂NH_a) [238°].

Actide C₁₀H₃Br₂(SO₂NIL₂) [238°].
Di-bromo-naphthalene sulphonic acid
C₁₀H₂Br₂(SO₂II). Got by sulphonating dibromo-naphthalene (Laurent, A. 72, 299).—KA'₂
—B₃A'₃.

BROMO-NAPHTHALIC ACID v. Bromo-oxy-(a)-NAPHTHOQUINONE.

BROMO (α)-NAPHTHOIC ACID C₁₁H₇BrO₂ i.e. C₁₀H₆Br.CO₂H [1:4']. [246°] (Eketrand, B. 19, 1135). Produced from its nitrile or by brominating (a)-naphthoio acid (Hausamann, B. 9, 1516). White needles (by sublimation). - KA' laq. - CaA'. I laq. S. I 5 at 20°. - BaA', 3uq. S. 1.7 at 21°. - AgA'.

Amide C, HeBr.CONH,: [211°]; flat needles. Nitrile C, HeBr.CN. [147°]. From (a)-Nitrile C, H, Br.CN.

naphthonitrile in CS, and Br.

Bromo (8)-naphthoic acid C₁₀H_eBr.CO₁H. [256°]. From (\$) naphthoic acid and Br (II.). Needles (by sublimation). KA' 2 aq. CaA', 3aq. O2 at 20°. —BaA', 3aq.
 Nitrite C₁₀H, Br.CN: [149]; that needles.

Tri-bromo-(β)-naphthoic acid C₁₀11,Br_s.CO₁11. [270°]. From (3)-naphthoic acid (1 mol.), Br (3 mols.), and I at 350°. Needles (by sublimation). BaA', (H1.).

Tetra-bromo-(a)-naphthoic anid

C₁₀H_aBr_bCO₁H. [239°]. From (a)-naphthoic acid (1 mol.) and Br (4½ mols.) at 350° (H.). Granules (from alcohol) or needles (by sublimation).-BaA',.

Tetra-bromo-(β)-naphthoic acid

C₁₀H₃Br₁.CO_.H. [260°]. Preparation and properties similar to those of the preceding acid (II.).

BROMO-(a)-NAPHTHOL. Ethyl ether C₁₀H_oBr(OEt). [18°]. From ethyl bromo (α)-naphthol and Br (Marchetti, C. N. 40, 87). Prisms, v. sol. ether.

Bromo - (B) - naphthol CtoH, Br(OH) [3:2] ?. [84°]. Prepared by adding Br in glacial acetic acid slowly to naplithol in glacial acctic acid (A. J. Smith, C. J. 35, 789). Needles. Sol. alcohol, ether, light petroleum and benzene. At 130° it begins to decompose, giving off 11Br. Oxidised by alkaline KMnO, to phthalic acid. PBr₅ gives di-bromo-naphthalene [68] and (β)bromo-naphthaleno (Canzoneri, G, 12, 424).

Acctyt derivative C10HaBr(OAc). (215)

at 20 mm. (C.).

Nitroso- derivative Cull (NO)Br(OII).

[65°]: green needles.

Di-bromo (α) naphthol C₁₀H₃Br₂(OH) [1:3:4]. [106°] (Fittig, A. 227, 244). Formed by brominating (a)-naphthol in HOAc (Bicdermann, B. 6. 1119) and in small quantity from di-bromo-(a)naphthylamine by the diazo- reaction (Meldola, C. J. 45, 161). Long needles (from alcohol). Powerful oxidising agent.

Reactions. -1. KMnO, gives phthalic acid. -2. Alcoholic KOH gives tri-oxy-naphthalene. --3. Combines with aniline forming a white crystalline salt. If this is beated for 10 minutes at 200°, and then allowed to cool, crystals of

 $C_{10}H_{\bullet}(NPhH) < \stackrel{\mathrm{OPh}}{N}$ or (β) -naphthoquinone dianilido (q. v.) are got (Meldola, C. J. 45, 156) .-- p-Totaldine forms the corresponding (β)-naphthoquinono di-toluide (q, v_*) . -5. (β)-naphthylamine forms the corresponding (β)-naphtho-

quinone di-naplithalide (q. v.).
Tetra-bromo-(\beta)-naphthol O₁₀H₄Br₄OH.[156'].
Prepared by adding excess of bromine to (\beta)naphthol dissolved in glacial acetic acid (A. J. Smith, C. J. 35, 791). White needles (from glacial acctic acid). Sol. CS, benzene and alkalis. Oxidised by KMnO, and COH to bromephthalio acid (anhydride [125°]). Hence it is C IIBr (C H BrOU).

Penta-bromo-(a)-naphthol C10H2Br2.OH [2:4:1':3':4':1]. [239°]. Formed by bromination phthalide.

of (α)-naphthol in presence of Al₂Br_a. Slender felted needles. Sl. sol. benzenc, xylene, and cumene, nearly insol. alcohol and ether. Dissolves in alkalis. By dilute HNO, at 100° it is oxidised to tetra-bromo-(a)-naphthoquinone [265°]; at 150° it is oxidised to di-bromo-phthalic acid [206]. — C₁₀H₂Br₂.ONa: long easily soluble needles. CulliBr OK: small colourle's needles (Blümlein, \hat{B} , 17, 2485).

Penta - bromo - (6) - naphthol C to H.Br. (O11). [237°]. Formed by bromination of (β) naphthol in presence of Al₂Br_a. White needles. Insol. alcohol, sl. sol. benzenc. It is oxidised by HNO₂. to tetra-bromo-(8)-naphthoquinone; on further oxidation it yields tri-bromo phthalic acid Cioll Br (ONa): long white silky needles (Flessa,

B.~17,~1479).

Bromo-(B)-naphthol (a)-sulphonic acid

 $C_{i_0}H_iBr(Off)SO_iH_i = [3:2:1]?$

Salts .- Formed by adding the calculated quantity of bromine to saturated solutions of the salts of (β) -naphthol (α) -sulphonic acid (Armstrong a. Graham, C. J. 39, 137).—KA'. S. 4 at 15 . Boiling HNO, forms phthalic acid. —CaA', raq.

BROMO-(a)-NAPHTHOQUINONE

Anilide C₁₆11,Br(NI1C₆11_)O₂. Formed by the action of anilino on bromo-oxy-(a)-naphthoquinone [197] in acetic acid solution. Red prisms. Sol, hot alcohol and hot acetic acid. By cold aqueous NaOH it is split up into its constituents (Baltzer, B. 14, 1902).

An isomeric anilide C10 II Br (NPh II)O2 [2:3:1:1] [194°] is formed by boiling di bromo-(a) naphthoguinone [218] with an alcoholic solution of anilinc. It is converted by KOII into bromo-oxy-(a)-naphthoquinone [202] [Miller, Bl.

[2] 43, 125). p-Bromo-anilide C, II, Br(NIIC, II, Br)O.

[240]. Prepared by bromination of (α)-naphthoquinonc-anilide, or by boiling a mixture of pbromo-anilin and bromo-oxy (a)-naphthoquinone with acetic acid. Red needles. Sol. benzene, sl. sol. alcohol. By alcoholic H.SO, it is decomposed into bromo-oxy-(a)-naplithoquinone and pbromo-aniline (Baltzer, B. 14, 1901).

Bromo-(8)-naphthoquinone C,II, CO.CO CH:CBr>.

[178°]. Obtained by bromination of (β) -naphthoquinone in acctic acid. Red needles or prismatio crystals. M. sol. warm alcohol, benzena, and acctic acid. Sublimable. Dissolves in dilute caustic aikalis with a brownish red colour, forming | hromo - oxy - (a) - naphthoquinone $[196^{\circ}]$ (Zincke, B. 19, 2495).

Di-bromo - (a)- naphthoquinone C10114Br2O,. [151°]. S. (alcoho!) 98 at 13'. Formed by the action of Br (7 pts.) and I (2 pts.) on (a)naphthol (1 pt.) in presence of water (Diehl n. Merz, B. 11, 1065). Yellow needles; may be sublimed. Alkalis from HBr and bromo-oxy-

naphthoquinone.

Di - bromo - naphthoquinone C1011, Br. O2 [f71°-173°] [f:4:1':4']? S. (95 p.c. alcohol) it at 16°. From di-bromo-naphthalene [82°], CrO, and glacial acctic acid (Guareschi, A. 222, 279). Yellow needles (from alcohol). Insol. water. Cannot be sublimed. Volatile with steam. CrO₃ does not oxidise it to di-bromo-Volatile with

Di-bromo-(8)-naphthoquinone C₁₀H₁Br₂O₂ [1:2:3:4]. [174°]. Obtained by the action of bromine upon bromo-(8)-naphthoquinone in hot acetic acid, or better upon (a)-amido (β)-naphthol or its sulphate. Thick red plates or tables. Sl. sol, alcohol and ether (Zincke, B. 19, 2196).

Di-bromo-naphthoquinone Cio [I, Br. O., [218]] From (α) naphthoquinone, Br, and I (Miller, Bl. [2] 38, 138). Prisms. Converted by KOH into

bromo oxy naphthoquinone [200°].

Anilide [194°]. Tetra-bromo-(α)-naphthoquinono C₁₀H₂Br₁O₂. [265°]. Yellow plates. Sl. sol. alcohol. Formed by oxidation of penta-bromo-(a)-naplithol with dilute IINO, at 100°. By further exidation at 150° it yields di-bromo-phthalic acid [206°] (Blümlein, B. 17, 2488).

Tetra-bromo-[a]-napl thoquinone CioII Br.O. [1':4':2:3:1:4]. [221°]. Formed by oxidising tetrabroino - nuphthalene [173°]. Orange - yellow prisms, v. sol. hot water (Guareschi, G. 16, 149).

Tetra-bromo-(β)-naphthoquinone C10 II, Br, O2. [164°]. Red granular crystals. Sl. sol. alcohol. Formed by exidation of peuta-brome-(B)-naphthol with dilute IINO, By further oxidation it is converted into tri-bromo-phthalic acid (Flessa, B. 17, 1481).

BROMO-NAPHTHOSTYRIL v. Inner anhydride of Bromo-amido naphthoic acid.

DI-BROMO-(aa)-DINAPHTHYL [215°]. From di-naphthyl and bromine-vapour (Lossen, A. 114, 77). Monoclinic prisms; v. sl. sol, alcohol.

Hexa-bromo-(a2)-dinaphthyl C20H3Brg. Rosin. C.,H.Br,. Henta-bromo- $(\beta\beta)$ -dinaphthyl Amorphous (Smith a. Poynting, C. J. 27, 851).

BROMO-NAPHTHYLAMINE C₁₀H_cBr(NII).

By reduction of bromo-nitro-naphthalene [85] of Jolin, itself got from (a)-bromo-naphthaleno by nitration. An oil. May be distilled with steam. Fe₂Cl_s gives a violet colour in its aqueous solution. Reduces AgNO, (Guareschi, A. 222. 299). Pessibly identical with Rother's (1, 4)bromo-naphthylamine [94°].

m-Bromo-(α)-naphthylamine C₁₀H₃Br(NH₂) [3:1]. [62°]. From the mitro- compound, zincdust, and HOAc (Meldola, C. J. 47, 509). Needles (from dilute alcohol).

Acetyl derivative C10HaBr(NHAc) [187]:

noedles.

o-Bromo-(3)-ne rathylamine C10Ha(Br)NI1 [1:2]. [63°]. Got by saponification of the acetyl derivative obtained by bromination of acetyl (8)-naphthyland no [131°]. Small white needles. Volatile with steam. It is a neutral body. Dilute HNO₃ gives phthalic acid (Meldola, C. J. 4B, 6). Whon NH, is displaced by II (α)-bromo-naphthalene results (M.).

C10H6Br.NH(Ac) Acetyl derivative

[135°]. Needles (Cosiner, B. 14, 59).

Bromo-(α)-naphtbylamine C₁₀H₀Br(NH₂).
[1:1' or 4']. [64°]. From bromo-nitro-naphtthalenc [122-5°] (Guarcschi, A. 222, 297). Volatile with steam. Plates (from boiling water). May be sublimed. KMnO, forms c-brome-phthalic aoid [165°-172°].—B'IICl.

m-Bromo-(B)-naphthylamine C10 HaBr(NH2) [1:3]. [72°]. From Liebermann's bromo-nitronaphthalene by reduction (Meldola, C. J. 47, 509). Converted into (1, 3)-di-bromo-naphthal-

one by the diazo- reaction.

C, H Br(NHAo). Acetyl derivative [187°]: needles.

C,oH,Br(NH2) p-Bromo-(a)-naphthylamine [1:4]. [94°]. From its acetyl derivative and KOH (Rother, B. 4, 850; Meldola, B. 12, 1961). Needles. On oxidation it gives phthalio acid. By diazo-reaction it gives (a) bromo-naphthalene. Br gives di-bromo-naphthylamino [119°].

Acetyl derivative C. H. Br. NHAo. [1920]. Prepared by slowly adding HCl to a solution of bromine in NaOH in which is suspended acetyl-(a)-naphthylamine. White needles (Prager, B. 18, 2159).

Di.bromo-(α)-naphthylamine C₁₀H₂Br₂(NH₂) [3:2'or3':1]? [102']. From its acctyl derivative. Needles; sol. boiling dilute acids. Removal of NII, gives di-bromo-naphthalene

[74°] (? [1:1']). Acetyl derivative C10 II Br. (NHAc) 221°]. From acetyl (3, 1)-bromo-naphthylamine and Br (Meldoln, C. J. 47, 514).

Di-bromo (a)-naphthylamine

 $C_{10}H_sBr_s(NH_s)$ [1'3:1]. [105°]. From its acetyl derivative. Needles. Oxidised by dilute HNO, to c-brome-phthalic acid [174°]. Converted into di-bromo-naphthalene [74°] by diazo- reaction.

A ectyl derivative C₁₀H, Br. (NHAe).

[222°]. From bromo-naphthylamine [62] by acetylation and bromination (McIdola, C. J. 47,

512; C. J. Proc. 1, 173).

Di bromo-(a)-naphthylamine C₁₀H.Br.(NIL) [1:3:4]. [119°]. From the acetyl derivative by hot aqueous KOII. Needles; does not combine with acids. Oxidation gives phthalic acid; the diazo- reaction gives (1, 3)di-bromo-naplithalene.

Acetyl derivative C10H Br. (NHAe). [225 ']. Formed by bromination of acetyl-(a). naphthylamine (Meldola, B. 12, 1961).

Li-bromo-(B) naphthylamine C1011 Br2(NH2) [121°]. Long colourless needles. Formed by the action of bromine in acctic acid solution upon (β)-naphthalene-azo-(β) naphthylaminc. Acetyl derivative: [2086] (Lawson, B. 18, 2121).

Tetra-bromo-naphthylamine

Acetyl derivative C10 II Br NllAe. [138°]. Formed by brominating acctyl-(1,2)-bromo- (β) naphthylamine in acctic acid solution (Meldola, C. J. 43, 8). Minute needles (from alcohol). Could not be saponified.

BROMO-NAPHTHALENE-DIAMINE.

Acetyl derivative CullsBr(NH.)(NHAc) [2:4:1] [c. 222°]. From brono-nitro-acetnaphthalide. Is not basic (Meldola, C. J. 47, 499).

Di-bromo-(1:1'or4')-naphthylono-diamine $C_{10}H$ Br(NII_2). From naphthylene-diamine hydrochloride (from (α) -di-nitro-naphthalene) and bromine-water (Hollemann, Z. 1865, 556).

DI-BROMO-(α)-DINAPHTHYLENE-OXIDÉ $C_{20}\Pi_{10} Br_2 O.~[287^\circ]$. Light-yellow crystals. Sl. sol. benzene and acetic acid. Prepared by the nction of Br on a CS₂ solution of (α)-dinaphthylene-oxide (Knecht a. Unzeitig, B. 13, 1725). $Di-bromo-(\beta)$ -dinaphthylene-oxide

C₂₀H₁₀Br₂O. [247°]. Yellow needles. Prepared by the action of bromine on a CS₂ solution of (B)-dinaplithylene-oxide (K. a. U.).

DI-BROMO-DI-NAPHTHYL-METHANE C.H., Br., [193°]. From di-naphthyl-methane anu Br (Grabewski, B. 7, 1605). Needles (from alcohol-benzene); not affected by boiling alcotrolic KOH.

BROMO-NICOTINE v. NICOTINE.

BROMO-O-NITRO-ACETOPHENONE O_sH_sBrNO_s i.e. [2:1] C_sH_s(NO₂).CO.CH₂Br. [56°]. By bromination of C_sH_s(NO₂).CO.Mo (Gevekoht, A. 221, 327). Needles (from benzoline).

Bromo-m-nitro-acetonhenone

[3:1] CaH (NO2).CO.CH2Br. Nitro-phenyl bromomethyl ketone. [96°]. Formed by nitrating brome-acetophenone. Needles (from dilute alcohol); v. sl. sol. ether. Oxidation gives mnitro-benzoic acid (Hunnius, B. 10, 2008).

Di-bromo-o-nitro-acetophenone C.H.(NO.).CO.CIIBr... [86]. Prepared like the above (G.). Attacks the cycs. Prisms (from

bonzoline). ω-Di-bromo m.nitro-acetophonono

C_aH₄(NO_a).CO.CHBr₂, m-Nitro-phenyl di-bromo-methyl ketone. [59°]. Formed by careful nitration of ω-di-bromo-acetophenone; or by further bromination of ω bromo-m-nitro-acctophenone. Yellowish tables. V. sol. most solvents (Engler a. Hassenkamp, B. 18, 2210).

BROMO-NITRO-O-AMIDO-BENZOIC ACID

C_aH_aBr(NO_a)(NII_a)CO_aH [1:27:4:5]. [272]. Formed by treating nitro-isatoic acid at 100° with bromine dissolved in glacial acetic acid (Dorsch, J. pr. [2] 33, 40). Long yellow needles. Sol. hot water, acctone, alcohol, glacial acctic acid and ether. Insol. chloroform and benzene.

Di-bromo-nitro-o-amido-benzoic acid C_aHBr₂(NO.)(NH₂)CO₂H. [c. 203°]. From nitro-isatoic acid and bromine in glacial acetic acid at 100° (D.). Plates. Sol. acctone, alcohol and glacial acetic acid, less sol. benzene, chloroform,

and other, insol. water.

Tri-bromo-nitro-o-amido-benzoic acid C_sBr_s(NO_{..})(NH_{..})(CO_{..}H). [196°]. From nitroisatoic acid and bromine (D.). Needles (ppd. by adding water to its solution in acctone). acetone, alcohol, ether, glacial acetic acid, ch'oroform and benzene.

(3:5:4:1) - BROMO-NITRO-AMIDO-PHENYL. ACETIC ACID C.H.(Br)(NO.)(NII.).CII..CO.II [3:5:4:1], [192°]. Prepared by saponification of the acetyl derivative of (3:5:4:1)-bromo-mitroamido-benzyl cyanide (Gabriel, B. 15, 1994). Long yellow needles. Sol. hot alcohol, ether and acetic acid, sl. sol. cold water, benzene, and chloroform.

Nitrile

 $C_aH_2(Br)(NO_2)(NH_2).CH_2CN$ [3:5:4:1]. nitro-amido-benzyl cyanide. Acetyl derivative: [191°]. Formed by nitration of the acetyl derivativo of (3:4:1) bromo-amido-phenyl-aceto-nitrile (Gabriel, B. 15, 1992). Slender yellow needles, sol. alcohol and acetic acid, sl. sol. cold water, v. sl. sol. ether and CS2.

BROMO-NITRO-ANILINE

needles; gives m-bromo-nitro-benzene by diazo-Br forms di-bromo-p-nitro-anilino reaction. .[203°].

Benzoyl derivative CaH.B. (NO.)(NIBz). [160°]. From benzoyl-p-nitro-anilino and Br

(Johnson, B. 10, 1709).

Bromo-nitro-aniline C.H.Br(NO2)(NH2)[1:3:4]. [111]. S. 014 at 20°; S. (alcohol) 10.4.

Formation.—1. From nitro-p-di-bromo-benzene and alcoholic NH₄ at 165° (K.; Meyer a. Wurster, A. 171, 59).—2. By nitrating p-bromo anilino in glacial HOAc (Hübner, A. 209, 357).

Buth a time of alcoholic NH on the activation of the control of th 3. By the action of alcoholic NII, on the mothyl derivative of (1,3,4)-brome-nitre-phenel.

Properties .- Orange needles; may be sublimed; searcely basic. Converted by diazo-re-

action into m-bromo-nitro-benzene [56°].

Acetyl derivative C.H.Br(NO2)(NHAc). Formed by nitrating acetyl-p-bromoaniline (II.). Annuonia and zinc-dust reduce it to C_aH_aBr(NHAc).N_c,C_aH_aBr(NHAe) [282°] (Matthiessen a. Mixter, Am. 8, 317).

Benzoyl derivative Call, Br(NO2)(NHBz). [138°]. Formed by nitrating benzoyl-p-bromoanilino or brominating benzoyl-o-nitro-unilino (Meinecke, B. 8, 561; Johnson, B. 10, 1710).

Bromo nitro aniline

C. H. Br(NO.)(NH.) [1.2:4]. [132]. Formed by nitration of p-bromanilino dissolved in 10 pts. of H.SO. Flat plates. V. sol. alcohol, ether, acetic acid, and chloroform, v. sol. water. By further bromination it yields tri-bromo-nitrani-line [103] (Nölting a. Collin, B. 17, 266).

Bromo-nitro-anilino

C_H_Br(NO2)(NH2) [1:4:5]. [151°]. Formed by the action of alcoholic NH3 upon (1.5.1)-dibromo-nitro-benzene [62°] or on (1,4,5)-bromo-di-nitro-benzene [56°] (K.; Wurster, B. 6, 1542). Orange needles. Gives by diazo-reaction p-bromonitro-benzene. Is not basic. Dilute 1180, (S.G. 1.38) slowly forms bromo-di-nitro-phenol [81°].

Bromo-di-nitro-aniline C.H.Br(NO.) (NH.) [1:3:5:6]. [154] (La); [144°] (K.). Formed by brominating distitre-aniline (K.), or by heating di-nitro methyl-amiline with HOAc and Br (Leymann, B. 15, 1231). Yellow needles. Converted by boiling KOII into bromodi-nitro-phenol (118°)

Bromo-di-nitro-aniline C.H.Br(NO.) (NH.). [160°]. From di-bromo-di-mitro-benzene [100°] and alcoholic NH3 at 100° (Austen, B. 9, 919). Orange scales.

Bromo-di-nitro-aniline C.H.Br(NO.).(NH.). [178]. From di-bromo-di-nitro-benzene [117] and alcoholic NH, (K.).

Bromo di nitro anilino. Benzoyl derivative C61LBr(NO2) (NHBz) [1:3:5:4]. [221°]. Small needles; formed by nitrating benzoyl bromo-nitro-aniline C₆H₃E (NO₂)(NHBz) [1:34], or benzoyl di-bromo-aniline (Johnson, B. 10, 1710).

Bromo di-nitro aniline. Benzoyl derivative Callabr(NO.)2(NHBz). [196°]. Formed by nitrating benzoyl-p-bromo-aniline (Meinecke, B. 8, 561), is probably ilentical with the precoding.

Di-bromo nitro aniline C.H.Br. (NO2)(NH2). [75°]. From di bromo-di-nitro-benzene [159°] and alcoholio NII, at 100° (Austen, B. 9, 622). Red necdles.

Di.bromo-o nitro-aniline C.H.Br. (NO.) (NH.) [1:3:5:6]. [127°] (Hentschel, J. pr. [2] 34, 426). Formation.—1. By brominating o-nitro-aniline or (1,3,4) bromo-nitro-aniline -2. By the action of alcoholic NH3 on (1,3,4,5)-tri-bromonitro-benzeno or the methyl other of (1,3,5,6)-dibromo-nitro-pheuol (K.).

Properties .- Orange needles. Acetyl derivative C.H.Br. (NO.)(NIIAc). [209°]. From acetyl-di-bromo-aniline by nitra-tion. Needles, v. sol. alkalis (Remmors, B. 7, 348).

Benzoyl derivative Coll.Br.2(NO2)(NHBz) [1:3:5:6]? [195]. Formed by brominating benzoyl-o-nitro-anilino (Johnson, B. 10, 1710). Yellow needles.

Di-bromo-p-nitro-aniline CaH_Br_(NO2)(NH2)

[1:3:5:2]. [207°] (L.); [203°] (K.).

Formation .- 1. By brominating p-nitro aniline or (1,3,6)-bromo-nitro-aniline. - 2. By the action of alcoholic NH, on tri-bromo-nitro-benzene [112°] or the methyl ether of (1,3,5,2)-dibromo-nitro-phenol (K.). - 3. From tri-bromoaniline in HOAc by warming with conc. HNO, for a short time (Losanitsch, B. 15, 474).

Properties.-Thin yellow needles. Displacement of N11, by I1 gives tri-bromo-benzene [112].

Tri-bromo-nitro-aniline CaHBra(NO.)(NH.) [1:3:5:4:6]. [103°]. From m-nitro-aniline and bromine-vapour (K.), or from (1,2,1)-bromonitro-aniline and Br in HOAc (Nölting a. Collin, B. 17, 266). Pale greenish-yellow needles in stellate groups; v. sol. alcohol. Converted by diazo- reaction into (1,3,5,1)-tri-bromo-nitrobenzene.

Tri-bromo-nitro-anilino CaHBr3(NO2)(NH2) [1:3:5:4:6]? [215°]. From its acetyl derivative (Remmers, B. 7, 351). Flat yellowneedles; sl. sol. alcohol. This body might be expected to be identical with the preceding.

Acetyl derivative C. 11Br. (NO.) (NHAc). [232°]. I'rom acetyl-(1,3,5,6)-tri-bromo-anilino (R.). Needles.

Di-acetul derivative Calibra (NO2) (NAc.). Formed by nitrating di-acetyl-tri-bromo-aniline.

Tri-bromo nitro-aniline C. HBr. (NO.)(NH.) [1:2:3:5:6]. [161°]. From (1,2,4)-bromo-nitroaniline [151°] and bromine vapour (K.), Lemonyellow needles (from alcohol). Converted by dinzo- reaction into (1,2,3,5)-tri-bromo-nitrobenzene [112°].

BROMO-NITRO-ANTHRAQUINONE

Cull_(NO.)(Br)O. [261° uncor.]. Prepared by nitration of tetra-bromo-anthracene. White needles. Sublimable. Sol. acetic acid, sl. sol. alcohol, ether, and chloroform. On reduction it gives amido-anthraquinoue (Claus a. Hertel, B. 14, 980).

Bromo-di-nitro-anthraquinone

 $C_{14}H_2Br(NO_2)_2O_2$. [213° uncor.]. Prepared by nitration of tri brown-anthracene with a mixture of funning IINO, and funning ILSO, Yellow needles. Sol. henzeno, chloroforis, and acctio acid, m. sol. alcohol and ether (Claus a. Diernfelluer, B. 14, 1333).

Di-bromo-nitro-anthraquinone

C, H Br (NO.)O. [215° uncor.]. Prepared by nitration of tetra-bromo-anthracene, Sublimable. Fine yellow needles. V. sol. hot acetic acid, less in alcohol and ether. On reduction with sodium-amalgam it gives amido-anthraquinone. Di-bromo-di-nitro-anthraquinone

C14H1Br2(NO2)2O2. [239° uncor.]. Prepared by nitration of tetra-bromo-anthracene with a mixture of fuming 11₂SO₄ and fuming HNO₃. Needles. Sol. acetic acid, benzene and chloroform, sl. sol. alcohol and ether (C. a. D.).

Tetra-bromo-di-nitro-anthraquinone C₁₄H₂(NO₂)₂Br₁O₂. [105°]. Prepared by nitration of dibrono anthracene-tetrabromide. Sol.

sublimable. On reduction it gives (a)-diamidoanthraquinone (Claus a. Hortel, B. 14, 981).

p BROM D-0-NITRO-BENZALDOXIM
C₇H₄N₂O₄Br i.e. [2:4:1] C₄H₄(NO₂)(Br)(CH:NOH)
[153°]. Prepared by heating o-nitro-p-diazobenzaldoxin with HBr (Gabriel a. Meyer, B. 14, 827). Fine needles. Sol. alcohol, ether, acetic acid, and hot benzene.

o-BROMO-NITRO-BENZENE CaH,Br(NOa) [1:2]. Mol. w. 202. [41°] (F. a. M.); [43°] (K.). (261° i.V.). Formed, together with a much larger quantity of the ρ -isomeride, by nitrating bromo-bonzone (Hübner a. Alsberg, A. 156, 316; Z. [2] 6, 369; Walker a. Zincke, B. 5, 114; Fittig a. Mager, B. 7, 1179). Yellow needles, more sol. alcohol than the p-isomeride.

Reactions. -1. SnCl. reduces it to o-bromoaniline.-2. Alcoholio NH, at 190 gives o-nitroaniline (W. a. Z.).—3. HNO₃ gives bromo-dinitro-benzene [72].—1. KCN and alcohol at 190° gives Call, Br.CN. -5. Hoating with KOHAq gives o-bromo-phenol.

m-Bromo-nitro-benzene C.II.Br(NO.) [1:3]. [56°]. (257° i.V.).

Form tion. -1. From m-nitro-aniline by the diazo- reaction (Gricss, T. 1861 [3] 712).— 2. From (1, 3, 4)-bromo-nitro aniline (Wurster, B. 6, 1513; 7, 416). -3. From nitre-benzene (10 g.), Fe Cla (1 g.), bromine (13 g.) in scaled tubes for 12 hours at 70° (Scheufelen, A. 231, 165). The yield is 80 p.c. of the theoretical .-4. By the action of a liet solution of cuprous bromido upou m-nitro diazo-benzene sulphate (from m-nitraniline) (Sandmeyer, B. 18, 1495).

Properties. -Yellow trimetric plates. Not attacked by KOHAq or alcoholic NII,

p-Brome-nitro benzene C.H.Br(NO.) [1:1].

[126°]. (256° i.V.).

Formation.—1. The chief product obtained by dissolving bromo-benzeno in fuming IINO, (Couper, A. 101, 226). -2. From p-nitro-aniline by the diazo- reaction. -3. From bromo- nitroaniline [151°]. -1. From bromo-benzene p-sulphonic acid and HNO₄ (Spiegelberg, A. 197, 257). 5. Formed by the action of precipitated Cu.O (1 mol.) upon p-bromo-diazo benzene nitrite (1 mol.) obtained by adding slowly a solution of 15 g. of NaNO, in 50 c.c. of water to a mixture of 17 g. of p-bromo-aniline, 20 g. HNO, (1:4), and 50 c.c. of water. The yield is small (Sandmeyer, B. 20, 1496).

Properties. -Long white needles, sl. sol. HOEt.

Reactions.—1. Resembles o-nitro-aniline in reactions 1 and 2 .- 2. Alcoholic KCN at 1903 gives m-bromo-benzonitrile (Richter, B. 4, 462). 3. Br at 250° gives p-di-, u-tri-, and s-tetra-bromo-benzenes (Ador a. Billiet, J. 1876, 370).

Bromo-di-nitro benzene C.H.Br(NO₂)₂ [1:3:4]. [59]. From m-bromo-nitro-benzene, IINO, and ILSO, (Körner, J. 1875, 332) Monoclinic plates (from ether-alcohol). Alcoholic NH, at 180° forms C, H, Br(NO.)(NII.) [1:4:3] [151°]. Boiling NaOMAq (S.G. 1-135) gives C.H.Br(NO.)(O11) [1:4:3] aud a little C_11 Br(NO_)(OH) [1:3:4] (Laubenheimer, B. 11, 1159).

Bromo-di-nitro-benzene CaHaBr(NO2)2 [1:2:4]. [72°]. From bromo-benzenc, furning HNO,, and H.SO, in the cold (Kekulé, A. 137, 167; Spiegelberg, A. 197, 257). Large yellow prisms. Alcoholic NII, forms di-nitro-auiline. KOHAq forms alcohol, ether, benzene, and acetic acid. Not di-nitro-phenol [111]. Sn and IICl gives itphenylene-diamine (Zincke a. Sintenis, B. 5, 791). Crystallises with benzene as (C.H.Br(NO2),2,C.H.

[65°]. Bromo-di-nitro-benzene C_oH₃Br(NO₂)₂. [87° Di-bromo-di-nitro-benzene [1580] is treated with alcoholic NH, at 100° and in the resulting C,H,Br(NO,),(NH,) hydrogen is substituted for NII, by the diazo-reaction (Austen, B. 8 1183). Not affected by alcoholic NII,

Di-bromo-nitro-benzene C.H.Br.(NO.) [1:2:4]. Mol. w. 281. [59°]. From o-di-bromo-benzene and HNO, (Riese, A. 161, 179). Monoclinic tables (Groth a. Bodewig, B. 7, 1563). May be reduced to di-bromo-aniline [80°].

Di-bromo-nitro-benzone C_aH_aBr₂(NO₂) [1:3:4]. Formed by nitrating m-di-bromo-benzcno (Meyer a. Stüber, A. 165, 176). Triclinic prisms (by sublimation, G. a. B.); volatile with steam. Converted by alcoholic into (1,4,3)-bromo-nitro-aniline. Converted by alcoholic NH, at 190° Reduction gives (1,3,4)-di-bromo-aniline. Alcoholic KCN at 250° gives the nitrile of di-bromo-benzoic aoid [2095] (R.).

Di-bromò-nitro-benzene C₈H₃Br₂(NO₂) [I:3:2]. 18**3°**1. Separates from the alcoholic motherliquors from which the preceding has crystallised. Prisms or lamino. Alcoholic NII, at 190° gives nitro-m-phenylene-diamino (Körner,

G. 4, 360).

Di-bromo-nitro-bonzene C, H, Br, (NO,) [1:4:5]. [85°]. From p-di-nitro-benzeno by nitration (Riohe a. Bérard, A. 133, 51). From m-bromonitro-benzene (14 g.), Br. (11.2 g.), and Fe Cla (4 g.) at 80° for 12 hours (Scheufelen, A. 231, 169). Yellowish-green tablets (from ether-alcohol). Alcoholic NII, at 210° gives (1,3,4)bromo-nitro-amiline. Sn and HCl gives p-di-bromo-aniline [51°]. Alcoholic KCN gives the nitrile of di-hromo-benzoic acid [152°]

Di-bromo-nitro-benzene CaHaBra(NOa) [I:3:5]. [105°]. From (1,5,3,6) or (1,5,3,2) di-bromonitro-uniline by diazo- reaction (Körner). Prisms or tablets (from other). May be reduced to di-

brome-aniline [57°].

Di-bromo-di-nitro-benzeno C.H.Br.(NO.). [H7°]. Formed by nitrating m-di-bromo-benzeno (Körner). Greenish-yellow needles, volatile with steam. Heating with KOHAq gives bromo-di-nitro-phenol [925].

Di-bromo-di-nitro-benzene C.H.Br.(NO.)... [58°]. Formed by nitrating o-di-bromo-benzene (Austen, B. 8, 1182). Prisms (from HOAc). By displacement of Br by NH₂ and H successively it may be converted into bromo-di-nitro-benzene [87°]

-benzene [37°]. C.H.Br.(NO.).. Di-bromo-di-nitro-benzene [120°]. Formed in small quantities in the pre-

paration of the preceding body (A.). C.H.Br.(NO.). Di-bromo-di-nitro-benzene

[159°]. Formed by nitrating p-di-bromo-benzeno (Austen, B. 9, 621). Small ncodles. Alcoholic NII, forms di-bromo-nitro-aniline [75°].

Di-bromo-di-nitro-benzen♂

 $C_8H_2Br_2(NO_2)_2$ [I:4:2:6]. [100°]. Formed in preparing the preceding (A.). Alcoholic NII3 converts it into bromo-di-nitro-anilino [160°].

Tri - bromo - nitro - benzene C. H.Br. (NO.) [1:3:4:6]. Mol. w. 360. [94°]. Formed by nitrating u-tri-bromo-benzene (Mayer, A. 137, 226). Palo yellowish green needles (from alco-Alcoholic NII, gives bromo-nitro-phol). phenylenc-diamine.

Tri-bromo-nitro-benzene

C₆H₂Br₅(NO₂) [1;2:3:5]. [112°]. From (1,5,3,6)-di-brome-nitro-aniline [203°] by displacing NH2 by Br, or from (1,2,3,5,4) tri-bromo-nitroanilino by displacing NH, by H (Körner). Triclinio crystals; a:b:c = 1.005:1:.4823 (La Valle, G. 10, 1). Reduction gives tri-bromo-aniline. Alcoholic NII gives di-bromo-nitro-aniline [203°].

Tri-bromo-nitro-bonzene

C_aH_aBr_s(NO₂) [1:2:4:6]. [120]. From (2,4,6,1) di-bromo-nitro-andine by dialo reaction (K-rner). Needles (from HOAc). Alcoholic NH2 gives the parent di-bromo-nitro-aniline.

Tri-bromo-nitro benzene C₀11₂Br₄(NO₂) [1:3:5:2), [125°]. (177°) at H min. Prepared by nitration of s-tri-bromo-bonzene with HNO, (1.5) (Wurster a. Berau, B. 12, 1821; cf. C. L. Jackson, B. 8, 1172). Formed also by diazo- reaction from (1,3,5,2,4)-tri-bromo-nitroanilino (Körner, G. 1, 122). Monoclinic prisms; a:b:c = 6518:1; 8695; η - 99°46' (Panebianco, G. 9, 354). Tin and HCl reduce it to ordinary tribromo-aniline. Alcoholic NII, at 170° gives (I,4,3,5)-bromo-nitro-phenylene-diamine.

Tri-bromo-nitro-benzene

C₆H₂Br₃(NO₂) [1:3:1:2]. [above 187°]. Formed in small quantity in preparing the isomeride [94°]. Sublimes at 187°.

Tri-bromo-di-nitro-benzene

C.HBr.(NO.). [1:2:1:3:5?]. [135°]. Formod by nitrating the preceding body (Mayer). Triclinic crystals; a:b:c--455:1: 457 (Panebianco, G. 9, 355). Alcoholic NII, gives bromo-di-nitrophenylene-diamine.

Tri-bromo-di-nitro-benzene

C₈HBr₃(NO_{.)}, [1:3:5:2:6], [192], Glistening needles. Prepared by nitration of s-tri-bromo-Glistening. benzene with IINO3 and H2SO4 (Wurster a. Beran, B. 12, 1821).

Tetra-bromo-nitro-benzene

C, 11Br, (NO.) [1:3:1:5:6]. [96° after several fusions]. Slender needles, [60°] (from alcohol). From u-tetra bromo-benzene by nitration. Formed also by heating C_u(NO₂)Br₄SO₃H with HCl (V. v. Richter, B. 8, 1427; Langfurth, A. 191, 202).

Penta-bromo-nitro-benzene C.Br. (NO.). (228°]. From u - tetra - bromo - benseue and fuming HNO, (R.). Monoclinic prisms (from benzene).

BROMO - NITRO - BENZENE SULPHONIC ACID U,H3Br(NO2)(SO3H) [1:1:2]. [1307-1350]. From brome-benzeno o-sulphonic acid and conc. HNO₃ (Editional, A. 186, 315, From p-bromonitro-benzene and fuming F. SO₄ (Augustin a, Post, B. 8, 1559). Flat yellow columns, v. o. sol. water. Reduction gives amido-bensene msulphonic acid; exchange of NO2 for Br gives p-di-bromo-benzene sulphonic acid. AgA'.--BaA', 5aq. S. (of BaA',) 5:3 at 16'.—CaA', 4aq.
—CaA', 6',aq (A. a. P.).—KA'.—NaA'.—NH,A'.
PbA', 5aq. ZnA', 7:q.
Chloride C,H,Br(NO.)(SO.Cl): [92°]; tables.

Amide $C_aH_a^{\prime}Br(NO_a)(SO_aNH_a)$. [205°].

Bromo-nitro-benzene sulphonic acid

C.H.Br(NO.)(SO.H) [1:6:2]? Formed in small quantity in preparing the above by nitratingbromo-benzene o-sulphonio acid (B.) .- BaA'. S. 156 at 8°. KA'.

Chloride C.H.Br(NO.)(SO.Cl): [97°]; tables.. Amide C, II, Br(NO2)(SO2NH2). [2150].

Bromo-nitro-benzene-sulphonio acid CaHaBr(NOa)(SOaH) [1:3:6]. Formed by heating nitro diazo benzene sulphonic acid (1:3:6) with HBr (Limpricht, B. 18, 2186).

Chloride: [75°], large yellow tables. Amide: [166°], small white tables.

Bromo-nitro-benzene sulphonic acid C_aH₃Br(NO₂)(SO₃H) [1:4:3]. Formed by nitrating brome-benzene m-sulphonic acid (Berndsen, A. 177, 95; Thomas, A. 186, 124). Reduced to amido-benzene o-sulphonic deid. Exchange of NO, for Br gives p-di-bromo-benzene sulphonic acid.—AgA' 1 aq.—BaA'..3aq. S. 24 at 7'.— CaA'..6aq.—KA'.—NH,A'.—I'bA'..3aq. Chloride • C. II.,Br(NO.)(SO.Cl). [83°].

Amide C₆H₃Br(NO₂)(SO₂NH₂). [170°].

Biomo-nitro-benzeno sulphonic aci-C.H.Br(NO.)(SO.11) [1:2:4]. Formed by nitrating brome-bezzene p-sulphonic acid (Goslich, A. 180, 93; Limpricht, B. 8, 456). Formed also by sulphonating o-bromo-nitro-benzene (A. a. P.; Andrews, B. 13, 2127). Reduction by HI at 120° gives amido benzene m-sulphonic acid.— BaA', aq. S. (of BaA',) 1.71 at 15° (A.).
—BaA', 1¹aq. S. (of BaA',) 1.46 at 9° (G.).—CaA', 2¹aq (A. a. P.).—CaA', 2¹aq. S. (of CaA',)
1.71 at 9° (G.).—CuA', 2¹aq.—KA'. S. 1.02 at 9° (G.). NH₄A'. S. 5.96 at 9°.—PbA', 2aq.— ZnA', 2aq.

Chloride CallaBr(NO2)(SO2Cl). [10°-50°] (A.); [57°] (G.).

Amide CallaBr(NO.)(SO.NH2): [177°]; plates. Bromo-nitro-benzene di-sulphonic acid

C₆H₂Br(NO₂)(SO₃H)₂. From nitro-benzeno msulphonic acid via di-nitro-benzene di-sulphonic acid and amido-nitro-benzone disulphonic acid (Limpricht, B. 8, 289). Trimetric tables (containing aq).

Di-bromo-nitro-benzene sulphonic acid From o.di. $C_6H_2Br_2(NO_2)(SO_3H)$ [1:2:4:6]. bromo-benzene sulphonic acid and fuming HNO₃ (Goslich, A. 186, 152). Reduction gives di-brome-aniline sulphonic acid.—15a\(\Lambda'_2\)3a\(\text{3a}\)4. S. 9 at 7°.—Ca\(\Lambda'_2\)4aq.—Ca\(\Lambda'_2\)6aq.—K\(\Lambda'_2\) NH,A'.—PbA', 3aq. S. 8 at 11°. Chloride C,H.Br.,(NO.,)SO,Cl.

Amide C₆H₂Br₂(NO₂)SO₂NH₂. [211°].

Di-bromo-nitro-benzene snlphonic acid $C_aH_2Br_a(NO_2)(SO_3H)$ [1:4:37:5]. From (1,4,5)di-bromo-benzene gulphonic acid and HNO, (Borns, A. 187, 358; Hübner a. Williams, A. 167, 121). Hygroscopic prisms which blacken at 100°.—BaA'₂ q.—BaA'₂ 1¹₂aq.—BaA'₂ 2¹₃q.—BaA'₂ 2¹₄q.—BaA'₂ 2¹₄q.—CaA'₂ 3¹₄q.—CaA'₂ aq.—CaA'₂ qq.—CaA'₂ qq. PbA', Baq. S. 10-3 at 10°

Chloride Call Bra(NO.) (SO.Cl). Oil (?). A mide C, H, Br, (NO,) (SO, NII,). [178°].

Di-bromo-nitro-benzene sulphonic acid C.H.Br.(NOz)(SO,II) [1:3:4:5]. From s-dibromo-benzene sulphonic acid and HNO, (Lenz, A. 181, 32). Tablets (containing xaq); not hygroscopic. By exchanging NO₂ for Br it is converted into (1,3,4,5)-tri-bromo-benzene sulphonic acid. BaA', 1¹,aq. S. 73 at 20³.— BaA', 4aq.—CaA', 3aq.—KA'aq. S. 1·09 at 20°.— NH,A',—PbA', 5aq. S. 1·20 at 20°. Chloride C,11,Br,(NO),(SO,Cl). [121°].

Amids C₄H₂Br₂(NO₂)(SO₂NH₂). Blackens

at 300%

Di-bromo-nitro-benzene sulphonio acid $C_4H_2Br_2(NO_2)SO_4H[1:3:4:6]$ [above 200°]. Formed by nitrating C.H.Br.SO.H [1:3:4] (Bässmann, A. 191, 235). Deliquescent needles or prisms (containing raq).—KA'. S. 1.4 at 21.5°.—BaA', 2aq. S. (of BaA',) 1.06 at 24°.—CaA', 6aq.—PbA', 4aq. Chloride C.H. (NO.)Br. SO.Cl. [115.5°]. Amide C. 11. (NO.)Br. SO.NH., Minute

tablets. Not melted at 240°.

Tri-bromo-nitro-benzene sulphonic acid A. 191, 196; Reinke, A. 186, 282; Bässmann, A. 191, 216). Hygroscopic, monoclinic prisms. Conc. HCl at 180° gives Il SO, and Conc. HCl at 180° gives H.SO and C.H.Br. (NO.) [125°]. - KA'. S. 76 at 5° (B.) CaA'₂2aq.—PbA'₂9aq. S. (of BaA'₂) ²07 at 1.5° (B.); 331 at 15° (L.). BaA'₂1¹₂aq.—CaA'₂2aq.—PbA'₂9aq. S. (of PbA'₂) 63 at 7° (B.); 93 at 10° (L.).—PbA'₁1¹₂aq.—PbA'₂PbO 7aq. PbA'.PbO 6aq. NH,A'.

Chloride C,HBr,(NO.)90.Cl. [145°].

 $Amide C_6HBr_3(NO_2)SO_2NH_2$

Tri-bromo-nitro-berzeno sulphonic acid CaHBra(NOa)(SOaH) [1:2:3:4:5]. From (1,2,3,5)tri-bromo-benzene sulphonic acid by nitration (Limpricht a. Lenz, B. 8, 1072, 1432; A. 181, 41). Lamine.—BaA', 4aq, S. 074 at 18'.—CaA', 3aq. S. 105 at 20'.—KA' aq. S. 16 at 18°. NH₄A' aq. PbA', aq. S. 14 at 20°. Chloride C₆HBr₃(NO₂)(SO₂Cl). [116°].

Amide C₆HBr₃(NO₂)(SO₂NH₂). [202°].

Tri-bromo-nitro-benzene sulphonic acid C, HBr, (NO,)(SO,H) [1:3:4:2:6]. [125°] or, anhydrous, [111]. From (1,2,4,5) tri-bromo-benz-ene sulphonic acid and HNO₃ (Spiegelberg, A. 197, 284) columns (containing 3aq).—Ag&'aq. S. (of AgA') '45 at 7°. BaA', 3aq. S. (of BaA'), '669 at 9°.—CaA', 4\{\}aq. S. (of CaA', 195 at 8°. KA', S. 19 at 8°.—NH, A', S. 168 at 6.5°. -PbA'₂ 6aq. S. (of PbA'₂) 853 at 7°. Uhloride C₆HBr₈(NO₂)(SO₂Cl). [143°].

Amide CallBrg(NOz)(SOzNIIz). Blackens

Tri-bromo-di-nitro-bonzene sulphonic acid C_aBr_s(NO_a)_aSO_sH [1:3:5:2:4:6]. [216°]. From Call Br SO II and conc. HNO at 100° (Bassmann, A. 191, 239). Colourless columns (containing 3aq). Not hygroscopic, but v. sol. water, sol. alcohol. - With water at 230° it gives C.HBr. (NO.). and H.SO.. Reduced by Sn and HCl to C.H.Br (NH.). SO. H.—N.11. A'aq.—KA'aq. S. (of KA') 48 at 24 . BaA', 9aq. S. (of BaA',) 83 at 21°.—CnA', 7, aq. IbA', 9aq. S. (of PbA',) 1.02 at 19.5°.

Chloride C.Br. (NO.) SO.Cl. [203°]. Amide C. Br. (NO2) SONH2. [200°].

Totra-bromo-nitro-benzene snlphonic acid CaBra(NO2)SO4H [1:2:3:5:4:6]. Got by nitrating C.HBr.SO.H. Crusts of needles (containing 4aq). V. sol. alcohol and water (Beckurts, A. 181, 220; Laugfurth, A. 191, 202). With conc. HCl at 200 it gives C. HBr. (NO.) and IL SO. - KA'1 aq. S. (of KA') '57 at 10.5°, - BaA', 9aq. S. (of BaA') '36 at 11° (B.); '100 at 14.5° (L.), -NH, A' aq. S. (of NH, A') 1.01 at 11°, --CaA', 8aq. S. (of CaA'2) 16 at 6°.—PbA'2 9aq. S. (of PbA'2) 06 at 6°

Chloride C.Br. (NO.) SO. Cl: [147.5°]; tablets. Amide: crystalline powder.

Tetra-bromo-nitro-bonseno sulphonic acid C.Br.(NO2)SO4H [1:2:3:4:5:6]. [173°]. From ctetra-bromo-benzene sulphonio acid and HNO, (Spiegelberg, A. 197, 297). Slender needlos (con-(Aphegorous) - BaA'₂4aq. S. (of BaA'₂)·22 at 12°.— BaA'₂9aq.—CaA'₂aq. S. (of CaA'₂)·28 at 13°.— KA'aq. S. (of KA')·17 at 11°.—NH₄A'. S. 46 at 11°.—PbA'₂2aq. S. 012 at 11°.

Chloride CaBr, (NO2)(SO2Cl): [173°]; prisms. Amide C. Br. (NO.) (SO. NII.). Blackens at

260°

BROMO-NITRO-BENZOIC ACID

[111°]. Formed, $C_aH_aBr(NO_a)CO_2H$ [1:4:3]. together with the isomeride [250°] by nitrating m-bromo-benzoic acid (Hübner a. Ohly, Z. [2] Monoclinic prisms.—NaA' 3aq.—NaA' 2½aq.— KA' 2aq.—BaA'₄4aq.—CaA'₂ 2aq.—MgA'₂ 4aq.— PbA'₄—CuA'₂. AgA.

Ethyt ether EtA': [55°]; monoclinic prisms. Bromo-nitro-bonzoio acid C₀H₂Br(NO₂)CO₂H [1:3:5]. [161°]. S. 057 at 11°. Forused from C₂H₂(NH₂)(NO₂)CO₂H, glacial acetio acid, Hisr (S.G. 1-19), and nitrous acid gas (Hesemann a. Köhler, A. 222, 166). Long needles (from water, benzene, ether, or CS₂), whetstone shaped crystals (from glacial acetic acid) or thin sixsided plates (from alcohol).— KA' laq.—BaA', 5 laq.—CaA', aq.—MgA', aq.—ZnA', 1 laq.—CdA', 4 laq.—SrA', —AgA', —PbA',

Bromo-nitro-benzoic acid CallaBr(NO.)CO.H [1:3:6]. [164°]. Formed by oxidation of C.H.Br(NO.)Mo by dilute HNO. (Schenfelen, A. 231, 173). V. sol. ether and dilute alcohol. Sl. sol. water. Reduced by Sn and HCl to m-bromo-

aniline, CO, going off. AgA'.

Bromo-nitro-benzoic acid CaHaBr(NOa)(COaH) [1:4:6]. [180°]. From C.H.MeBr(NO.) [1:2:5] and dilute HNO3 (Scheufelen, A. 231, 181) or by nitrating o-bromo-benzoic acid (Burghard, B. 8, 560). Almost insol. cold water, sl. sol. hot water, v. sol. ether and dilute alcohol. Alcoholic NII, at 130° gives p-nitro-aniline and (1, 4, 6)-amidonitro-benzoic acid.—BaA', 5 ! aq. Ethyt ether EtA': [66°]; needles.

Bromo-nitro-benzoic acid C,H3Br(NO2)CO2H [1:2:4], [199°]. Formed by nitration of p-bronno-benzoic acid [248] (Hübner, A. 113, 248; Raveill, A. 222, 177) and by oxidation of the corresponding bronno-nitrotolucne (Schenfelen, A. 231, 183). Long needles (from water) or plates (from dilute alcohol); v.

sol. ether, sl. sol. water. Reduction gives mamido-henzoic acid.

Salts. -AgA'. -BaA', 1aq. -MgA', 6aq. Ethyl ether EtA': [74°]; prisms.

Bromo-nitro-bonzoic acid C₆H₃Br(NO.)CO₂H [1:2:3]. [250°]. From mbrome-henzoic acid by nitration. Separated from its isomeride [141°] by being less soluble in water (Hübner, A. 113, 231; A. 222, 101). Monoelinio octahedra (irom cilier). NaA'aq.—, BaA', 4aq.—MgA', 6aq.

Ethyl ether Eth'. [80°]; prisms.

Di-bromo-nitro-bonzoic acid $C_{\rm e}H_{
m 2}Br_{
m c}({
m NO}_{
m c}){
m CO}_{
m c}H$ [3:4:2or6:1]. [162°]. From di-brome-benzoic acid [239°] by nitration (E. F. Smith, A. 222, 188). Colourless needles; reduction gives anthranilic acid.

Salts.—PbA',.—NaA' 3aq.—KA'.—BaA', aq. —CaA', 3laq.—MgA',

Di-bromo-nitro-benzole acld

C_sH₂Br₂(NO₂)(CO₂H). [162°]. Formed by nitrating the di-bromo-benzoic acid [223°-227°] obtained by brominating benzoic acid (Angerstein, A. 158, 13). Needles (from water). Reduction gives di-bromo-amido-benzoic acid [1963] and then authranilic acid. NaA' 3aq. -BaA' 2aq. This acid is perhaps identical with the preceding.

Di-bromo-nitro-benzoic acid

 $C_d H_2 Br_d (NO_2) CO_2 H_2 [3:5:2:1], [233^{\circ}].$ Formed by mitration of CaHaBr.CO.H (Hesemann a. Köhler, A. 222, 173). Long colourless needles; may be sublimed.— $\operatorname{BaA'}_2\operatorname{Iaq}$.— $\operatorname{CaA'}_2$.— $\operatorname{AgA'}$.— $\operatorname{KA'}$.

BROMO-NITRO-O-BENZYL-PHENOL $C_{13}H_{16}BrNO_{26}$ [105°-110°]. From potassium nitro-o-benzyl-phenol sulphonic acid and Br (Rennic, C. J. 49, 410). Yellow scales (from

alcohol). ⊸KA'.

Bromo-nitro-p-benzyl-phonol Ph.CH_aC_aH_aBr(NO_a)OH [1:3:5:1]. [65°].

Formation .- 1. From potassic bromo-benzylphenol sulphonate and dilute HNO₃ (1:1) (Rennie, C. J. 41, 223).—2. From potassic nitro-benzyl-phenol sulphonate, C₂H₄O₂, and Br.— 3. From nitro-benzyl-phenol, C.H.O., and Br. -4. From benzyl-phenol by first brominating and then nitrating.

Properties. - Crystalline scales (from alcohol). KA'. Red scales. HNO, oxidises it to bromodi-mitro phenol, $C_6H_2Br(N\ddot{O}_2)_2OH[1;2;3;5][118^\circ]$.

BROMO-NITRO-BUTANE C.H. BrNO. i.e. C,H,CHBr(NO₂). (181° cor.). From nitro-butane, potash, and Br (Züblin, B. 10, 2085). The three following compounds are prepared in a similar way (Z.).

Di-bromo-nitro-butane CaH, CBr, (NO.). (2010

Bromo-di-nitro-butane C,H,CBr(NO2)2. Not volatile.

Bromo-di-nitro-iso-butane

(CH₃)₂CH.CBr(NO₂)₂. [38]. Solid resembling camphor. V datile with steam. Readily decomposed by alkalis forming dinitro isobutane. BROMO-m-NITRO-CINNAMIC ACID

[3:1] C₆H₄(NO₂).C₂HBr.CO₂H. [212]. Formed by heating the dibromide of m-nitro-benzylidenemalonic acid (Stnart, C. J. 49, 361).

Bromo-p-nitro-cinnamic acid [1:1]C_aH₄(NO₂).C₂HBr.CO₂H. [146°]. V.sol. alcohol, other, chloroform. Sl. sol. hot CS. More sol. in cold water than its isomeride [205].

Salt, -BaA'. Boiled with water gives nitro-

phenyl-acetylene, CO, and Bal r... Ethyl ether Eth. (63). Prisms. From C.H. (NO.) CHBr. CHBr. CO. Et and alcoholic KOH (C. L. Müller, A. 212, 131).

Bromo-p-nitro-cinnam, acid [4:1] C_oH₁(NO₂) C HBr.CO.H. [205°]. Slender silky needles (from water). Sl. sol. cold water, insel, cold CS. V. sol. alcohol, other, chloro-

form, or benzoline.
Salt.-BaA'2. Decomposed by boiling into

nitro-phonyl-acetylene, CO, and BaBr, Ethyl ether EtA'. [93°]. Needles. From di-exo-bromo-p-nitro-phenyl-propionic ether by alcoholic KOH (C. L. Müller, A. 212, 131).

Di-bromo-p-nitro cinnamic acid

[4:1] CaH, (NO.). CBr: CBr. CO2H. [c. 180°]. From p-nitro-phenyl-propiolic acid and Br (Drewson, A. 212, 157).

Ethyl other Etk'. [86°]. V. sol. bonzene, chloroform or glacial HOAo, sl. sol. benzoline. BROMO-NITRO-CINNAMIC ALDEHYDE

C,H,(NO2).CH:CBr.CHO. [97°]. Long yellowish ncedles. Formed together with the isomeride [136°] by nitration of a-bromo-cinnamic aldehyde.

Phenyl-hydrazide. [131]. Large yellow

plates (Zineke a. Hagen, B. 17, 1816).

Bromo-nitro-cinnamic aldehyde C,II,(NO2).CII:CBr.Cl10. [436°]. Yellowish needles. Formed as above.

Phonyl-hydrazide. [151°]; red crystalline solid, sl. sol. alcohol (Z. a. II.).

DI-BROMO-NITRO-o-CRESOL

 $\{1:I:x:2\}$ C₆H(CH₃)(NO₂)(Br)₂(OH) Formed by bromination of nitro o- ercsol C.H4(C113)(NO2)(O11) [1:4:2]. Yellowish needles. V. sol, alcohol and other, nearly in ol. water (Nölting a. Collin, B. 17, 270).

Di-bromo-nitro-p-cresol

 $C_0 \text{IIMe}(\text{NO}_2)(\text{OH}) \text{Br}_2 [1:2:4:7:7]. [83°].$ nqueous nitro-cresol, [78°], and bromine-water. Long yellow needles (from alcohol). Insol. cold water, v. sl. sol. hot water, v. sol. alcohol or ether (E. Knecht, A. 215, 89; B. 15, 1071).

 $\begin{array}{c} Salts. \leftarrow C_0 IIMe(NO_2)(ONa)Br_2 2!aq. \quad Red\\ needles (from alcohol). - C_0 IIMe(NO_2)(OK)Br_2 aq. \end{array}$

DI-BROMO-NITRO-CUMENE C. II., CHBr. CBr(NO.).CII. Di-lophenyl - propylene. [77°-78.5°]. Di-bromo-nilrophenyt - propylene. [77°-78'5°]. From I'h.CH:C(NO.).CH, and Br (Priebs, A. 225, 362). ('olourless prisms (from light petroleum). Not decomposed even by hot aqueous NaOH, thus differing markedly from the corresponding dibromo-nitro-ethyl-benzene.

Bromo-nitro-4-cumene

C₀llMe₃Br(NO₂) [1:2:4:5:x]. [192°]. Formed by nitration of bromo-pseudo-cumence [73°] by fuming HNO, Needles, Sol. benzene, sl. sol. alcohol (Kelbe a. Pathe, B. 19, 1548).

Bromo-di-nitro-\psi-cumene C_sMe₃Br(NO₃)₂ [1:2:4:3:5:6]. [181°]. Formed by nitration of bromo-pseudo-cumene [1:2:4:3']. Long yellowish needles. Sl. sol. hot alcohol, nearly insol. cold (Kelbe a. Pathe, B. 19, 1551).

Bromo-di-nitro-\psi-cumene

 $C_aMe_aBr(NO_a)_2$ [1:2:4:5:3:6]. [214°]. Formed by nitration of brome-pseudo-cumene [73°] with fuming HNO3 and conc. H2SO1. Microscopio tables. Sol. benzenc, sl. sol. hot alcohol, nearly irrol. cold alcohol (Fittig, Λ . 147, 14; Kelbo a. Pathe, B. 19, 1518).

BROMO NITRO iso CUMENOL *C. H1.BrNO. i.e. C.H2.(C.H.)BrNO.)(OH) [1:5:3:2]. Bromo uitro-isopropyl-phenel. [33°]. From bromo-iso propyl-phenol and 11NO, (Fileti, G. 16, 123). Palo yellow needles (from diluto HOAc).

Bromo-nitro-iso-cumenoI

C₀H₂PrBr(NO₂)(OH)[1:3:5:2]. [88°]. From nitroisopropyl phenol and Br (F.). Nacreous tables

(from dilute alcohol); volatile with steam.

BROMO-NITRO-CUMYL-PROPIONIC ACID

C₁₂H₁₄BrNO₄ i.e. C₄H₃(C₄H₂)(NO₂).CHBr.CH₂.CO₂H. [127°]. From o nitro-eso-propyl-cinnamic acid and HBr (Einhorn a. Hess, B. 17, 2020).

Di-bromo-nitro-cumyl-propionio acid C₃II₃(C₃II₇)(NO₂).CIIBr.CIHBr.CO₂H. [171°]. From o-nitro-eso-propyl-cimamio acid and Br (Widman, B. 19, 260).

Di-hromo-nitro-cumyl-propionio acid. [184°]. From m-nitro-eso-propyl-cinnamic acid and Br (Widman, B. 19, 418)

BROMO-NITRO-CYMENE C₁₀H₁₂BrNO₂ i.e. 1..Me(C₂H₁)Br(NO₂) [1:4:3:x]. Formed by $C_0H_2Me(C_3H_7)Br(NO_2)$ [1:4:3:x]. nitrating the brome eymene derived from thymel (Mazzara, G. 16, 193). Oil, volatile with steam.

Bromo-di-nitro-cymene

C₀11 MePrBr(NO₂)₂ [1:4:2?:?:?]. [98°]. Formed by nitrating bromo-cymene (229°). Monoclinio prisms (Gerichten, B. 11, 1092). May be identical with the following.

Bromo-di-nitro-cymene

C_oHMePrBr(NO₂)₂ [1:d:3:?:?). [94°]. Formed by nitrating the bromo-cymene derived from thymol (M.). Slender yellow needlos.

Bromo-nitro-isocymene

 $C_{a}H_{a}(C_{a}H_{a})(CH_{a})(Br)(NO_{a}) = [4:2:1:?].$ Long red needles. Prepared by nitration of (1:2:1)-bromo-isocymene (Kelbe, B. 15, 40).

Bromo-nitro-m-isocymene (?) C10 II 12 (NO2) Br. [83°]. From di-bromo-m-isocymene by nitration (Kelbe a. Czarnomski, A. 235, 284).

Bromo-di-nitro-isocymene Call(Call)(CII3)(NO.) (Br). [55°]. Short thick needles. Prepared by nitration of (β) -bromo-isocymene (Kelbe, B. 15, 42).

BROMO NITRO ETHANE C.H.BrNO. i.e. CH3.CHBr(NO2). (147°). Formed by dissolving nitro-ethane (q.v.) in aqueous caustic potash and adding bromine: Cl1, Cl1 k(NO.) + Br₂ = CH, CHBr(NO.) + KBr (Meyer a. Wurster, B. 6, 91; Tseherniak, B. 7, 916; A. 180, 126). - Pungent oil. Forms unstable salts.

Bromo-di-nitro-ethane CH, CBr(NO.).. From Br and potassium dinitrocthane (Ter Meer, A. 131, 15).-Oil, volatile with steam; decomposed by K₂CO₃ which forms CH₃.CK(NO₂)₂.

Di-bromo-tetra-nitro-etbane CBr(NO₂)₂, CBr(NO₂)₂. From ethylene bromide and fuming 1INO₃, or from C₂(NO₂)₄K₂ and Br. Unstable liquid; forms with potash a compound C.Br. (NO.) 2KOH, m. sol. hot water, which explodes at about 180°. Ammonium sulphide converts it into C₂K₂(NO₂)₄. SO₂ forms NH₃, IIBr, and IICN. Aqueons K₂SO₃ forms yellow crystals C₂(NO₂)₄K₂3K₂SO₄ (Villiers, C. R. 94, 1122; 98, 431).

Di - bromo - nitro-ethane CH3.CBr2(NO2). (165°). Formed by adding potash to a mixture of nitro-ethane (q. v.) and the calculated quantity of bromine (V. Meyer, B. 7, 1313). Indifferent oil, insol. KHO.

BROMO-NITRO-ETHENYL - NAPHTHYL-ENE-DIAMINE

 $C_{10}H_4Br(NO_2) < N_H > C.CH_3 [4:x:\frac{2}{1}].$ [242°].

Formed by nitration of ethenyl-(4:2:1)-bromonaphthylene-diamino (Prager, B. 18, 2162). Yellow needles sl. sol. alcohol, v. sol. 11NO, Ac

ωα-DI-BROMO O-NITRO - ETHYL - BENZENE C₈H₁Br₂NO₂ i.e. [2:1] C₆H₄(NO₂).CHBr.CH₂Br. o-nitro-styrene dibromide. [52°]. From o-nitro-styrene and Br (Einhorn, B. 16, 2213).

ωα-Di-bromo-m-nitro ethyl-benzene [3:1] C_nH₁(NO₂).CHBr.CH₂Br. [79°]. From m-nitro-styrene and Br (Prausnitz, B. 17, 598).

ωα-Di-bromo-p-nitro-cthyl-benzene [4:1] C,11,(NO,),CHBr,CH,Br. [73°]. From pnitro-styreno and Br (Basler, B. 16, 3006).

ωa-Di-bromo-ω-nitro-ethyl-benzene C.H., CHBr. CHBr. O. [86°]. From ω -nitro-phenyl-ethylene and Br (Erdmann, B. 17, 414). Also from w-nitro-ethyl-benzene and Br (Priebs, A. 225, 341). Monoclinic crystals, a:b:c=1.257:1:1.396; L=83.51'. Cold aqueous NaOII givos bromo-nitro-styrone.

ωa-Di-bromo-ωo-di-nitro-ethyl-benzene [2:1] C_sH_s(NO_s).CHBr.CHBrNO_s. [91]. From wo-di-nitro-phenyl-ethylene and Br (Priebs, A. 225, 352). White needles, v. sl. sol. ligroin.

ωα. Di-bromo-ωp-di-nitro-ethyl-beazene [4:1] C.H. (NO.). CHBr. CHBr. NO. [1033]. From wp-di-nitro-phenyl-ethylene and Br (P.). Plates.

DI-BROM 9-NITRO-ETHYLENE C2HBr2(NO.). [1120]. From sodium tri-nitro-rosorein C2H(NO.)2(OH)(ONa) in aqueous solution, and bromino vapour (Merz a. Zetter, B. 12, 2046). Prisms (from CHCl3); does not combine with Br. Zn and HCl give ethylamine.

DI-BROMO-DI-NITRO-FLUORE3CEÏN C20HaBr2(NO2)2O5. From di-bromo-fluorescein and HNO, or from di-nitro-fluorescein and Br. Yellow needles; is not fluorescent.

Acetyl derivative [256'] (Baeyer, A. 183, 61).

BROMO-NITROFORM v. BROMO-TRI-NITRO-METHANE.

BROMO-NITRO-HYDROCINNAMIC ACID v. BROMO-NITRO-PHENYL-PROPIONIC ACID.

BROMO NITRO MESITYLENE C.H., BrNO. i.e. C. HBrMe, (NO2). [51]. Formed by mitrating bromo-mes tylene (Fittig a. Storer, 1.147, 7).

Brome di-nitro-mesitylene C.B. Mag(NO.) .. [194°]. From di-bromo-mesitylene by finning Needles (Süssenguth, A. 215, 248).

BROMO-NITRO-METHANE CIL Br(NO.). (141'). Formed by the action of bromine on sodium nitro-methane (Tscherniak, B. 7, 916; A. 180, 128 v. Nitro-Methane). Pungent oil. The bromine and nitroxyl render its hydrogen displaceable by sodium: it is a strong acid.

Bromo-di-nitro-methane CfiBr(NO.). From di-bromo-di-nitro-methane and alcoholic KOII (Villiers, Bl. [2] 37, 452; Losanitsch, B. 16, 51); or from (α)-di-bromo camphor and conc. HNO, (Kachler a. Spitzer, M. 4, 558). Oil.

Salt. - CKBr(NO.) .: S.G. 11 1.25; triclinic crystals which explode at about 147. tion by sodium-amalgam gives HCN, HBr, and NH, (Villiers, Bt. (2) 11, 282). Ammonium sulphido gives di-nitro-methane.

Bromo-tri-nitro-methano CBr(NO.), Bromo-[about 12°]. S.G. 2.8. From nitroform. nitroform and Br in sunlight; or from mercuric nitroform and Br (Schischkoff, A. 119, 247). Decomposes at 140°, but volatile with steam.

Di-bromo-nitro-methane Clibr (NO). (155°-160°). Formed by adding bromine to potassic bromo-nitro-methane:

 $CHKBr(NO_s) + Br_s = CHBr_0(NO_s) + KBr_s$ Bromopicrin, insoluble in potash, is formed at the same time (Tschorniak, A. 180, 130). Very pungent oil, volatile with steam, soluble in caustic soda.

Di-bromo-di-nitro-methane CBr. (NO.) [c. 0°]. Formed by the action of cone. IINO, on tri-bromo-aniline, ethylene bromide, bromophenol, or di-bromo-p-toinidine (Losanitsch, B. 15, 472; Villiers, Bl. [2] 37, 452). Greenishyellow, pungent oil, volatile with steam. Alkalis form salts of bromo-di-nitro-methane.

Tri-bromo-nitro-methane CBr₂(NO₂). Bromopicrin [10°]. S.G. 13:2.811.

Formation .- From nitro-methane, bromine, and KOH (V. Meyer a. Tsoherniak, A. 180, 122).

Preparation. - CaO (4 pts.), H2O (50 pts.), Br, (6 pts.) and pierie acid (1 pt.) are mixed in the order named and the product is distilled (Stenhouse, P. M. [4] 8, 36; Groves a. Bolas, A. 155, 253; C. J. 23, 153).

Properties .- Pungent prisms, may be distilled in vacuo. Converted by Br into CBr.

BROMO-DI-NITRO-METHYL-ANILINE C.H.Br(NO.)2NHMe [1:3:5:6]. [147°]. From di-nitro-methyl-aniline and Br. Yellow crystals; boiling aqueous KOH gives bromo-di-nitro-phenol (Norton a. Allen, B. 18, 1996).

Bromomitro-di-mothyl-aniline $\mathbf{C}_{a}\mathbf{H}_{3}\mathrm{Br(NO_{2})NMe_{2}}$ [4:3:1]. [72]. Long crystals.

Formed together with other products by the action of nitrons acid upon p-hromo-di-methylaniline (Koch, B. 20, 2460).

AMINE C₁₃H₁₀NBr(NO₂)₂, [1942]. Light yellow

Formed by bromination of di nitro.

BROMO-DI-NITRO-METHYL-DI-PHENYL-

tables.

methyl-di-phenyl-amine (Leymann, B. 15, 1236).

gives C_mH_aBr_z [81°] (Jolin, Bl. [2] 28, 515).

BROMO-NITRO-NAPHTHALENE $C_{ie}H_{i}Br(NO_{e})$ [1:4]. [85°]. From (a) bromonaphthalone and HNO_{a} . Yellow needles. PBr,

Bromo-nitro-naphthaleno $C_{16}H_1Br(NO_2)$ [1:1' or 4']. [122:5']. S. (93 p.o. alcohol) 337 at 15:7°. From (a)-nitro-naphthalene and bromine (Guareschi, A. 222, 291). Yellow needles (from alcohol). KMnO, gives bromo-phthalic acid [1749-176 "

Bromo-nitro-naphthalene Coll. Br(NO.) [3:1]. [131]. From (2,4,1)-bromo-nitro-(a)-naphthylamine by the diazo- reaction (biebermann a. Scheiding, A. 183, 262; Meldola, C. J. 47, 508). Straw-coloured needles. Exchange of NO. for Br gives (1,3)-di-bromo-naphthalene [61"].

Bromo-nitro-naphthaleno Cull Br(NO,) [1:3]. [132]. From (a)-naphthylamine by bromination, nitration, diazotisation &c. (Liebermann, B. 8, 1108; A. 183, 262). Yellow needles; Sn and HCl give (β) -naphthylamine.

Bromo-di-nitro-naphthalene

Coll Br(NO.). [170]. Long glistening needles. Formed together with the following isomeride by nitration of (a)-brome-naphthalene with furning HNO, (15). Not attacked by boiling with aqueous NaOH. On oxidation with dilute HNO, it gave a small quantity of (3)-nitro-phthalio neid (Merz a. Weith, B. 15, 2710).

Bromo-di-nitro-naphthalene C₁₀H Br(NO₂)₂ [143]. Tables or prisms. Formed as above. Notattacked by boiling aqueous NaOlf. On oxidation with ditute HNO, it gave a little (a)nitro-phthalic acid (Merz a. Weith, B. 15, 2710).

Bromo tetra-nitro naphthaleno $C_{10}H_3{\rm Br}({\rm NO}_2)_{\rm p}$ [190°]. Needles. S. (benzene at 18°) 3.7. Formed by further nitration of bromo-di-nitro-naphthalenc [170] by heating with a mixture of HNO, and H.SO,. It dissolves in caustic alkalis forming tetra-nitro-naphthol. NH, converts it into tetra-nitro-naphthylamine, and aniline gives the phonyl derivative of the latter. On oxidation with diluto HNO, it gives di-nitro-phthalic acid [227°] (Merz a. Weith, B. | Ba(C,H,Br,NO4), 2aq: red needles (Weselsky, 15, 2712).

Bromo-tetra nitro-naphthalene

C₁₆H₃Br(NO₂)₄. [245°]. White glistening needles. Nearly insol. ordinary solvents. Formed by nitration of bromo-di-nitro-naphthalene [143°] with a mixture of HNO, and H2SO,. It is attacked by alkalis with difficulty. NH3 converts it into tetra-nitro-naphthylamine and anilino gives the phenyl derivative of the latter. On oxidation with dilute HNO, it gives di-nitrophthalic acid [200] (Merz a. Weith, B. 15,

Di-bromo-nitro-naphthalene C₁₀11₃Br₂(NO₂). [96.5°-98°]. One of the products of action of Br on nitro-naplithaleuc. Small yellow needles (from alcohol) (Guareschi, A. 222, 286).

Di-bremo-nitro-naphthalene

 $C_{10}H_3Br_c(NO_a)$ [1:4:17]. [116:5°]. From (1,4)-di-bromo-naphthalene and $11NO_3$ (S.G. 1·4) in the cold (Jolin, BL, [2] 28, 515). PCl, gives tribromo-naphthalene [85°].

Di-breme-nitro-naphthalene C₁₀H.Br. (NO.). [100°-105°]. From di-bromo-naphthalene [68°] and HNO₃ (S.G. 1·4) (Canzoneri, G. 12, 427).

Tri-breme-di-nitro-naphthalene

C₁₀H₃Br₃(NO₂)₂. From (1, 2, 4)-tri-bromo-naphthaleno and fuming HNO₃ (Prager, B. 18, 2164). BROMO - NITRO - (α) - NAPHTHOIC ACID

C₁₀H₃Br(NO₂)CO₂H [1:4:4']. [260°]. Formed by nitration of bromo-(a)-naphthoic acid [246°]. Small yellowish prisms (from alcohol). Its ammonium salt forms glistening plates, el. eol. cold water (Ekstrand, B. 19, 1135).

BROMO-N1TRO-(a)-NAPHTHOL

 $C_{10}H_0Br(NO_0)(OH)[2:4:1]$. [136°]. From (2, 4, 1)bromo-nitro-acetyl-(a)-naphthylanine and conc. NaOH. Silky needles (from alcohol); oxidation gives phthalic acid.

Salts.— $C_{10}H_2Br(NO_2)(ONa)aq$: red needles. - $(C_{10}H_2Br(NO_2)O)_2Ba$ 3aq.

Methyl ether C,H Br(NO2)(OMe). [115°]. Pale yellow silky needles (Meldola, C. J. 47, 497).

Brome-nitre-(α)-naphthel C₁₀11, Br (NO₂)OH.

[142°]. From acetyl-bromo-(a)-naphthylamine by nitration and saponification (Biodermann a. Remmers, B. 7, 538).

BROMO-N1TRO-(a)-NAPHTHYLAMINE

C₁₀H₅Br(NO₂)(NH₂) [2:4:1]. [197°]. From the acetyl derivative by dissolving in conc. H₂SO₄ and ppg. with water. Orange needles; gives phthalic acid on oxidation (Meldola, C. J. 47, 497; 43, 9).

Acetyl derivative C10H Br(NO2)(NHAe). f225°). From acetyl-(a)-naplithylamine by nitration and bromination. Pale ochreous

needles.

Bromo-nitro-(a)-naphthylamine

 $\mathbf{C}_{10}\mathbf{H}_{5}\mathbf{Br}(NO_{2})(N\dot{\mathbf{H}}_{2})$ [4:2:1]. [200°]. From acetyl-(a)-bronio-(a)-naplithylamine by nitration and saponification (Liebermann a. Scheiding, A. 183, 258). Oxidised by dilute HNO, to plithalic acid. Elimination of NH, gives bromonitro-naphthalene [132°]. Conc. IIBrAq at 130° gives (1, 2, 4)-tri-bromo-naphthalene.

Acetyl derivative C10 H Br(NO2)(NHAc). $[232^{\circ}].$

DI-BROMO-NITRO-ORCIN C, H, Br, NO, i.e. C. MeBr. (NO.)(OH). [112°]. From (8)-nitro-orcin and Br. Yollow lamine (from alcohol).— B. 7, 444). BROMO-NITRO-O-OXY-BENZOIC ACID

C₆H₂Br(NO₂)(OH)CO₂H [5:3:2:1]. Bromo-nitro-salicylic acid. [175°]. Yellow noedles. Formed by nitration of bromo-salicylio acid in acetic: acid solution.—A'2Ca xaq: V. sol. water.—A'2Ba: yellow needles. $-C_6H_2Br(NO_2) < {}^{CO.O}_O > Ba 2aq$: red crystals.— $C_8H_2Br(NO_2) < {}^{CO,O}_{O} > Pb$: nearly

insoluble pp. (Lellmann a. Grothmann, B. 17,

Bromo-nitro-o-oxy-benzoic acid

C₆H₂Br(NO₂)(OH)CO₂H [3:5:2:1]. [222°]. Colourless needles. V. sol. alcohol, ether, and hot water. Formed by bromination of nitro salicylic acid in acctio acid solution.

Salts.—A'_Ba 4aq: long yellow needles.—A'_Ca 6aq: yellow prisms (Lellmann a. Groth-

mann, B. 17, 2724).

Bremo-nitre-o-exy-benzoic acid. Methul derivative C.H.Br(NO2)(OMe)(CO.H). From methyl bromo-isopropyl - phenyl oxide and HNO, (S.G. 1.3) (Peratoner, G. 16, 420). A dibromo-nitro-o-oxy-benzoic acid is also formed.

Bromo-nitre-p-oxy-benzeic acid. Methyl derivative Call_Br(NO2)(OMe)(CO.H) [1:3:2:5]. [182°]. Bromo-nitro-anisic acid. Formed by nitrating bromo-anisic acid.

Ethyt ether EtA'. [85°]: needles (Balbiano, G. 14, 241)

DI-BROMO-DI-NITRO-DI-OXY-D1-PHENYL SULPHONE

C₁₂H₂Br₂N₃SO₈ i.e. (C₄H₂(NO₂)(OH)Br).SO₂. [285°]. From di-nitro-di-oxy-di-phenyl sulphone in CS₂ and Br (Annalicin, B. 9, 660). Yellowish needles. — C₁₂II₄Na₂Br₂N₂SO₈ 2aq: orange needles.

BROMO-NITRO-OXY-PIPERIDINE-v-CAR-BOXYL1C ETHER C₅H₂(Br)(NO₂)(OH)N.CO₂Et [157°]. Colourless prisms; sol alcohol. Formed by the action of Br in HOAc on nitro-dehydropiperidine-v-earboxylic other (Schotten, B. 16, 646).

BROMO-NITRO-PHENANTHRENE v. PHEN-ANTHRUNE

BROMO-o-NITRO-PHENOL

C₄H₃Br(NO₂)(OH) [1:4:3]. [44°]. Formed, together with the following body [88°] by boiling (1, 3, 4)-bromo-di-nitro-benzene with aqueous KO11 (Laubenheimer, B. 11, 1160). Prisms; volatile with steam.—NaA': scarlet needlos.— BaA'2 aq: red neodles, sl. sol. water.—CaA'2 2aq. AgA'.

Bremo - o - nitro - phonol CaHaBr(NOa)(OH)

[1:3:4]. [88°].

Formation .- 1. From p-bromo-phenol and 11NO3 (Hübner a. Brenken, B. 6, 170; Körner, G. 4, 388) .- 2. From o-nitro-phenol (45g.) and Br (52g.) (Brunck, Z. 1867, 203).-3. From bromo-di-nitro-phenol (v. sup.).

Properties.—Yellow monoclinic laminm (from

alcohol) a:b:c = 2.941:1:1:625. $\beta = 64^{\circ} 2'$ (Arzruni, Z. Kryst. 1, 436); may be sublimed; v. sol. alcohol and ether, slightly volatile with steam. Reduced by Sn and HCl to bromo-amido-phonol

(Schütt, J. pr. [2] 32, 61).
Salts. -Na(C, H, BrNO,): red needles with golden-green lustre, v. sol. water.-KA 2aq.-

BaA', -AgA'.

Methyl ether CollisBr(NO2)OMe. [88°]. From the silver salt and MeI by boiling; or from the potassium salt, MeOH and MeI at 110°. V. sol. hot alcohol or hot ether, v. sl. sol. water (Staedel, A. 217, 55; B. 11, 1750).

Ethyl ether $C_{u}H_{3}Br(NO_{a})OEt$. [13°] (S.); [47°] (H.). From the potassium salt, Et1 and alcohol at 100°. Formed also by nitrating o-

bromo-phenetol.

Benzyl derivative

C₆H₃Br(NO₂)(OC₇H₂). [84°]. Yellow needles. Insol. water, v. sol. alcohol and glacial acetic acid, sl. sol. benzene, ether or chloroform (Roll a. Hölz, J. pr. [2] 32, 57). Reduced to bromoamido-phenol, when treated with Sn and HCl, benzyl chloride splitting off.

Bromo-m-nitro-phenol $C_oH_a(Br)(NO_o)(OH)$ [?:3:1], [147°] (L.); [110'] (P.). Prepared by bromination of m-nitro-phenol (Pfaff, B. 16, 612; Lindner, B. 18, 612). Yellow needles. Sublimable. Sl. sol. hot water, CS, and petroleumether, insol. cold water. On reduction with tin and HCl it gives m-amido-phenol, the Br atom being eliminated.-KA' 2aq: red crystals, sol. water and alcohol. - NaA'aq: yellowish - red erystals, sol. water and alcohol.—BaA', Jaq.

Methyl ether A'Me: [101°]; white needles, v. sol. alcohol and ether, on reduction [101°]; white with tin and HCl it gives meanisidine.

Ethyl ether EtA'. [57°]: prisms.

Bromo-p-nitro-phenol C, 11, Br(NO,)(OH) [1:3:6]. [102°]. Formed by brominating p-nitro-phenol (Brunck, Z. 1867, 201). Satiny needles (from ether or alcohol); m. sol. water... Ba(C,H3BrNO3)26aq: orange needles, in. sol. water.

Methyl ether MeA'. [106°]. From the potassium salt, McI and McOH at 110°. While needles (from alcohol). V. sol. hot alcohol or ether, m. sol. hot water (Staedel, A. 217, 66).

Ethyl ether EtA'. [98°] (S.); $[55^{\circ}]$ (H.). Formed like the preceding (S.). From p-nitrophenetol and Br (Halloch, B. 14, 37). Yellow needles (from alcohol). V. sol. alcohol or ether.

Benzylether C.H.Br(NO,)(OC,H.). [126°]. Nearly coloniess plates (from alcohol). Insol. water, sol. alcohol and other (R. a. H.). Reduced by Sn and HCl to brome-p-amide phenol and C,H,C1.

Bromo-di-nitro-phenol C₆H₂Br(NO₂)₂(OII) [1:3:5:4]. [86] (K.); [71] (Austen); [76] (Armstrong). Formed by nitrating p-bromophenol in HOAc, or by brominating and nitrating o-nitro-phenol (Körner, A. 137, 205; Armstrong a. Prevost, B. 7, 922). Formed also by brominating di nitro-phenol [61°] (Körner, G. 4, 305); and by boiling di-nitrated p-di-bromobenzene with aqueous KNO₂ (Austen, Am. S. [3] 16, 46). Yellow monoclime prisms, a:b:e-**2.795:1:1.778**; $\beta = 67^{\circ} 53'$ (Arzruni, loc. cit.). Water and Br at 100° change it into the isomeride [118°] (Armstrong, C. J. 28, 520). 11NO, forms pierio acid.

Salts.-NH,A': silky red needles; sol. boiling water and alcohol. - NH, A'aq. -BaA', : yellow needles, sol. hot water. - CuA'2: short brown needles, insol. water.—KA': long red needles, sl.

sol. water. -AgA'. - CuA'₂8uq.

Ethylether EtA' [66°]: small needles, sol. alcohol and hot water; saponified by cold [91°]. Formed by heating m-nitro-phenol (1 Vor. I.

NaOHAg (Schoonmaker a) Van Mater. Am. 8. 187).

Bromo-di-nitro-phenol CaH, Br(NO,) (OH). [91.5°]. Formed by nitrating m-bromo-phenol. and also from di-bromo-di-nitro-phenot [117°] and boiling aqueous KOH (Körner, G. 4, 305). Prisms (from alcohol or ether). The K salt forms yellow needles.

Methyl other MeA' [109].

Bromo - di - nitro - phenoI CallaBr(NOa)aOH [1:3:5:6]. [118°].

Formation.—1. From (1,3,4)-di-nitro phenol and Br (Laurent, Rev. Scient. 6, 65). -2. By nitrating o-bromo-phenol (Körner, 6, 1, 394).—3. From o-nitro-phenol by bromination and nitration.-4. By boiling bromo-di-nitra-aniline [144°] with aqueous KOH (Körner).-5. By nitrating brominated phenol disulphonic acid or di-brominated phenol p-sulphonic acid (Armstrong a. Brown, $C.\ J.\ 25,\ 861,\ 865, -6.$ By warming the isomeride [c. 76°] with Br and water (Armstrong, C. J. 28, 520). -7. From picric acid, water, and Br (Armstrong, B. 6, 650). 8. By nitrating tri-bromo-phenol (Armstrong a. Harrow, C. J. 29, 477) .- 9. From bromo nitrobenzyl-phenol in HOAc by HNO_a: benzyl being displaced by NO₂ (Rennie, C. J. 11, 225).

Properties.—Yellow prisms. Needles (from

Salts.—KA'aq: tlat yellow needles, sl. sol. cold water.—KA'1 aq. BaA₂'3 laq. BaA'. laq. -BaA', 5aq: yellow needles, sl. sol. water. --— CaA'₂ 7aq. — CaA'₂ 8aq. — CaA'₂ 12aq. — NH₁A' 2aq. — NaA' 1'aq. — PbA'₂ 2aq. Methylether MeA' [48°]. From bromo-

anisic acid and HNO₃. Yellow prisms, sol. alcohol and other, insol. water (Balbiano, G. 14, 235).

Di-bromo-o-nitro-phenoI Cull_Br.(NO2)(OII) [1:3:5:6]. [117:5°].

Formation .-- I. From o-nitro-phenol and Br (Brouck, Z. 1867, 203; Körner).—2. From (1,3,4,)-di-brom--phenol by nitration (K.).—3. By nitrating di-bromo-phenol sulphonic acid (Armstrong a. Brown, C. J. 25, 863).

Properties. - Golden monoclinic prisms (from alcohol); a:b:e = 515:1::591; $\beta = 65^{\circ}23'$ (Arzruni, Z. Kryst. 1, 436). Volatile with steam; may be sublimed; v. sl. sol. water. Heated with bromine at 100° it gives some of the isomeride [141°] together with t. tra-bromo-quinone and (1,3,6) bromo-nitro-phenol (Ling, C. J. 51, 147).

Salt .- KA': scarlet needles, v. sl. sol. cold water.

Methylether MeA'. [77°]. silver salt and Mel; alcohol'e NII, converts it into di-bromo-nitro-ar dine [127°] whence HNO gives di-bromo-nitro-benzene [105°] (K.).

Ethylether Eth. [46°]. From the silver salt, Etl and alcohol as 100' (Staedel, A. 217, 58). V. sol. benzene, alcohol or ether, insol. water.

Benzylether C, H2Br2(NO2)(OC,H,). [65°]. Yellow crystals (from ether). Sol. henzene, chloroform, and glacial acetic acid, insol. water (R. a. H., J. pr. [2] 32, 57). Reduced by Sn and HCl to di-bromo-amido-phenol and benzyl ehloride.

Di-bromo m-nitro-phenol CaH2Br2(NO2)OH.

mol.) with bromine (2 mols.). Yellowish plates. | p-bromo-benzoic acid and a little p-nitro-benzoic V. sol. alcohol, sl. sol. water.

Salts .- KA' aq: casily soluble orange-red needles .- AgA': sparingly soluble red powder.-BaA', 6aq : very soluble red needles.

Ethylether C.H.Br.(NO.)OEt: [110°]: yollowish needles, sol. hot alcohol (Lindner, B. 18, 613).

Di-bromo-p-nitro-phenol C₆11₂Br₂(NO₂)(O11) [1:5:3:6]. [112°] (Lellmann a. Grothmann, B. 17, 2731).

Formation .-- 1. By brominating p-nitro. phenol or its sulphonic acid (Brunck, Z. 1867, 201; Post a. Brackebusch, A. 205, 94). 2. By nitrating tri-bromo-phenol dissolved in 110Ac (Armstrong a. Harrow, loc. cit.) .- 3. In small quantity by heating the isomeride [117°] with Br (Ling, C. J. 51, 147).

Properties. - Prisms, sl. sol. water, v. sol.

alcohol and ether.

Salt. -BaA'. 10 aq : yellow needles, effervesoing to a red powder. BaA', Baa, AgA'.

Methyl ether McA': [123°] (Körner, G. 4, From di bromo-anisic acid and HNO, the CO.II being displaced by NO. (Balbiano, G. 14, 9). Pyramidal needles; converted by NII, into di-bromo-p-nitro-aniline [203°].

Ethyl ether [108°]. columns (Staedel, A. 217, 67). Long quadratie

Benzyl ether C. II. Br. (NO.) (OC. II.). [94°]. Nearly colourless needles (from alcohol). Insol. water, sl. sol. alcohol, ether, benzene, ehloroform, and glacial acetic acid (R. a. II.). Reduced by Sn and HCl to di-bromo-p-amido-phenol (q. v.) and benzyl chloride.

Tri-bromo-nitro-phenol CaHBra(NO.).OH [6:4:2:3:1]. [85°] (L.); [89°] (D.). Formed by heating m-nitro-phenol with Br (3 mols.) at 100° (Lindner, B. 18, 614). Colourless crystalline powder; v. sol. alcohol, other, and beazene, v. sl. sol. hot water.

Salts .- A'NH : sparingly soluble microscopic needles. - A'Kaq: v. sol.water. - 'A' Br Saq: sparingly soluble orange-yellow crystals,-A' Ba aq. -A' Mg*: easily soluble red plates.

o-Nitro-benzoyl derivative $C_0HBr_3(NO_2)O.CO.C_0H_1(NO_2)$: [129° cor.]; very small colourless needles.

m-Nitro-benzoyl derivative

 $C_6HBr_3(NO_2)O.CO.C_6H_4(NO_2)$: [154° cor.]; glis-- tening colourless needles; S. (90 p.c. alcohol) *253 at 14° (Daccomo, B. 18, 1167).

Ethyl ether EtA': [79°]; prisms.

BROMO-NITRO-PHENOL SULPHONIC ACID $C_aH_aBrNSO_ai.e.C_aH_aBr(NO_a)(OII)SO_aII[1:3:6:5].$ Formed, together with di-bromo-nitro-phonol, by brominating (1, 4, 3)-nitro-phenol sulphonic acid (Posta, Brackebusch, A, 205, 92).—CaA'₂3aq. -BaA', 3aq. -C, 11, BrNSO, (PbO11), 21, aq.

Bromo-phenol sulphonic acids have been obtained by Armstrong (C. J. 25, 857; B. 7, 401, 924) and Post (B. 7, 169) by the action of nitrie acid on various brominated phenol sulphonic acids, and by the action of bromine on (1, 2, 5) nitro-phenol sulphonic acid.

BROMO-NITRO-DIPHENYL C, II, Brno. i.c. [4:1] C₀H₄Br.C₀H₄(NO₂) [1:4]. [173°]. (abovo 360°). Formed by heating diphenyl (1 pt.) with conc. HNO, (1 pt.), or from amido-nitro-diplienyl by the diazo reaction (Schultz, A. 174, 218). Long white needles (from toluene). CrO, gives

acid.

Bromo-nitro-diphenyl C₁₂H₄BrNO₂ i.s. [4:1] C₆H₄Br.C₆H₄(NO₂) [1:2]. [65°]. (o. 360°). Formed together with the preceding. Monoclinio columns. columns. CrO₃ gives p-bromo-benzoio aoid (Schultz, A. 207, 348).

Di-bromo nitro diphenyl CaHaBr.CaHaBr(NO.). [127°]. Formed by nitration of di-bromo-diphenyl in acctic acid solution (Lellmann, B. 15, 2837). Yellowish crystals; v. sol. alcohol, benzene, and acctic acid.

Di-bromo-di-nitro-diphenyl C12H6Br2(NO2)2 From pp-di-bromo-diphenyl and fnming HNO, (Fittig, A. 132, 206; S.). Hair-like crystals (from benzone). Su and KCl give di-bromo-diamido-diphenyl [89°].

Di.bromo-tri-nitro-diphenyl

C₆H₃Br(NO₂).C₆H₂Br(NO₂). [177°]. Formod by nitration of di-bromo-diphenyl with cold furning HNO₂ (1.55) (Lellmann, B. 15, 2838). Small colourless needles. Sol. benzono, st. sol. alcohol.

BROMO-NITRO-PHENYL ACETIC ACID C.H.Br(NO.)CH2CO2H [1:2:4]. [114°]. Formed by nitration of a mixture of o- and p-bromophenyl-acetic acids (Bedson, C. J. 37, 97). Flat greenish-yellow needles. Sol. hot, insol. cold water. V. sol. alcohol and ether. K.Cr2O, and H_SO, give C_HBr(NO.)CO.H [199°].

Salts.—BaA', aq. Its aqueous solution gives white pps. with AgNO3, Pb(OAc)2, and a blue

pp. with Cu(OAc)...

Methyl ether. McA'. [41°]. Needlos. Ethyl ether. Oil.

(a).Bromo-nitro-phenyl-acetic acid C₆H₃Br(NO₂)CH₂CO₂H. [169°]. Formed by nitration of mixture of o- and p-bromo-phenylacetic acids (Bedson). Yellowish-white, branching needles. V. sol. alcohol and ether, insol. cold, sol. hot water.

Salt.—BaA', Juq. Its aqueous solution gives white pps. with AgNO₃ and Pb(OAc), but a green pp. with Cu(OAc)2.

Methyl ether [68°]. Flat needles. Ethyl ether. Yellowish needles.

(B)-Bromo-nitro-phenyl-acetic acid C, H, Br(NO.) CH., CO.H. [162°]. Formed togother with the two preceding (Bedson). Small yellow prisms.

BROMO-DI-NITRO-DI-PHENYL-AMINE C12HaBrN3O1i.e.C6H2Br(NO2)2.Nfl.C6H3. Phenylbromo-nitro-phenyl-amine. [120°]. Formed by warming bromo-di-nitro-benzene [100°] with anilino (Austen, B. 9, 920). Orange hair-like ncedles (from alcohol).

Bromo di nitro di phenyl amino C.H.Br.NH.C.H.(NO.J., Bromo-phenyl-di-nitro-phenyl-amine. [153°]. From (1, 2, 4)-bromo-

di nitro-benzene and di-p-bromo-di-phenyl-urea at 170°. Yellow needles (from ether) (Willgerodt, B. 11, 60%).

Bromo-tri-nitro-di-phonyl-amins Call, Br(NO.) .. NII.CaH, NO. [158°]. From bromodi-nitro-di-phenyl-amine [120°] and HNO, (A.).

Di-bromo-di-nitro-di-phenyl-amine C₁₂H₂Br₂(NO₂)₂N. [196°]. Formed by bromination of di-nitro-di-phonyl-amine (Leymann, B. 15, 1236).

Di-bromo-tetra-nitro-di-phenyl-amine C_sH₂Br(NO₂)₂·NH.C_sH₂Br(NO₂)₂· [2350-2427], From NMe(C,H2Br2)(C,H4Br) and HNO, (Gnehm, B. 8, 929). Lamine (from HOAc).

Tri-bromo-di-nitro-di-phenyl-amine O₁₂H₂Br₂(NO₂),N. [210°]. Formed by nitrating tetra - bromo - di - phenyl - amine (C_uH₂Br₂),NH (Gnehm a. Wyss, B. 10, 1323).

BROMO-NITRO-PHENYL BENZYL OXIDE

v. Bromo-nitro-phenol, Benzyl ether.

DI - BROMO - NITRO - PHENYL - CARBAMIC ACID. Methyl ether [I:3:4] C_eH₂Br₂(NO₂).NII.CO₂Me. [152°]. From methyl-(1,3,4)-di-bromo-phenyl-carbamate and HNO₃ (Hentschel, J. pr. [2] 34, 425). Prisms (from alcohol). NII, forms (1,3,1)-di-bromo-

aniline.

BROMO-NITRO-m-PHENYLENE-DIAMINE C_eH_eBrN_aO₂ i.e. C_eH₂Br(NO₂)(NH₂)₂ [1:2:3:5]. From tri-bromo-nitro-benzene and alcoholic NH₄ for some days at 175° (Körner, G. 4). Orange needles; decomposes at 163°. Converted into p-bromo-benzene by the diazo- reaction.

Bromo-nitro-p-phenylene-diamine

C₆H Br(NO₂)(N1Î₂)₂ [1:1:2:5]. [156°]. From tribromo nitro-benzene [94°] and alcoholic NII, at 110° (Körner, G. 4). Pyramidal needles. Converted by diazo- reaction into p-bromo-benzene.

BROMO - NITRO - PHENYL - ETHANE v.

Bromo-nitao ethyl benzene.

Di-bromo-di-nitro-s-di-phenyl-ethane

C.H.(NO.).CHBr.CHBr.C.Hr.NO. [above 300°]. Di-bromide of di-p-nitro-stilbene. Split up by heat into 2HBr and di-nitro-tolano (Elbs a. Bauer, J. pr. [2] 31, 315).

Di-bromo-di-nitro-s-di-phenyl-etbane

 $\mathbf{C_{i,H_{in}}}$ $\mathbf{Br_{o}}(\mathbf{NO_{o}})_{2}$. Di-bromo-di-nitro-di-benzyl, [205°]. Formed by nitrating di-p-bromo-di-benzyl (Fittig a. Stelling, A. 137, 260). Sword-Di bromo di nitro di benzyl. shaped crystals (from benzene).

BROMO-NITRO-PHENYL ETHYL OXIDE v.

Ethyl-Bromo-nitro-fuenol.

aa-DI - BROMO - am - DI-NITRO - ω - PHENYL-

METHYL-CARBINOL.

 $Ethyl\ ether\ CBr_2(NO_2).CH(C_aH_1NO_2).OEt.$ [99] .- 1. From the compound of alcohol with am-di-nitro-cinnamic ether (q, v_i) by simultaneous treatment with aqueons NaOH and Br (Friedländer a. Lazarus, A. 229, 237).—2. From ω -mdi-nitro-styrene, alcohol, aqueous NaOH and Br. White plates (from dilute alcohol). Insoluble in aqueous NaOll.

Methylether CBr.(NO.).CH(C.H.NO.)OMe. [146°]. Formed in a similar way from the compound of methyl alcohol with the same body, or from ω-m-di-nitro-styrene, methyl alcohol, Br.

l aqueous NaOll. White plates.

BROMO - NITRO - PHENYL - METHYLand aqueous NaOl1.

KETONE v. Bromo-nitro-acetophenone

p-BROMO-o-N1T30-B-PHENYL-PROPIONIC ACID C₆H₃(Br)(NO₂)C₆H₄CO₂H [4:2:1]. Bromonitro-hydrocinnamic acid. [112]. Prepared by the action of IIBr on the diazo- compound from p-amido-o-mitro-hydrocimamic acid. Formed together with the (4,3,1) isomeride by nitration of p-bromo-hydrocinnamic acid. Flat feathery crystals. By reduction with tin and HCl it gives p-bromo-hydrocarbostyril (Gabriel a. Zimmermann, B. 13, 1682).

p-Brono-m-nitro-6-phonyl-propionic acid

O_cH₂(Br)(NO₂).O₂H₁.CO₂II [4:3:1]. [90'-95'].

Long glistening needles. Prepared as above.

On reduction with tin and HCl it gives p-bromo-

m-amido-hydrocinnamio acid (Gabriel a. Zimmermann, B. 13, 1683).

β-Bromo-o-nitro-β-phenyl-proplonio acid CaH₁(NO₂).CHBr.CH₂.CO₂H. [140°]. From onitro-cinnamic acid, 110Ac, and 11Br at 100° (Einhorn, B. 16, 2208). Monoclinio crystals; v. sol. ordinary solvents, sl. sol. benzene.

Reactions. - I. Boiling water forms indoxyl. 2. NaOHAq forms nitro-cinnamic acid. -3. Cold Na.CO3Aq forms the lactone of σ-nitro-β-oxyphenyl-propionic acid.—1. Hot Na CO Aq gives nitro cinnamie acid, nitro-oxy-phenyl-propionio acid, and o-nitro-styrene.

B-Bromo-m-nitro-B-phenyl-propionic acid C.II. (NO.).CHBr.CH., CO.II. [96]. Prepared by heating an acetic acid solution of m-nitro-cinnamic acid with Illir at 100'. V. sol. alcohol. sl. sol. toluene, insol. petrolemn-ether.

Reactions. - 1. By boiling with water it chiefly gives m-nitro-styrene. 2. An excess of alkali converts it back into m-nitro ciunamie acid.-3. By adding the powdered acid to an aqueous solution of 1 mol. of Na.CO., it yields 30 p.c. of m-nitro-styrene, 20 p.c. of m-nitrocinnamic acid, and 10 p.c. of m-nitro-\$-oxy-\$phenyl-propionic acid. If the powdered acid is added to a cold solution of Na₂CO₃ the \$\beta\$-lactone is formed (Prausnitz, B. 17, 595).

B-Bromo-p-nitro-phenyl-propionic acid

C.H.(NO.).CHBr.CH.,CO.H. [172]. Prepared by heating p-nitro-cinnamic other with IIBr at 100'. Prisms; sol. hot alcohol, sl. sol. water and benzene.

Reactions .- I. Long boiling with dilute II,SO, (25 p.o.) reconverts it into p-nitro-sinnamic acid.—2. Heated with water it yields p-nitro-β-oxy-phenyl-propionic acid (72 p.c.) and p-nitro-styrene (28 p. c.) - 3. Cold aqueous KOII gives p-nitro oxy-phenyl-propionic neid and its lactone. Alcoholic KOII yields almost entirely p-nitro-cimeamic acid.—1. Aqueons NII, yields the lactors which by oxcess of NII3 is converted into the corresponding amido- acid.

Ethyl ether A'Et. [81°]; colourless plates

(Basler, B. 16, 3001).

Di-bromo-o-nitro phenyl-propionic acid CoH4(NO2)CHBr.CHBr.CO2H. Dibromide of onitro-cinnamic acid. [c. 180]. From Br and o-nitro-cinnamic acid (Baeyer, B. 13, 2257). Needles or plates; sol. he. water. NaOHAq gives o-nitro-phenyl-propiolic acid. Zinc-dust and NaOH give indol...

Methyl ether MeA'. [99']. Ethyl ether EtA'. [71° uncor.] (M.) From σ-nitro-cinnamic ether and Br (Müller, A. 212, 130). Alcoholic KOH converts it into onitro-phenyl-propiolis acid. Heated with water at 120 ' it gives o-nitro cinnamie acid.

Di-bromo-p-nitro-phenyl-propionic acid Call (NOa) CHBr. CHBr. CO. H. [218"]. From pistro-cinnamic acid and ir (Drewson, A. 212, 151). Rhombic prisms (from glacial HOAc). M. sol. water or glacial HOAc, v. sol. alcohol or other, sl. sol. benzene, v. sl. sol. benzoline. Aqueous NaOII forms p-nitro-cinnamio and p-

nitro-phenyl-propiolic acids.—CaA'₂.

Ethyl ether EtA'. [II1°]. From p-nitrocinnamic ether and Br in CS₂ (C. L. Müller, A. 212, 129). Columns (from CS₂); v. sol. hot alcohol, ether, or benzoline. Alcoholic KOH converts it into a mixture of two isomerio bromo-pnitro-cinnamic ethers, and p-nitro-phenyl-propiolic acid. Heated with water at 120° it gives p-nitro-cinnamic acid.

BROMO-NITRO-PHTHALIC ACID

C₆H₂Br(NO₂)(CO₂H)₂ [1:4:2:3]. From di-bromonaphthalene [820] and HNO3.—Na2A" (Guareschi, A. 222, 277).

· α-BROMO-ω-NITRO-PROPANE

CH₃,CH₂,CHBr(NO₂). (c. 160°). Formal together with di-bromo-nitro-propane by the action of potash and Br on nitro-propane (V. Meyer a. Tscherniak, A. 180, 116). Oil, sol. potash.

a-Broho-a-nitro-propane CH , CBr(NO.), Cll , (150°). From CH2, CH (NO2), CH3. Insol. potash. Di-bromo-nitro-propano CH. CH. CBr. (NO.). (184°-186°). From bromo-nitro-propane, Br, and

potash. An oil, insol. potash.

TRI-BROMO DI-NITRO-PROPIÒNIC ACID C₃HBr₃N₂O₆ i.e. CBr₃.C(NO₂)₂.CO₂H. From tribrome-phloroglucin and conc. HNO, (Benedikt, A. 184, 255). Silky scales, insol. cold water, v. e. sol. alcohol and other; decomposed by boiling water.

BROMO - NITRO - PROPYL - BENZENE

Bromo-nitro-cumene.

BROMO-NITRO-PROPYL-PHENOL v. BROMO-

DI-BROMO-DI-NITRO-PYRROL

CBr:C(NO2) C.HBr.2N,O. i.e. [169°]. CBr:C(NO₂)

Formed by dissolving di-bromo-nitro-pyrryl methyl ketono [206°] in a mixture of fuming HNO, and conc. H.SO, at -18', and precipitating in water. Silky teatlets (containing aq). At the ordinary temperature it is converted into dibromo-maleimide (Ciamician a. Silber, B. 20, 699; (J. 17, 262).

DI-BROMO-NITRO-PYRRYL METHYL KE-

CBr:C(NO₂) CBr.C-CO.CH, [206°], Long needles.

Formed by the action of cold nitric acid upon di-bromo-pyrrylene-di-methyl-di-ketone CBr:C-CO.CH,

NH CBr.C CO.CH₃, [171°], which is formed by passing bromine-vapour into pyrrylene di-methyl di-ketone (Ciamician a. Silber, B. 20, 699).

Di-bromo-nitro-pyrryl methyl ketone

CBr : CBr NH [175°]. From nitro-pyrryl C(NO.):C-COCH, methyl ketone [197°] and Br. Needles.

BROMO-NITRO QUI NOLINE C., H., N(Br)(NO2) [133°]. Formed by nitration of bromo quinoline. Long glistening needles. Volatile undecomposed. Sol. alcohol and other, sl. sol. water. Weak base. -B', H, Cl, PtCl, : short orange-yellow prisms (La Coste, B. 15, 1918). DI-BROMO-NITRO-QUINONE

 $C_6HBr_2(NO_2)O_2[6:2:3:4:1]$. [246°]. Yellow plates; v. sl. sol. hot water and cold alcohol. Formed by the action of a mixture of HNO₃ and H₂SO₄ upon the propionyl derivative of tri-bromo-

BROMO-NITRO-RESORCIN Ethyl ether C_s11₂Br(NO₂)(OH)(OEt). [114°]. From eth From ethyl (I, 2, 4)-nitro-resorcin and Br (Weselsky, M. 1. 898).

phenol (Guarcschi a. Daccomo, B. 18, 1174).

Di-bromo-nitro-resorcin CaHaBraNO C₆HBr₂(NO₂)(OH)₂. [147°]. From (1, 2, 4)-nitroresorcin [115°] in ether and Br (Weselsky, A. 164, 7). Golden laminæ.—Ba(C₈H₂Br₂NO₄)₂ 4aq.

Ethyl $C_0H\dot{B}r_2(NO_2)(OEt)(OH)$ ether [1:5:3:2:6]. [69°]. From ethyl nitro-resorein and

Di-bromo-nitro-resoroin C, IIBr (NO2)(OH)2 [1:3:5:4:6]. [117°]. From (1, 2, 6)-nitro-resorcin and Br (W.)

Bromo-di-nitro-resorcin C.HBr(NO2) (OH)2. [193°]. Formed by nitrating di-bromo-nitrosoresorcin or by brominating di-nitro-resorcin (Fèvre, Bl. [2] 39, 590; C. R. 96, 790; Typke, B. 16, 555). Orange needles; sl. sol. boiling alcohol. Ka' aq: red needles.—Na,A"2aq.— BaA" 3aq.—(NH),A"aq.
Acctyt derivative: [135]; prisms.

BROMO-NITRO-SALICYLIC ACID v. Bromo-NITRO-OXY-BENZOIC ACID.

DI-BROMO-N1TROSO-RESORCIN

C₀HBr₂(NO)(OH)₂. From nitroso-resorcin and Br (Fèvre, Bt. [2] 39, 591). Yellowish needles (containing 2aq) which turn brown at 138 and decompose at 150°; insol. benzene, sl. sol. cold water, v. e. sol. alcohol.

BROMO-NITROSO-THYMOL

C₆HMePrBr(NO)(OH). [c. 135°]. From nitrosothymene and Br (Mazzara a. Descalzo, G. 16, 196).

BROMO-NITRO-STYRENE Ph.CBr:CHNO2. From di-bromo-nitro-phenyl-ethane PhCBrH.CHBrNO2 and aqueous NaOH (Priebs, A. 225, 313). Golden needles or plates (from light petrolemm). Smells something like hay; when freshly ppd. from alcoholic solution by water it is solublo in alkalis, hence its constitution is as above, rather than Ph.CH:CBr(NO2). DI-BROMO-DI-NITRO-THIOPHENE

C₁SBr₂(NO₂)₂. [134°]. Light-yellow crystals. V. sol. hot alcohol. Formed by nitration of dior tri-bromo-thiophene (Kreis, B. 17, 2074; Losenberg, B. 18, 3029).

TRI-BROMO-NITRO-THIOPHENE

C₁SBr₂(NO₂). [106°]. Formed by nitration of tri-bromo-thiophene. Felted yellow needles. V. sol. ether, sl. sol. alcohol (Rosenberg, B. 18, 3028).

BROMO-NITRO-THYMOL

 $C_aHMePrBr(NO_a)(OH)$. [101°]. From bromonitroso-thymoland K_aFeCy_a (Mazzara, G. 16, 196).

BROMO-NITRO-TOLUENE

C_bll₃Me(NO₂)Br [1:2:3]. Oil. From bromo-nitrom-tolniding by nitrous gas and alcohol (Nevile Winther, C. J. 37, 630).

Bromo-nitro-toluene C₆H₃Me(NO₂)Br [1:3:4]. [32°], (256°). S.G. 12 I-631. From the corresponding nitro-toluidine by the diazo-perbromide reaction (Nevilo a. Winther, C. J. 37, 442). Formed also by nitrating p-bromo-toluene (Wroblewsky, A. 168, 176) and by treating mnitro-tolueno with Br and FeBr, at 70° (Scheufelen, A. 231, 180).

Bromo-nitro-toluene C₆H₃Me(NO₂)Br [1:2:4]. [45°]. (257°). From the corresponding nitro-toluidine [78°] by the diazo-perbromide reaction (Beilstein a. Kuhlberg, A. 158, 340; Nevile a. Winther, C. J. 37, 441). Formed also, together with the preceding, by nitrating p-bromo-toluene (W.). Large monoclinic tables.

C₆H₃Me(NO₆)Br Bromo - nitro - toluene [1:20r6:3]. [55°]. (267°). Formed by nitrating m-bromo-toluene (W.; Grete, A. 177, 246). Trimetric crystals; on reduction it gives bromoo toluidine.

Bromo-nitro-toluene C₄H₃Me(NO₃)Br [1:4:2]. [77°]. From C₆H₂(CH₃)Br(NO₂)(NH₃) [1:2:4:5] (Nevile a. Winther, C. J. 39, 85). From p-uitro-toluene, FeBr₂, and bromine (Scheufelen, A. 231, 171). Also from diazo-nitro-toluene piperidide C₂H₃Me(NO₂).N₂·NC₃H₁₀ and boiling HBrAq (Wallach, A. 235, 248). Needles.

Bromo-nitro-tolueno C₄11,Mc(NO₂)Br [1:3:6]. [78°]. From m-nitro-toluene, FeBr₂, and bromine (Scheufelen, A. 231, 179). From (3, 1, 6)-nitro-

o toluidine (N. a. W.).

Bromo-nitro-tolnone C_aH_aMe(NO.)Br [1:3:5]. [81°]. (N. a. W.); [86°] (W.); (270°). Formed (a) from brono-nitro-p-tolnidine [65°], (b) from nitro-(5, 1, 2)-bronno-tolnidine [143°], or (c) from bromo-nitro-o-tolnidine [181°] by the usual methods (Nevile a. Winther, C. J. 37, 431; Wroblewsky, A. 192, 203). Hence it can be prepared from a mixture of accetyl o- and p-tolnidines by successive bromination, nitration, saponification, and diazotisation.

Bromo-di-nitro-toluene C_oH₂Me(NO₂)₂Br. [104°]. From *m*-bromo-tolueno and fuming

lino, (Grete, A. 177, 258).

Di-bromo-nitro-toluené C₀1I₂Me(NO₃)Br₂ [1:4 or 6:2:3]. [c. 57]. From C₀1I₃(ClI₃)Br₂[28] by nitration (Nevile a. Winther, C. J. 37, 434).

Di-bromo-nitro-tolnene C_aH₂Me(NO₂)Br₂ [1:4:2:6]. [58°]. From C_aH(CH₂)(NH₂)(NO₂)Br₂ [124°-130°] by cthyl mitrito (Nevile a. Winther, C. J. 37, 445). Also from (2,4,1)-bromo-nitro-toluene, FeBr₂, and Br (Scheufelen, J. 2:31, 178).

Di-bromo-nitro-toluene C_aH_aMe(NO₂)Br₂ [1:5:3:4]. [63°]. From bromo-nitro-toluidine, C_aH_a(CH_a)(NO₂)(NH₂)Br [1:5:4:3] by diazo-perbromido reaction (Nevile a. Winther, C. J. 37, 447). Colourless plates (from alcohol).

Di-brome - nitro-toluene C_aH₂Me(NO₂)Br₂ [1:3:2:5]. [70°]. From bromo-nitro-e-toluidine, [143°], by diazo-perbromide reaction (Nevile a. Winther, C. J. 37, 448).

Di-bromo - nitro - tolnene C_eH_{*}McBr₂(NO₂) [1:4:6:2?]. [80]. By nitrating di-bromo-toluene from di-bromo-m-toluidine, [75°] (Nevile a. Winther, C. J. 37, 441).

Di. bromo - nitro - tolnene C₆H₂Me(NO₂)Br₂ [1:6:3:4]. [87°]. Formed by nitrating the corresponding di-bromo-toluene. Converted by reduction and diazo-reaction into (2, 4, 5, 1)-tribromo-toluene [113°] (Nevilc a. Winther, C. J. 39, 83).

Di -bromo - nitro -toluene C, H, Mc(NO.) Br., [1;4:2:5]. [88°]. From C, H, (CH.) (NO.) (NI.) Br., [181°] by the diazo-perbromide reaction. Formed also by nitrating C, H, (CH.) Br., [1:2:5]. Convorted by reduction and diazotisation into (2, 4, 5, 1)- tri-bromo-toluene [113°] (Nevile a. Winther, C. J. 37, 445; 39, 83).

Di . bromo . nitro . toluene C₆H Mc(NO₆)Br₂ [1:3:5:6]. [106°]. From bromo-nitro-o-toluidine, [181°], by exchange of NIL, for Br (Nevilo a. Winther, C. J. 37, 433).

ω-Di-bromo-nitro-toluene C_cH_s.CBr₂(NO₂). Phenyl-di-bromo-nitro-methane. Colourless oil. Formed by the action of bromine upon an aqueous solution of the di-sodium salt of nitro-benzylidene-philialide CoH. C(ONa) CNa(NO2) & H3. Volatile with

steam (Gabriel a. Koppe, B. 19, 1145).

Di-bromo-di-nitro-toluene C_uHMe(NO₂)₂Rr₂, [158³]. Formed by nitrating C_uH₃(C11₂)Br₂, [39⁵] (Nevile a. Winther, C. J. 37, 437).

Di-bromo-di-nitro-toluene C_aIIMe(NO.)₂Br₂ [105°]. Formed at the same time as the preceding (N. a. W.).

Di-bromo di-nitro-toluono C,11Mc(NO.),Br₂ [1:2:2:2:6]. [161°]. By nitration of (2, 6, 1)-di-bromo-toluene (N. a. W.).

Tri - bromo - nitro - toluene C HMc(NO₂)Br₃ [1:4:2:5:6], [106°]. From di-bromo-nitro-m-toluidine, [125 -130°], by the diazo-perbromide reaction. White needles (Nevile a. Winther, C. J. 39, 85.

Tri - bromo - nitro - tolueno $C_6 \mathrm{HMe}(\mathrm{NO}_2)\mathrm{Br}_4$, [107°]. [1:x:2:3:4]. Formed by nitrating tribromo-toluene, [44°] (N. a. W.).

Tri - brome - nitre - teluone C_e IIMe(NO.)Br_s [1:3:2:1:6]. [215°]. Formed by nitrating tribrone teluone [70°] (Wroblewsky, A. 168, 195).

Tri-bromo-di-nitro-toluene C_sMe(NO₂)₂Br₄ [1:??:2:3:4]. [217°-220°]. Formed by nitrating tri-bromo-toluene [44°] (Nevilo a. Winther, B. 13, 975).

Tetra-bromo-nitro-toluene C₀Me(NO₂)Br₄ [1:4:2:3:5:6]. [213°] (N. a. W.); [227°] (S.). Formed by nitration of C₄H(CH₂)Br₆ [117°] (Nevile a. Winther, C. J. 37, 450). From (2,4,1)-bromo-nitro-toluene, bromine, and FeBr₂ (Scheufelen, A. 231, 179).

Tetra-bromo-nitro-tolueno C_oMe(NO₃)Br₄ [1:2:3:4:5:6]. [212°]. From tetra-bromo-toluene, [111°].

Tetra-bromo-nitro-toluene C_aMe(NO_a)Br₄ [1:5:2:3:4:6]. [216]. By nitration of tetra-bromo toluene, [108].

BROMO - I. TRRO - TOLUENE SULPHONIC ACIDS Call McBr(NO.) SOall. The six following acids of this constitution have been described.

I. Formed by nitrating o-bromo-toluene sulphonic acid (Müller, A. 169, 42; Pagel, A. 176, 299). Deliquescent.—PbA'₂2aq.—NaA'aq.—KA'.—BaA'₂2aq.

II. By the action of funning IINO, on (2,1,4)-bromo-toluene sulphonic acid or on (2,1,4)-o-toluidine sulphonic acid; in the latter case the resulting diam-nitro-toluene sulphonic acid is boiled with HBrAq (Hayduck, 4, 172, 219; 174, 347). Minute needles; may be reduced to (1,3,4)-m-toluidine sulphonic acid. "BaA', 3aq.

Chloride C.H. MeBr(NO.)SO.cl. [220°].

Amide C.H. MeBr(NO.)(SO.NH.). Does not melt below 200°.

HI. From m-bromo-(olueno sulphonic acidand HNO₃ (Wroblewsky, A. 168, 169). - CaA'₂4½aq₀ -BaA'₂3½aq. -PbA'₂3aq.

IV. Formed by nitrating (3,1,2 or 6)-m-bromo-tolueno sulphonic acid (Weckwarth, A. 172, 200).—NaA'.—Ca V. 3aq.—BaA'_2 3'.aq.

V. Formed by intrating (4,1,2) -p-bromotoluene sulphonic acid (Hässelbarth, A. 169, 22), Deliquescont lamine.— AgA'.— BaA'_22aq.— CaA'_26aq.—PbA'_23aq.—SrA'_27aq.

VI. Formed by hitrating (4,1,8)-p-bromotoluene sulphonio acid (H.). Deliquescent ueedles.—BaA'2aq.—PbA'23aq.—SrA'25aq.

Di-bromo-nitro-toluene-di-sulphonic acid CeHBr. (NO.) Mc.SO.3H. From p-bromo-toluene di-sulphonio acid and boiling tuming HNO₃ (Kornatzki, A. 221, 197).—KA'aq.—BnA', 3!aq.
BROMO-NITRO-m-TOLUIC ACID C, H. BrNO₃

i.e. C. II MeBr(NO.)(CO.II). [176°], From bromo-m-toluie acid and IINO, (Fittig, A. 147,

84).—CaA', 3aq. -BaA', 3aq. *
Bromo-nitro-p-toluic acid

C₆H₂MeBr(NO₂)(CO₂H) [4:2:x:1]. [200°]. S. 1 at 15°. Formed by boiling bromo-cymenc (from thymol) with HNO, (S.G. 1.3). Lamina. -BaA', 4aq (Fileti a. Crosa, G. 16, 297).

Bromo-nitro-p-toluic acid $C_sH_sMeBr(NO_s)(CO_s11) + 4:3:x:11.$ [270°-180° From bromo-p-toluio acid and fuming IlNO, (Landolph, B. 5, 268). Needles (from water). BaA', aq. BROMO-NITRO-o-TOLUIDINE

 $\mathbf{C}_{6}\mathbf{H}_{2}\mathbf{Me}(\mathbf{NH}_{2})(\mathbf{NO}_{2})\mathbf{Br}$ [1:2:3:5]. [139°] (W.); [143°] (N. a. W.). Formed by nitrating bromoacetyl-o-toluidine, C_cH₃Mc(NHAc)Br [1:2:5], and removing acelyl (Wroblewsky, A. 192, 206; Nevile a. Winther, C. J. 37, 131). Gives, by nitrous gas and alcohol, CaH3Me(NO2)Br, [810] whence C_aH₃Me(NH₂)Br, [35]

Bromo-nitro-o-toluidine

 $C_a H_2 Mc(NH_2)(NO_a) Br [1:2:5:3]$. [181° cor.]. By brominating C, H, Me(NH,)(NO,) [1:2:3], [128] (N. a. W.). Converted by nitrons gas and alcohol into Call Mc(NO)Br, [81°] whence C_eH₃Me(NH..)Br [36°].

Bromo-nitro-m-toluidino

C₆H₂Me(NH₂)(NO₂)Br [1:3:6:5]. [88°]. Formed by nitrating bromo-acetyl-m-tolnidine, and then removing acetyl by H.SO, (2 vols.) and water (1 vol.) (Nevile a. Winther, C. J. 37, 630).

Bromo-nitro-m-toluidine

C_H_Me(NH_)(NO_)Br[1:3:2.6], [163°], Is formed in small quantity in the preparation of its i:omeride [181°].

Bromo-nitro-m-toluidino

 $C_0H_2Me(NH_0)(NO_0)Br = \{1:5:4:2\} = [181^\circ], From$ the acctyl derivative by saponification.

Acetyl derivative [110'-121']. Formed by nitration of bromo-acetyl m-toluidine (Nevile a. Winther, C. J. 37, 441).

Bromo-nitro-p-tolnidine

C.H.Mc(NH.)(NO.)Br [1:4:5:5]. [0::5]. Got by saponitying its acqtyl derivative. Orange needles. Converted by nitrons gas and alcohol into bromonitro-toluene [86°] (cf. Hand, A. 234, 157).

Acetyl derivative [2]1°]. From bromoacetyl-p-toluidine and HNO. Or from acetyl-p-toluidine by successive nitration and bromination (N. a. W.). White needles (from alcohol or diluto acetic acid) (Wroblewsky, A. 192, 202).

Di-bromo-nitro-m-toluidine

C₆HMc(NH₂)(NO₂)Br₂ [1:5:4:2:6]. [124°-130°]. From the acetyl derivative of bromo-nitro-mtoluidine [181] by heating with H.SO, (2 vols.) and water (1 vol.) and subsequent treatment with bromino (Nevile a. Winther, C. J. 37, 444).

BROMO-NITRO-m-XYLENE C. H. Me2(NO2)Br. (260°-265°). From bromo-m xylene and cold fuming HNO. Liquid (Fittig, A. 147, 31). -Di-brome-nitro-o-xylene $C_eHMe_2(NO_a)Br_a$ [1:2:8:4:5]. [141°]. Obtained by nitration of di-brome-o-xylene $C_eH_2(CH_3)_2Br_2$ [1:2:4:5] with cold furning HNO₂. Colourless needles (from alcohol) (Töhl, B. 18, 2561).

Di-bromo-nitro-m-xylene C.HMe_(NO2)Br20 [108°]. From di-bromo-m-xylene and IINO,

Needles (F.).

Di - bromo - nitro - p - xylene C. IIMc2(NO2)Br20 [112°], From di-br HNO₃ (F.), Needles. From di-bromo-p-xylene and fuming

Di-bromo-di-nitro-o-xylene C₆(CII₃)₂Br₂(NO₂)₂ [I:2:1:5:3:6]. [c. 250°]. Small needles. Nearly insol. cold alcohol. Formed by nitration of dibromo-o-xylenc C.H.(CH3),Br2 [1:2:4:5] (Töhl, B. 18, 2561).

BROMO-NITRO-XYLENE SULPHONIC ACID $C_aHMe_aBr(NO_a)(SO_aH)$ [1:3:6:x:4]. From nitrom-xylidine sulphonic acid by diazo- reaction (Sartig, 4.230, 341; B.18, 2190). Rhombic plates, v. sol. water and alcohol. -BaA'23aaq. -KA'aq.

BROMO-NONYLIC ACID v. BROMO-ENNOIO

DI-BROMO-OCTADECANE C181136Br2. Octadecylene bromide. [24']. Silvery plates. sol. alcohol. Formed by the addition of Br (1 mol.) to octadecylene (Krafft, B. 17, 1373).

BROMO-OCTANE v. OCTYL BROMIDE.

Di-bromo-octane CaH16Br2 Octylene bromide. From Br and octylene derived from castor oil (Rubien, A. 142, 297) or that from paraffin (Thorpe a. Young, Pr, 21, 193). Non-volatile oil.

Tetra-bromo-octane C, II, Br. Caprylidene tetra-bromide. From bromo-octylene and Br. Oil. BROMO-OCTONENE C, H, Br. (2017). From C, H, Br, (v. supra) and alcoholic KOH (R.).

BROMO-OCTYL-BENZENE $C_0H_1(C_0H_1)$ Br. (285°-287°). Formed by bromination of octylbenzene. Oil (Ahrens, B. 19, 2719).

BROMO-OCTYLENE C. H. Br. (185°). From di bromo-octane and alcoholic KOII (Rubien, A. 112 297). With Br it gives an oily tri-bromedecáne.

Di-bromo-octylene C₈II₁₄Br₂. S.G. 10 1.568. Conylene bromide. From conylene and Br (Wertheim, 4, 123, 182).

BROMO-OCTYL-THIOPHENE

C₄SH₂(C₈H₁₇)Br. (285°-290°). Colourless oil solidifying to plates at 5°. V. sol. ether, insol. Colourless oil water. Formed by shakir; octyl-thiophene with bromine-water (Schweinitz, R. 19, 644).
BROMO-OLEIC ACID C₁₈H₃₃BrO₂. From di-

bromo-stearic acid and alcoholic KOII (Over-

beck, A. 140, 47).

Di-bromo oleic acid C18H32Br2O20 From

stearolic acid and Br (O.).

BROMO-ORC:N C,11 Mc(OH) Br. From ordin and bromine-water (Lamparter, A. 134, 258). Crystals; m. sol. hot water, v. o. sol. alcohol and other. Solutions are ppd. by lead subacetate.

Di-bromo-orori

Mothyl derivative Call(CH3)(OMe)(OH)Br2. [146°]. White needles. Prepared by bromination of the mono-methyl ether of orein (Tiemann a. Streng, B. 14, 2002).

Di-methyl derivative C.11(C112)(OMe), Br₂. [160']. Colourless plates. Sol. alcohol, other, and benzene, insol. water and ligroin. Prepared by bromination of the di-methyl ether of orein (B. 14, 2001).

Tri bromo-orcin C₀(OH₂)Br₃(OH)₂ [103⁵]. From orcin and Br (Stenhouse, Tr. 1848, 87; Laurent a. Gerhardt, A. Ch. [3] 24, 317; Lamparter, A. 134, 257; Hesse, A. 117, 311; Stenhouse a. Groves, A. 203, 298). Is formed by heating penta-bromo-orein with formic acid. Needles; insol. water, sol. alcohol and other.

Diacetyl derivative [143]. White needles. Formed by the action of Ac.O on penta-bromo orein (Claussen, B. II, 1440).

Penta - bromo - orcin C.H.Br.O. C.McBr. (OBr). ? [126]. From orein and excess of bromine-water. Triclinic crystals (from CS.). At 160° it gives off Br., leaving C. H. Br. O. (Stenhouse, A. 163, 180; Liebermann a. Dittler, A. 169, 252).

Bromo-B-orein v. Bromo-Bettoecin.

DI.BROMO OXAL-ETHYLINE v. DI-BROMO-METHYL-ETHYL-GLYOXALINE.

BROMO-OXINDOLE v. OXINDOLE.

BROMO-OXY-ACRYLIC ACID. Phenylderivative CaH.BrOai.c. CHBr:C(OPh).CO.H. [138]. From phenyl-oxy-rencobronic acid CHO.CBr.C(OPh).CO.H and KOH (Hill a. Stevens, Am. 6, 190). Needles (from water); v. e. sol. alcohol and other.—KA'. `BaA', 5aq. CaA', 5aq. - AgA'

BROMO - OXY - AMIDO - BENZOIC ACID. Methyl derivative $C_{10}H_aBrNO_a$ C, H, Bi (OMe) (NH.,) CO, H. [185]. Bromo-amidoanisic acid. From the corresponding nitro-acid. Neodles, sl. sol. water.—CaA'₂ 5 [aq.—BaA'₂ 2aq.—HA'IICl [186"] (Balbiano, G. 14, 245).

BROMO-DI-OXY-ANTHRAQUINONE

C₁₁H₁BrO₄ i.e. C₁₁H₂O₂(OH)₂Br. Bromo-alizarin. From alizarin (3 pts.) and Br (2\frac{1}{2} pts.) in CS₂ at 190° (Perkin, C. J. 27, 401). Turks of orange needles; may be sublimed. KOHAq forms a blue solution, exhibiting the same absorption bands as alizarin. HNO3 forms phthalic acid. The same bromo-alizarin, or an isomeride, is formed by treating tri-bromo-anthraquinone with KOH. It melts at 280° (Dichl, B. 11, 190).

Bromo-tri-oxy-anthraquinone

C₁,H₁O₂(O11)₃Br. Bromo purpurin. [276°]. From Br and purpurin, or its carboxylic acid, or by warming di bromo-purpurin (v. infra) with conc. HoSO, Red needles (Plath, B. 10, 615, 1619; Schunck a. Roemer, B. 10, 554).

(B. 1,3,2)-Di-bromo-oxy-anthraquinone $\mathbf{C}_{t4}\mathbf{H}_{6}\mathbf{Br}_{2}\mathbf{O}_{3}$ i.e. $\mathbf{C}_{8}\mathbf{H}_{4}(\mathbf{C}_{2}\mathbf{O}_{2})(\mathbf{O}_{6}\mathbf{HBr}_{2}(\mathbf{OH}), [208])$. Formed, together with di-bromo-phenol by heating tetra-bromo-phenol-phthalein with excess of II₂SO₄ at 150° (Bacyer, A. 202, I36). Slender • vellow needles; its alcoholic solution shows reddish fluorescence. Its solution in alkalis is reddish-brown. NaOH at 200 gives alizarin.

Acetyl derivative C, H, AcBr, O, [190].

Di-bromo di-oxy-anthraquinone C, H,O, (OH) Br. Di-br mo-alizaria. [170]. Prepared by the action of Br in presence of I on alizarin. Small brownish-red needles. Combines with mordants (Diehl, B. 11, 190).

Di-bromo-di-oxy-anthraquinone

C_{1.}H₄O₂(OH), Br₂, Di-bromo-purp [227°-230°] (P.); [231°] (S. a. R.). Di-bromo-purpuroxunthin. From purpuroxanthin and Br (Plath, B. 9, 1205). From muniistin and Br (Schunck a. Roemer, C. J. 33, 424). Orange needles (from HOAe). Warm sonc. H.SO, forms bromo-purpurin. Selt.—(NH4)2A".

* Tri-bromo-tri oxy-antamquinone

C₁₄H₂O₂(OH)₃Br₃. Tri-bromo-flavepurpurin. [284°]. From flavopurpurin in HOAe and Br. Orange needles. Its alkaline solutions are orange (Schunck a. Roemer, B. 10, 1225).

Tetra-bromo-di-oxy-anthraquinone

H2O2(OH)2Brp. Tetra-bromo-alizarin. From alizarin and excess of iodine bromide at 180°. Does not combine with mordants (Diehl, B. 11, T91).

BROMO-O-OXY-BENZOIC ACID C. II BrO. i.e. C_aH₃Br(OH)CO₂H [3:2:1]. Brom> selleytic acid. [181] (L. a. G.); [220] (H. a. E.). From the corresponding bromo amido benzeic acid by exchange of NH₂ for OH (Hübner & Emmerling, Z. 1871, 709) or from (3, 5, 2, 1) brome-amideoxy-benzoic acid by climinating NII, (Lellmann a. Grothmann, B. 17, 2725). Needles, v. sl. sol. cold water, v. c. sol. alcohol. Fe Cl, gives a violet colouration. - CaA', 12aq: v. sol. water. BaA'23 aq: prisms.—PbA'2 (II.). -PbC, H, BrO,

Bromo-o oxy-benzeic acid C.H.Br(Oll)(CO.H) [5:2:1]. [165°]. From salicylic acid and Br or PBr₂ (Gerhardt, A. Ch. [3] 7, 217; Cahours. 4. Ch. [3] 10, 341; 13, 99; Henry, B. 2, 275; H. a. E.). Also from the corresponding amido-m-bromobenzoic acid (H. a. E.). Needles (from water). FogCl, gives a violet colouration. BaA', Saq. PbA'... $PbC_{s}H_{g}BrO_{g}-CuA'_{g}+AgA'_{s}$

Methyl ether MeA'. [38] (Henry); [61°] (Peratoner, G. 16, 405). (265°). From methyl salicylate and Br or PBr_s. Trimetric prisms or needles. Coloured violet by Fe,Cl,

Methyl derivative C. H. Br(OMe)CO.H. [119°]. — BaA', 3aq. — CaA', Laq. $-\Lambda_{\mathbb{R}}\Lambda'$ aq. Methyl ether C.H.Br(OMe)CO.Me. -[304].(295 ') (P.).

Ethytderivative C. II Br(OEt)CO.II. [130]. BaA', 4w; --CaA', 2aq. Methylether C₆H, Br(OEt)CO₂Mc. [49]. (301). Propyl derivative C₆H₃Br(O'r)CO₂H.

[62°]. Methyl other C.H.Br(Ofr)CO Me. (323 ').

Isopropyl derivative C. H. Br(OPr)CO.H. [101°]. Methyl other (301°).

Bromo-p-oxy benzoie acid. Methyl derivative C. H.Br(OMe)CO.H [3:1:1]. Bromo-

anisic acid. [214]. [218' cor.]. Formation.—1. From anisic acid and Pr (Laurent; Cahours, A. 56 311; Salkowski, B. 7, 1013). -2. By oxidising the methyl ether of . bromo-p-cg/soi (Schall a. Dralle, B. 17, 2531).

Properties. Needles; may be distilled or sublimed. Insol. water.

Salts.—A,A'.—BaA', 33,.q.—BaA', 4aq.—CaA', 6aq.—CnA', 24aq.—M,A', 5aq.—NnA', 2aq.—PbA', 3aq.—ZnA', 3aq.—Ethyi ether C, H,Br(CMe)(CO,Et). [74°]

(Crespi, G. 11, 419).

 $C_aH_aBr(OMe)(CO_2NH_a).$ Amide insol. water.

Bromo-p-oxy-bouzoic &id. Methyl derivative C.H.Br (OMe) CO.H. [212°]. Ethyl bromoanisate [740] is converted into an isomerido [60°] by heating with NaOEt at 180° for 20 hours; on saponification it yields the acid which crystallises in needles, sl. sol. water. Potash fusion forms protocatechnic acid. IINO, gives the methyl ether of (2,4,6,1) - bromo - di - nitrophenol.

Salt .- ZnA', 4aq (Balbiano, G. 11, 409).

Ethyl ether EA. [60°] (v. supra). This acid is possibly identical with the preceding.

| Fe₂Cl₄ turns its solutions yellowish-red. Sodium amalgam forms p-oxy-henzoic acid.—CaA. 289.

Bromo-di-oxy-benzoic acid. Methyl deri-C₆H₂Br(OH)(OMe)CO₂H [x:4:3:1]. Bromo-vanillic acid. [193]. From its acetyl derivative. Needles (containing aq). Acetyl derivative CaHaBr(OAc)(OMc)COaH. [167]. From acetyl-vanillic acid and Br (Matsmoto, B. 11, 138).

Di-methyl derivative

 $G_{\rm s}H_{\rm p}{\rm Br(OMe)_2CO_2H}$ [x:4:3:1]. Bromo-veratric acid. [184]. From veratric acid and Br (M.).

Methylene derivative C.H.Br(O.CH.)CO.H or C.H.(O.CHBr)CO.H. Bromo piperoriplic acid, [205°]. From bromopiperonal and KMnO, (Fittig a. Mielck, A. 172,

Bromo-di-oxy-benzoic acid

 $C_0H_2Br(OH)_2CO_2H = [x:1:3:5], [253°],$ From s-di-oxy benzoic acid and bromine water (Barth a. Senhofer, A. 164, 115). Needles (containing aq). Potash-fusion forms gallic acid. FegCla gives a yellowish brown colour.-CaA', 8aq.-Ag.A".

Bromo-di-oxy-benzoic acid CallaBr(OH)2CO.H [x:2:6:1]. [184°, anhydrous]. From c-di-oxybenzoic acid in ether and Br (Zchenter, M. 2, 480). Prisms (centaining aq). Fe₂Cl₆ gives a violet colour to its aqueons solution.—AgA'aq. $\operatorname{BaA'}_{2}7_{2}^{1}\operatorname{aq}$. $-\operatorname{CuA'}_{2}4_{2}^{1}\operatorname{aq}$. $-\operatorname{PbA'}_{2}3\operatorname{aq}$. $-\operatorname{KA'}1_{2}^{1}\operatorname{aq}$. Bromo tri oxy benzoic acid

C. HBr(OH), CO. H. Bromo-gattic acid. [above 200°]. From gallic acid and Br (Hlasiwetz, A. 142, 250; Grimaux, Z. 1867, 431). Monoclinic; sl. sol. water.

Di-bromo-o-oxy-benzoic acid

C.H.Br. (OH)CO.H [5:3:2:1]. Di-bromo-salicylic acid. [219°] (R.); [223°] (L.a. G.). From salicylic acid and Br or from (3,5,2,1)-bromo-amido-salicylic acid by the diazo- reaction (Cahcurs, A. Ch. [3] 10, 339; 13, 102; Rollwage, B. 10, 1707; Lollmann a. Grotlmann, B. 17, 2727). a violet colour with Fe, Cla. Heated with dilute H2SO4 it gives (3,5,2)-di-bromo phenol [36°]. -BaA', 4aq.

Methyl ether Coll_Br_(OH)CO_Mc: [149°]; from methyl salicylate and Br (Peratoner, G. 16, 405). Long needles, sl. sol. alcohol.

Methyl derivative C.H.Br.(OMc)CO.H:
[194°]. — Salt BaA(2!aq. Methyl ether
C.H.Br.(OMe)CO.Me: [53°]; needles.

Ethyl derivative G.H.Br. (OKt).CO.H: 6°]; white needles. Methyl ether CaH2Br2(OEt).COMe: [43°]; needles.

Di-bromo-o-oxy benzoic acid $\mathbf{C}_{\mathbf{t}}\mathbf{H}_{\mathbf{J}}\mathbf{Pr}_{\mathbf{t}}(\mathbf{OH})\mathbf{CO}_{\mathbf{J}}\mathbf{H}$ [9:3:20r6:1] [218°]. From (4,3,1)-di-bromo-benzoic acid [229°] by nitration, reduction, and dinzotisation (Smith, B. 10, 1706). Gives a violet colour with Fe₂Cl₆.

Di-bromo-o-oxy-benzoic acid C₆H₂Br₂(OH)CO₂H. [221]. Formed as a by-product in converting (5,2,1)-bromo-nitro-benzoic acid [250°] into di-bromo-benzoic acid by the diazo- reaction (Hübner a. Lawrie, B. 10, 1706). Fo2Cl gives a violet colour.

Di-bromo-p-oxy-benzoic acid

C_uH₂Br₂(OH)CO₂H. [268°]. From di-bromoanisic acid and conc. HI (Alccei, G. 15, 242). One of the products of the dry distillation of sodium di-bromo-anisate (Balbiano, G. 13, 69). Long needles, insol. water, sel. alcohol and ether.

amalgam forms p-oxy-benzoic acid.—CaA', 3aq.

Methyl derivative C,H,Br,(OMo)CO,H [3:5:4:1]. $Di ext{-}bromo ext{-}anisic acid [207°](R.); [214°]}$ (C.). From anisio acid, Br, and water at 1205 (Reinecke, Z. 1866, 366; Crespi, G. 11, 425). Converted by prolonged action of Br and water into tri-bromo-anisol [87°].—NaA' 3aq.—AgA'.-BaA'24 aq. Ethyl ether C.H.(OMe)(CO.Et): [88°]; plates.

Di bromo-di-oxy-benzoic acid

CaHBr₂(OH)₂CO_.H. [214°]. From (3, 2, 1)-dioxy-benzoic acid and Br (Zehenter, M. 2, 475). Needles (containing aq); m. sol. hot water. Fe₂Cl₆ turns its solution violet; conc. H₂SO₄ gives a green colonr. Heating with vater forms di-bromo-resorcin. — KA' 3 aq. — CaA' 28 aq. — PbC.H.Br.O. -CuA'.aq. -AgA'.

Di bromo tri oxy benzoic acid

C.Br.(Oll),CO.H. Di-bromo-galli acid. [140°] (G.); [150] (E.). From gallie acid and Br (Grimaux, Z. 1867, 431; Etti, B. 11, 1882). Fc.Cl, gives a blue-black colour in its aqueous solution.

Tri-bromo-o-oxy-bonzoic acid

C.HBr. (OH)CO.H. Tri-bre .w - salicylic acid. From Br and salicylic acid Small prisms, insol. water (Cahours, A. Ch. [2] 13, 104). Tri-bromo m-oxy-ben oic acid

C. HBr. (OH) CO. H. [147" From m.oxy-benzoio acid (1 mol.) and Br (3 mols.) (Werner, Bl. [2] 46, 276).

Tri bromo-di-oxy-benzoic acid

C₀Br₃(OH)₂CO₂H. [183]. From (5, 3, 1)-dioxy-benzoic acid and Br Barth a. Senhofer, A. 159, 225). Tables (from water). Potash-fusion reproduces s-di-oxy-benzoic acid.

BROMO - o - OXY - B INZOIC ALDEHYDE $\mathbf{C}_{7}\mathbf{\Pi}_{1}\mathbf{BrO}_{2}$ *i.e.* $\mathbf{C}_{6}\mathbf{H}_{3}\mathbf{Br}(\mathbf{C.H})\mathbf{CHO}_{2}$ Bromo - sali cytic aldehyde. [997]. From salicylic aldehyde and Br or PBr, (Löwig, P. 46, 57, 383; Piria, A. Ch. [2] 69, 281; Henry, B. 2, 275). Laminæ; insol, water, sol. alcohol and other. Combines with KHSO3.

*Br(OMe)CHO. Methyl derivative [114]. From methyl-salicylic aldehyde and Br (Perkin, A. 145, 304). Flat prisms (from alcohol). Ethyl derivative C.H.Br(OEt)CHO: [68°]; prisms.

Bromo-p-oxy-benzoic aldohyde

C_aH_aBr(OH)CHO. [180°]. From p-oxy benzoic aldehyde and Br. V. sol. alcohol and ether, v. sl. sol. water: Combines with KHSO, (Herzfeld, B. 10, 2198).

Di-bromo-o-oxy-benzoic aldehyde

C.H.Br. (OH)CHO. Di-bromo salicylic-alde nyde. Prisms. From salicylic aldehyde and Br (Hee lein, J. pr. 32, 65).

Phenyl-hydrazide

C.H.Br₂(OH)CH;N.HPh: [148°]; v. sol. alcohol, benzene, ether, and CHGl_{st} insol. water. The mons-acctyl-derivative C, H, Br, (OAc)CH: N. HPh forms fine needles [1884], nearly insoluble in ether. The di-acetyl derivative C. H. Br. (OAc) CH: N. AcPh crystallises in white needles, [158], easily soluble in ether; it is formed by brominating the di acetyl derivative of the phenyl-hydrazide of salicylic aldehyde (Rossing, B. 17, 3008).

Di-bromo-p-oxy-benzoic aldehyde C₆H₂Br₂(OII)CHO. [181°]. From p-oxy-benzoio aldehyde (1 mol.) and Br (2 mols.) (Wernor, Bl. [2] 46, 277).

BROMO.OXY.BUTYRIC ACID C.H.BrO. [102°]. From di-bromo-butyric acid and baryta. Laminæ (Petrieff a. Eghis, J. R. 7, 179). -BaA'2.—AgA'.

Bromo-oxy-bntyric acid

CHa.CHBr.CH(OH)CO.H or CH3.CH(OH).CHBr.CO.H. An uncrystallisable syrup obtained as a residue when aB-di-bromobutyric acid is distilled with water (C. Kolbe, J. pr. 133, 389; cf. Erlenmeyer a. Müller, B. 15, 49). Bromo-oxy-butyric acid

CH, CHBr.CH(OH).CO.H or CH₂.CH(OH) CHBr.CO₂I1. [90°]. From Bmethyl-glycidic acid O CH.CH. and HBr (Melikoff, Bl. [2] 43, 116). Prisms. Probably identical with the preceding. Bromo-oxy-iso-butyric acid

CH_Br.C(OH)Mc CO.H. [101°]. Formed by boiling di-bromo-iso-butyric acid with water, and extracting with ether (K.). Also from IIBr and a-methyl-glycidic acid O CH2. Needles; sol. hot benzene, insol. CHCl3 and CS2. Not affected by boiling water. Reduced by the action of sodium amalgam on its aqueous solution, kept neutral by ILSO, to oxy-iso-butyric acid, [79°].

BROMO OXY-CINNAMIC ACID v. Bromo-

COUMARIC ACID.

BROMO-(B. 4)-OXY-(Py. 4)-ETHYL-QUINO-LINE TETRA-HYDRIDÈ C,H,Br(OH)EtN.

Ethyl ether [35°]; long monoclinic prisms, a:b:c=0.7902:1:0.5828. Formed by bromination of ethyl-kairine (ethyl ether of oxy-ethyl-quinoline tetra-hydride), or by ethylation of the ethylether of bromo-oxy-quinoline-tetra-hydride. Tho picrato forms yellow needles [174°] (Fischer a. Renouf, B. 17, 762)

DI-BROMO-OXY-INDONAPHTHENE

 $C_9H_4\mathrm{Br}_2\mathrm{O}~\textit{i.e.}~C_9H_4 \overset{\mathrm{CO}}{<}\mathrm{CBr} \overset{\mathrm{CBr}}{>}\mathrm{CBr}. \quad \textit{Phenylena-discontinuous}$ bromo-acetylene ketone. [123°]. Obtained by heating di-bromo-cinnamie acid C.H., CBr; CBr, CO, H

with cone. H.,SO, Yollow needles.

Oxim C, H,Br,(NOH): [195°]; yellow needles.

Anilide: [170°]; red needles.

Di-bromide C, H,OBr,: [124°]; prisms

(Roser, B. 20, 1273).

BROMO-OXY-MALEÏC ACID Phenyl derivative CO.H.CBr:C(OPh).CO.H. [104]. From the phenyl derivative of exy-mucobromic acid and Ag₂O (Hill a. Stevens, Am. 6, 187). Needles. $-Ag_2A'$

p-BROMO-6-OXY-MESITYLENE C. II, BrO i.e. C_bH₂(CH₃)₂Br(CH₂OH) [5:3:4:1]. p-Bromomesityl alcohol. [66°]. Obtained from p-w-dibromo-mesitylene (p-mesityl bromide) by treatment with KOAc and saponification of the acetate. Pointed needles. V. c. sol. alcohol, ether, and benzene, sl. sol. cold petroleum-other, insol. cold water. Decomposes on distillation with separation of H₂O and formation, amongst other products, of p-bromo-mesitylenic aldehyde C.H.(CH3).Br(CHO). By oxidising agents it is readily converted into p-bromo-mesitylenic acid [214°] (Schramm, B. 19, 213).

eso-Bromo-ω₁ω₂-di-oxy-mesitylene C₆H₂Br(CH₃)(CH₂OH)₂. [121°]. S. 3½ at 100°. From the corresponding tri-brome-mesitylene (200°-215°) by boiling with water and PbCO. (Colson, A. Ch. [6] 6, 98; C. R. 97, 177). Pearly scales; v. sl. sol. cold water, m. sol. alcohol. Boiling HClAq forms CaH2Br(CH3)(CH2Cl)2 [75°]

BROMO-OXY-8-METHYL-CUMARILIC ACID

 $C_0H_2Br(OII)$ < CMe $<math>C_0CO_2H$. [221°]. Formed by boiling bromg-β-methyl-umbelliferon-dibromide C_bH₂Br(OH) CMeBr.CHBr with alcoholio KOII. Colourless needles. V. sol. alcohol and ether, sl. sol. benzene, insol. water. Cold H2SO, gives a colourless solution which becomes violet on heating. FegCla gives a yellow colouration with the alcoholic solution

(Peelmann a. Cohen, B. 17, 2131). BROMO . OXY METHYL ETHYL PYRIMID . INE C_2H_3 .C $\langle N.C(OH_3) \rangle$ CBr. [195°]. Formed by bromination of oxy-methyl-ethyl-pyrimidine. Long colourless glistening needles. Sl. sol. water. C, II, N, Br(OII) aq: very soluble long white needlos (Pinner, B. 20, 2362).

BROMO-OXY-DI-METHYL-PYRIMIDINE

 $CII_{r}.C \stackrel{\mathrm{N.C(OH)}}{\leqslant_{\mathrm{N.C(CH)}}} > CBr$. Formed by bromination of oxy-di-methyl-pyrimidino. The hydrobromide (B'IIBr) forms colourless needles, m. sol. water, v. sol. alcohol (Pinner, B. 20, 2361).

(Py. 2,3,1)-BROMO-OXY-METHYL-QUINOL-

 $C_{10}\Pi_8NOBr$ i.e. $C_6H_1 < CMe:CBr \\ 1 : C.OH$? Bromo.

oxyquinaldine, or bromo-quinoxyl. [o. 258°]. Formed by the action of cold cone. H₂SO₄ upon the amlide of bromo-aceto-acetic acid CII_xC(OH):CBr.CONHPh. Also from (Py. 3, 1)oxy-methyl-quinoline and bromine-water (Knorr, B. 17, 2875; A. 236, 91). Fine silky needles. Sl. sol. alcohol. ether, and chloroform. Dissolves in aqueous acids and alkalis.

Tri-bromo (Py.1) oxy-(Py.3) methyl-quinoline C₂H. MeBr₃(OII)N. [275°]. Formed by bromination of (Py. 1, 3) oxy-methyl-quinoline. Insol. alcohol (Conrad a. Limpach, B. 20, 949).

Bromo-(Py.3)-oxy-(Py.1,1)-di-methyl-quinoline C. H. E. NO. Bromo-methyl-lepidone. [172°]. From the corresponding oxy-dimethyl-quinoline and bromine-water (Knorr, A. 236, 110). Spherical aggregates of needles (fymu alcohol). Insol. water and NaOHAq, v. scl. dilute acids. BROMO-OXY-(α)-NAYHTHOIC ANHYDRIDE

 $\mathbf{C_{11}H_{5}BrO_{2}\textit{i.e.}}\mathbf{C_{10}H_{.0}Br} \overset{\mathrm{CO}}{\underset{O}{\bigcup}} \text{[192°].}$ browination of oxy-naphthoic auhydride dissolved in CS, Small white

needles (Ekstrand, B. 19, 1139). Bromo-oxy-(a)-naphthoquinone

CO.C(OH) C.H. [197°]. ∖CO.ČBr

Formation .- 1. From di-bromo-(a)-naphthoquinone [151°] by boiling with aqueous NaOH or Na,CO,; the yield thing 60 p.c. of the theoretical (Diehl a. Merz, B. 11, 1064).—2. From oxy.(a)-naphthoquinone and Br.—3. Prepared by the action of alcoholic H2SO, on di-bromo-(a)naplithoquinone-anilide, p-bromo-aniline being simultaneously produced (Baltzer, B. 14, 1901). By the action of alkali upon bromo-β-naphthoquinone (Zincke a. Gerland, B. 20, 1515). -5. By boiling brome-amide-(a)-naphthogninene C,H, CO,CBr with dilute alkalis (Z.). 6. From bromo oxy-(a)-naphthoquinone-imide $C_uH_i < \frac{CO}{C(NH)CBr}$ by boiling with cone. HCl or by treatment with alcoholic NaOH (Z.).

Properties. - Yellow needles; v. sl. sol. water, sl. sol. ether, v. sol. alcohol. Oxidation gives phthalic acid.

Salts. KA'aq: red needles.—BA. S. 07 at 13°. -- AgA'.

Bromo-oxy-(a)-naphthoquinone. [202°]. From the anilide [1974] of di-bromo-naphthoguinono [218°] by boiling with aqueous Na CO, (Miller, Bl. [2] 43, 125). Oxidises to phthalic acid; it should therefore be identical with the preceding.

BROMO - OXY - (a) - NAPHTHOQUINONE -CO - C(OH). [c. 265°]. Formed C(NII).CBr

by boiling bromo amido - (a) - naphthoquinoneimide C_sH₄<CO - C(NH₂) with dilute NaOII.

Formed also by the action of NH₃ upon bromo-(β)-naphthoquinoue. Brownish - red glistening needles. By boiling with cone. HCl or by treatment with alcoholic NaOII it is converted into bromo-oxy-(a) - naphthogninone. The sodium. salt forms red needles; the salts of the heavy metals are sparingly soluble pps.

Acetyl derivative: [270°]: red hair-like needles (Zincke a. Gerland, B. 20, 1514).

BROMO-OXY-NAPHTHOQUINONE SUL-

PHONIC ACID C₁₆II BrSO₆ i.e. C₁₆H₃O₆Br(OH)(SO₆H). From (β)-naphthol suiphonic acid and Br, di-brome oxy-naphthoquinone being also formed in small quantity (Armstrong a. Graham, C. J. 39, 138; Armstrong a. Streatfeild, C. J. Proc. 1, 232).— BaC H BrSO

BROMO-OXY-NICOTINIC ACID v. BROMO-OXY-PYHIDINE CARBOXYLIC ACID.

BROMO-OXY-OCTOIC ACID Call Brog i.e. CII, CHBr.CII, CH(CO_II).CII, CH(ÖH).CH, Bromo-oxy-di-propyl-acetic acid.

CH. CH.CH. Lactone CH, CHBr.CH, CH

S.G. 15 1.394. From di-allyl-acetic acid and HBr, the compound (CH, CHBr, CH,), C11.CO, H being probably first formed (Hjelt, A. 216, 73). Oil. Insol. cold water, v. sl. sol. warm water. Insol. cold NaOH. Boiled for a long time with water or aqueous Na2CO3 it appears to form tho

lactone CII,:CH.CH,.CH CH2.CHMo

OCTENOIC ACID. Tri-bromo-oxy-octoic acid. Lactone

CH.CH.CH.Br CH2Br.CHBr.CH2.CH From

di-allyl-acetic acid and bromine in ohloroform (Hjslt, A. 216, 76). Oil. V. sol. other. Insol. oold NaOHAq. Boiled with aqueous Na, CO, it forms (CH., (OH). CH(OH). CH.), CH. CO, Na.

Tri-bromo-di-oxy-octoio acid. Lactone

C_sH₁₁Br₂O₃ i.e. CH2Br.CHDr.CH., C(OH).CH., CH(CH.Br), O.CO.

From so-called 'di-allyl-oxalic acid' and bromine (v. Oxy-octinoic acid) (Schatzky, J. pr. [2] 31, 485).

Tetra-bromo-oxy-octoic acid C.II. Br.O. i.e. (CII_Br.CHBr.CH_).C(OH).CO_H. From so-called 'di-allyl-oxalic acid' and Br (Saytzeff, A. 185, 189). Oil: readily splits up into HB7 and the preceding lactone.

Ethyl other Eth'. Oil (Schatzky, J. R.

DI - BROMO - HEXA - OXY - DIPHENYL. Methyl ether CmH2Br2(OMe). [140°]. From the methyl ether of hexa-oxy-diphenyl and Br (Ewald, B. 11, 1623). Needles (from alcohol or HOAc); cone. ILSO, forms a blue solution.

Tetra-bromo-di-oxy-diphenyl C, H, Br, O, i.e. C.H.Br. (OH).C.H.Br. (OH). [264°]. From dioxy-diphenyl and Br (Magatti, B. 11, 2267; 13, 225). Also by reduction of bromo-rosogninone (Baeyer, B. 11, 1301). Fuming IINO₃ forms becaused real scales of (C.H.Br.O)₂. Tetrabrownish-red scales of (C, H2Br2O)2. bromo-diphenyl-quinone.'

Acctyl derivative C12H,Ac,Br,Or [245°];

ncedles.

Tetra-bromo-tetra.oxy-diphenyl

C12H2Br1(OH) . Tetra-bromo-diresorcin. From tri-bromo-resogninone CaHBr3Og(?) and HS or Sn and HCl. Needles (from HOAc). Turns brown at 230° and decomposes at 280°. Insol. water, v. sol. alcohol and other. Sodinm amalgam gives diresorein. Red-hot zinc-dust gives diphenyl.

Acelylderivative C12H2Br (OAe)4 [195]; needles (from alcohol) (Benedikt, M. 1, 352; B. 11, 2170).

Deca-bromo-tetra-oxy-diphenyl C. Br. (OBr) Formed by adding Br and IICl to a solution of diresorcin in aqueous potash (Benedikt a. Julius, Unstable crystals, gives off Br M. 5, 179).(2 mols.) at 185°. SO, reduces it to Cp. Br. (OH).

BROMO DOXY-PHENYL-ACETYLENE Methyl derivative $C_{\theta}H_{3}Br(OMe).CCH.$ [75°]. Formed by heating the methyl derivative of tri-bromo p-oxy-phenyl-propionic acid C_bH_aBr(OMe).CHBr.CHBr.CO₂H with aqueous KOH (30 p.c.). Plates. Gives an unstable greenish-yellow compound with ammoniacal CuCl₂ (Eigel, B. 20, 2538).

DI-BROMO-DI-OXY-DI-PHENYL-AMINE CoHa(OII).NH.CoH.Br.OII [4:6:2:1]. Leuco-dibromo-quinone-phenol-imide. [170°]. Colourless prisms. V. sol. all ordinary solvents, except water. Formed by reduction of di-bromo-quin-one phenol-imide (Mölilan, B. 16, 2848). a · ξ · eso · eso · TETRA - BROMO · o · OXY · β.

PAENYL-BUTYRIC ACID. Methyl derivative C.H.Br.(OMe).CHBr.CMeBr.CO.H. [c. 200°]. l'rom (a) or (β) methoxy-phenyl-crotonic acid and bromine vapour (Perkin, C. J. 39, 434). Crystalline powder (from eliloroform).

Tetra-bromo-di-oxy-di-phenyl-methane C₁₃H₈Br₁O₂ i.e. CH₂(C₆H₂Br₂,OH)₂. [225°]. From di-oxy-di-phenyl-methane and bromine-water In ethereal solution it forms an unstable crystalline compound with hydrio bromide C, H, Br, O2 (Beck, B. 10, 1837).

BROMO-OXY-PHENYL-METHYL-PYRAZOLE

 $C_eH_s.N < CO-CHBr > Bromo-phenyl-methyl$ pyrazolone. [0.130°]. From oxy-phenyl-methylpyrazole and Br in glacial acetic acid (Kuorr, A, 238, 176). Sol. alkalis, and dilute acids; insol. water. Sl. sol. ether, v. sol. glacial HOAc and chloroform. In alcoholic solution it slowly forms pyrazole-blue. Fe.Cl, forms pyrazolo-

Di-bromo-oxy-phenyl-methyl-pyrazole

C₁₀H₈N₂OBr₂ i.e. Pan < CO - CBr₂ >. Di-bromophenyl-methyl-pyrazolone. [80°]. From oxyphenyl-methyl-pyrazole (I pl.) and Br (2 pts.) in acctic acid solution (Knorr, A. 238, 177). Sol. alcohol, HOAc, ether, and CHCl₃; insol. water, alkalis, and acids. Not attacked by Fe, Cl, Reduced by Sn and HCl or fuming HI to oxyphonyl-methyl-pyrazole.

BROMO-OXY-PHENIL-METHYL-PYRIMID.

1NE C₁₁H_pN_zBrO i.e. C_gII_s. N.C(OII) CBr.

[260°]. Formed by bromination of oxy phenylmethyl-pyrimidine. Glistening needles (Pinner, B. 20, 2361).

BROMO-O-OXY-PHENYL-PROPIOLIC ACID.

Methyl derivative

C₆H₃(OMe)Br.C:C.CO₂H. [168°] (with decomposition). From the methyl derivative of tribrome-oxy-phenyl-propionic acid (q. v.). Short white needles (from benzene).

 $\begin{array}{l} \textbf{BROMO} - o \cdot \textbf{OXY} + \beta \cdot \textbf{PHENYI}_c \cdot \textbf{PROPIONIC} \\ \textbf{ACID} \cdot \textbf{C}_o \textbf{H}_a \textbf{BrO}_a \cdot i.e. \cdot \textbf{C}_o \textbf{H}_a \textbf{Br}(\textbf{OH}) \cdot \textbf{CH}_a \textbf{CO}_a \textbf{H}. \end{array}$ Bromo-melilotic acid. [142]. From its anhydrido by boiling with water. Rectangular tables (from chloroform). Sol. alcohol, sl. sol. water. Changes on melting into its auhydride.

O. Anhydride C₆II₄BrCO.[106°]. From melilotic anhydride and Br in CS, in the cold (Fittig a. Hochsletter, A. 226, 361). Thick prisms (from ehloroform). Bromine is not taken out by boiling alkalis. Sol. alcohol and chloroform, sl. sol. CS. Slowly converted by boiling water into bromo-melulotic acid.

a-Bromo. β-oxy-β-phenyl-propionio acid C.H. CH(OH).CHBr.CO.H. [122°]; [125°, anhydrous]. From aβ-di-bromo-β-phenyl-propionic acid by boiling with water (Glaser, A. 147, 81). Thin lamine (containing aq). Boiled with very dilute Na₂CO₃ it gives phenyl-acetic aldehyde:

Ph.CH(OII).CHBr.CO₂H =

Ph.CH . CH.CO2H + HBr =

Ph.CH.CH(OH) + IIBr =o.do

 $Ph.CH:CH.OH + CO_2 + HBr =$ Ph.CH..CHO + CO, 4 HBr. The yield is 75 p.c. of the theoretical, but some phenyl-glyceric acid is also formed:

 $Ph.CH \cdot CH.CO_{2}H + H_{2}O =$

Ph.CH(OH).CH(OH).CO,H (Erlenmeyer, B. 13, 308). Salt. - AgA'.

ββ-Bremo-a-oxy-α-phenyl-propionic acid CHBr₂.CPh(OH).CO₂H. Di-bromo-atrolactic acid. [167°]. Propared by dissolving di-bromo. pyruvio acid and benzeno in cold H.SO. Long needles or four-sided tables. Sol. benzene and CS., sl. sol. cold water. By boiling with water it decomposes into CO., IIBr, and ω bromacetophenone (C.II., CO.CH.Br). On reduction it gives atrolactic acid (Böttinger, B. 14, 1285).

Bromo-di-oxy-phenyl-propionic acid Melhylene other Coll, Bro, or

 $CH_2 < 0 > C_0 H_2 Br. CH_2 . CH_2 . CO_1 H.$

pipero-propionie acid. [140°]. From sodium bromo-(β)-hydro-piperate and KMnO₁ (Wein-stein, A. 227, 44). Monoclinie crystals (from cther) sol. alcohol, sl. sol. water .- CaA'

Di-bromo-o-oxy-phenyl-propionic acid C.H.Br.O. Di-bromo-melilotic acid. [115°]. From melilotic acid and Br (Zwenger, A. Suppl. 5, 116). Needles; may be distilled = BaA', 5aq α-β-Di-bromo-o-oxy-phenyl-propionic acid.

Di-bromide of coumaric acid. Methyl derivative

C₆H₄(OMe).CHBr.CHBr.CO...H. f162° S. (CHCl.) 2.7 at 17°. From the methyl derivative of commaric acid C.II, (OMe) CH CH.CO.II and Br. V. sol ether. Decomposed by aqueous alkalis. With bromine vapour it gives rise to C.H.Br. (OMc) CHBr. CHBr. CO.H [c, 202]. Cryslals (from benzene) (Perkin, C. J. 39, 420; Fittig a. Ebert, A. 216, 157). Strong potash (I:I) forms C, II, (OMe)C, HBr, CO, H [171].

Di methyl ether C.H.(OMe)CHBr.CHBr.CO.Me.

(a) compound [125]. S. (CS.) 34. (β) compound [68]. S. (CS.) 4.

These two compounds are formed together by acting on the isomeric methoxy-phenyl-acrylatesof methyl with bromine in CS. But the (v). isomeride gives chiefly that melting at 125° while the (β) isomeride forms chiefly the other (Perkin C. J. 39, 424). Alcoholic potash converts both into methoxy-phenyl-bromo-acrylic acid.

Ethyl derivative

С. H. (ОЕЙ). СПВт. СПВт. СО. П. [155-]. S. (СS.) 103 at 18°. From the ethyl derivatives of coumarie and of coumarinic acids by Br (F. a. E.). Small crystals (from CS2)

Di-ethyl ether C₆H₄(OEt)G: (Br.CHBr.CO.Et. [78°]. F₁ C₆H₄(OEt)G: (CH.CO.Et and Br in CS. (P.). From

eso-Di-bromo-p-oxy-B phenyl-propionic acid HO.C.H.Br. CH. CH. CO.H. Distromo-hydro-p-countaric acid. [108°]. From aqueous hydrop-coumaric acid and cold bromine-water (Stöhr. A. 225, 64). Needles (from acetic acid).

Salts. - (NII,)2C. H. Br.O. - Ag C. H. Br.O. aβ-Di-bromo-p- xxy-phenyl propionic acid C.H.(OH).CHBr.CHbr.CO.H. p - Coumaricacid-di-bromide.

Methyl derivative

C₆H₄(OMe).CHBr.CHBr.CO₂H: [149]; colourless crystals. Formed by combination of the methyl derivative of p-commaric acid with Br.

Di-methyl elher C_eH₄(OMe).CHBr.CHBr.CO₂Mc: [118°]; m. sol, ether and eliloroform. Formed by combination of the di-methyl ether of p-commaricacid with bromine. When boiled with aqueous potash solution (30 p \clubsuit .) it is converted into the methyl derivative of ω -bromo p-vinyl-pheno C.H.(OMe).CH:CHBr (Valentini, G. 16, 424 Eigel, B. 20, 2536).

Di-bromo-di-eso-oxy-aa-di-pheayl-propionic acid C₁,H₁₂Br₂O₄. Di-bromo-di-phenopropionic acid. Formed by bromination of di-phono-propionic acid CII₃.C(C₆II₄OH)₂.CO₂H. Amorphous powder. Sol. alcohol, insol. water.

Di-aectyt derivative C₁₃H₁₆Br₂(OAe)₂O₂; insoluble light yellow powder (Böttinger, B. 16,

Tri-bromo-p-oxy-phonyl-propionic acid C.H.Br(OH).CHBr.CHBr.CO.H. Bromo - pcommaric acid-di-bromide. [188°]. Obtained by the action of bromine upon p-commario acid. Needles. By alcoholic KOII it is converted into tri - bromo - oxy - ethyl - benzene C, H3Br(OII).CHBr.CII,Br.

Methyt derivative C_aH_aBr(OMe).CHBr.CHBr.CO₂H : needles. Formed by the action of bromine upon the methyl derivative of p-coumaric acid C.H. (OMc).CII:CH.CO.H. By heating with aqueous KOII (30 p.c.) it is converted into bromomethoxy-phenyl-acetylene C.H.Br(OMe).C:CII (Eigel, B. 20, 2534).

o-a-B-tri-bromo-o-oxy-phenyl-propionic acid Methylderivative·C_aH_aBr(OMe)CHBr.CHBr.CO₂H. [185°-188°]. S.

(chloroform) 42. From methyl-o-coumarie acid and bromine vapour (Perkin, C. J. 39, 417). White nodules (from benzenc).

Boiled with sodium acetate it gives off CO2 forming the methyl derivative of o-exo-dibromoo-vinyl-phenol, CallaBr(OMe)CzII2Br, a viseid oil. Strong KOII (1:1) gives the methyl derivative of brome-oxy-phenyl-propiolic acid (q.v.).

Tetra-bromo-o-oxy-phenyl-propionic acid

Methyl derivative

C₈H₂Br₂(OMe).CHBr.CHBr.CO₂H. [202°]. From the preceding and Br (P.).

DI-BROMO-DI-OXY-DI-PHENW SUL-PHIDE. S(C_aH₃Br.OH)₂. [173°]. From p-bromo-phenol and SCl₂ in CS₂ (Tassinari, G. 17, 91). Amorphous, reduced by zinc-dust to S(C.H.OII)... [128°]. DI-BROMO - DI-OXY - DI-PHENYL - SUL -

Di-methyl derivative SO₂(C₆H₃(OMe)Br)₂.[166°]. From SO₂(C₆H₄.OMe) Small plates, v. sol. boiling alcohol (Annaheim, A. 172, 48).

Diethyl-derivative SO2(Calla (OEt)Br)2.

₹183°].

Di-isoamyl Arivative

SO₂(C₀H₃(OC₃H₁₁)Br)₂. [100°]. Tetra-bromo-di-oxy-di-phenyl sulphone SO2(C.H.Br2.OH)2. [279°]. From di-oxy-diphenyl-sulphono and Br. Thick monoclinic prisms (from alcohol).

TETRA-BROMO-OXY-PHENYL-VALERIC ACID. Methyl der sative

C_aH₂Br₂(OMe)CHBr.CEtBr.CO₂H. [159°]. From (a) and (β) methoxy-plienyl-angelic acids and bromine vapour. Crystallised from light petro-leum (Perkin, C. J. 39, 437).

Di-bromo-di-oxy-phenyl-valeric acid Methylene derivative C121112Br2O4 i.e. CH. CO>C. H3.C. HBr2.CO2H. Di-bromo-piperhydronic acid. [136°-140°]. From (a)-hydro-

piperic acid and Br (Fittig a. Mielck, A. 172, 159; Woinstein, A. 227, 33). Warm NaOHAq gives piperio scid. Sodium-amalgam gives hydropiperic acid.

Tetra-bromo-di-oxy-phenyl-valerio acid Methylenc derivative

CH₂<0>C₆H₂.CHBr.CHBr.CHBr.CHBr.CO₂H.

Tetra-bromo-piperhydronic acid. [160°-165°]. From piperic acid and Br (F. a. M.). Alkalis give HBr and piperonal CH₂O₂.C₅H₃.CHO. Boiling water produces IIBr and 'di-bromo-piperinide 'C₁₂H₃Br₂O₄ [136°]; this body crystallises from alcohol in prisms, insol. water and alkalis, converted into piperonal by boiling aqueous Na₂CO₃. Further treatment with water converts di-bromo-piperinide into bromo-oxy-piperinide C12H2BrOs [132°], which separates from alcohol in monoclinic crystals, insol. aqueous Na_zCO_z

BROMO-DI-OXY-PHTHALIDE, Di-methylderivative C10H9BrO4 i.e.

 $C_s\Pi_sBr(OMe)_2 < \stackrel{CO}{CII}_2 > O [x:6:5:_1^2].$

Bromo-pscudo-mcconinc. [142°]. White floceulent solid (Salomon, B. 20, 887).

BROMO-OXY-PIPERINIDE v. Tetra-bromo-DI-OXY-PHENYL-VALERIC ACID.

a-BROMO-β-OXY-PROPIONIC ACID C₃H_.BrO₃ i.e. CII₂(OH).CHBr.CO₂II. Bromo-hydracrylie acid. Formed by warming silver aβ-di-bromo-propionate with water (Beckurts a.

Otto, B. 18, 236. Syrnp; converted by moist Ag₂O into glyceric acid. Salt.—ZnA'₂.

β-Bromo-a-oxy-propionic acid CH.Br.CH(O11).CO.11. β-Bromo-lactic acid. [90°]. From exy-acrylic acid and HBr (Melikoff, B. 13, 958). Prisms (from ether); miscible with water.

Di-cthyl ether CH₂Br.CII(OEt).CO₂Et. From CH₂Br.CIIBr.CO₂Et and NaOEt (Michael, J. pa [2] 35, 136).

aβ-Di-bromo-a-oxy-propionic acid

CILBr.CBr(OII).CO.II. Di-bromo-tactie acid. [98]. From acrolein dibromide and cold diluto HNO, (Linnemann a. Peni, B. 8, 1101).

ββ-Di-bromo-α-oxy-propionic acid CHBr₂.CII(Oil).CO₂II. Di-bromo-laetic acid. From the nitrilo and HClAq. Syrup.

Nitrite CHBr., CH(OH).CN. From dibromo-aldehyde and HCN. Oil (Pinner, A. 179, 71; B. 7, 1501).

βββ-Tri-bromo oxy-propionic acid

Tri-bromo-lactie acid. CBr₃.CH(OH).CO₂H. [143°]. From bromal, HCN, and HCl (Pinner, B. 7, 1501; Wallach, A. 193, 50).

Ethyl ether EtA' [46°]; prisms. Nitrile CBr₂CH(OII).CN. From bromal hydrate and conc. HCNAq. Prisms, v. sol. water.

Tri-bromo-ethylidene ether v. Bro-

Tri-chloro-ethylidene ether CCl. GH(C3I1Br3O3)2 [134°]. Formed by heating tuo acid with chloral.

DI-BROMO-DI-OXY-PROPYL-BENZENE v. Baomo-Eugenor.

Di-bromo-tri-oxy-propyl-benzone. Di-bromo-

propyl-pyrogallol. Di-methyl derivative $C_{11}H_{14}Br_{2}O_{3}$ i.e. $C_{a}(C_{2}H_{1})Br_{2}(OH)(OMe)_{2}$. [109°]. Acetyl derivative $C_{a}(C_{2}H_{1})Br_{2}(OAc)(OMe)_{2}$ [102°].

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Methyl di-acetyl derivative $C_6(C_3H_7)Br_2(OAc)_2(OMe)$. [79°]. These compounds are formed by brominating the corresponding derivatives of tri-oxy-propyl-benzene (Hofmann, B. 11, 331; Brezina, M. 4, 492; Pastrovich, M. 4, 185).

DI-BROMO-DI-OXY-DI-PROPYL-MALONIC ACID (CH_Br.CH(OH).CH2)2C(CO2H)2.

Di-lactone CH2Br.CII.CII2 CH_CH.CH_Br. [130°]. O.CO \co .ò

From di-allyl-malouic acid in glacial HOAc by Br (Hjelt, B. 15, 625; A. 216, 61). The tetrabromido (CH_Br.CHUr.CH_2)_C(CO_2H)_2 is first formed, but splits off 2HBr. Small plates (from alcohol). Insol. cold water, sl. sol. boiling water, v. sol. warm alcohol, sl. sol. ether. When boiled with baryta it ought to form Ba(CO₂)₂C(CH₂.CH(OH).CH₂OH)₂ but this splits off BaCO3 forming the lactonic acid:

CH2OH.CH.CH CH.CH, CH(OH).CH,OH. Ò--CO

DI-BROMO-OXY-PYRIDINE C. HalbraNO i.e. C_sH₂Br₂(OII)N. Prepared by heating piperidino with Br and water to 200° (Holmann, B. 12, 981). Glistening scales. Sl. sol. water, other, and Sol. aqueous acids and alkalis.-(B'HCl)₂PtCl₄: long needles.—C₅H₂AgBr₂ON: white pp.

Methyl derivative: [193°]; long needles. Di - bromo - oxy - pyridine C,II,Br,(OH)N. [207°]. Long white needles. Formed by adding bromine-water to a solution of oxy-pyridino [107°] (Königs a. Geigy, B. 17, 591).

OGH_Br2(OH)N. Di-breme-exy-pyridine [c. 200°]. Formed by the action of brominewater upon (β)-oxy-pyridine [125°]. Colourless needles. V. sol. water and alcohol, nearly insol. benzenc. Fe₂Cl₃ gives a violet colouration.

Salts.—B'HBr: small white silky n edles. $B'_2H_2SO_4$: easily soluble plates. $-B'_1H_2C_2O_4$: needles sl. sol. alcohol. The pierate forms yellow needles (Fischer a. Renout, B. 17, 1898).

BROMO - OXY - PYRIDINE - CARBOXYLIC ACID C. H. N(Br)(OH)CO. H[1:x:2:5]. Bromo-oxynicotinic acid. [296]. Obtained by saponification of the methyl-ether, which is formed by tho action of aqueous NII, upon the methyl-ether of bromo-cumalic acid. Sl. sol. hot water, nearly insol. ether, alcohol, and acetic acid.

Methyl ether C,II,N(Br)(OII)CO.Me. [222°]. Slender glistening needles. Sol. bot water and hot alcohol.

Phenyl derivative of the methyl-ether C.H., N(Br)(OPh)CO, Mc. [183°]. Formed by the action of aniline on the methyl-ether of camalic acid in alcoholic solution. Distils without de-composition; white glistening needles, sol. alcohol and other, insol. water (Pechmann a. Welsh, B. 17, 2398).

DI-BROMO OXY QUINOLINE C_pH₄Br₂(OH)N. [195°]. Prepared by the action of bromine-water on a solution of oxy-quinoline (Bedall a. Fischer, B. 14, 1367). Wh. te silky needles. Sol. alcohol, ether, benzene, CS., nsol. water, ligroïn, and dilute acids.

V. also Bromo-carbostyril. BROMO - (B. 4) - OXY - QUINOLINE TETRA-HYDRIDE. Ethyl ether C₉H₂Br₂(OEt)N.

[45]. Obtained by adding bromine to a cooled chloroform solution of (B.4)-ethoxy-tetra-hydroquinoline. Long tricline crystals. The hydrochlorido crystallises in felted noedles, the sulphate in colourless plates, and the exalate in prisms. The pierate forms sparingly soluble yellow needles [108]. The nitrosamine forms glistening plates [86] (Fischer a. Renonf, B.

C_aBr₂(ÔH)₂O₂ [1:4:2:5:3:6]. Bromanilic acid. Formation.—1. By dissolving di-, tri-, or tetra-bromo-quinone in potash (Stenhouse, A. 91, 311; Sarauw, A. 209, 115). - 2. By heating the sodium salt of di-oxy-qu'none-di-p-earboxylic acid with cone. HBr .- 3. Together with tetra-bromo-quinono by heating (1,3,5,2)-tribrome phenol with pyrosulphuric acid at 115°; the reaction is anomalous as the two Br should remain m to one another (Salzmann, B. 20, 1997), v. also Di-chloro-di-oxy-quinone.

Properties. - Monosymmetrical needles or bronzy plates. Converted by Br into hexa-bromo-acctone. A nentral solution of the Na salt gives the following reactions :- CaCl2: brown pp.—BaCl₂: yellowish-brown pp.—FeSO₄ and NiSO4: greenish-groy pp. -Fe Cla: brownish-black. -Co(NO3)2: brown. -I'b(OAc)2: reddish-brown.— CuSO; greenish-brown.—AgNO, and Hg₂(NO₃)₂; red. HgCl₂: no pp.

Salts. - Na2A" 4aq: asymmetric crystals. -K.A" 2nq: asymmetric crystals. - KA"aq (Hantzsch, B. 20, 1303; Hantzsch a. Schniter, B. 20, 2040, 2279).

Di-breme-di-exy-quinene. Di-methyl derivative C6Br2(OMe)2O2. [175] (Hofmaun, B. 11, 332).

Tri-brome exy-quinene C₆(OII)Br₂O₂ [207°]. From oxy-hydroquinouo and Br (Be th a. Schreder, M. 5, 593). Orange grains; sol. alcoaol and CHCl.

DI BROMO OXY-TOLUIC ACID. Methul ether C. HBr Mc(OMe).CO. H [?:?:4:2:1]. [194°]. From the methyl derivative of di-bromo-thymol by oxidation (Paternò a. Canzoneri, G. 10, 233). DI-BROMO-OXY-TOLUQUINONE

Formed in small C₆McBr₂(OII)O₂. $[197^{\circ}].$ quantity by the action of dilute KOH on tribromo-toluquinone (Spica a. Magnanimi, G. 13,

BROMO . OXY. TOLYL. PHENYL . KETONE. CARBOXYLIC ACID

 $C_{_{0}}\Pi_{_{4}}(\overline{CO}_{_{2}}\Pi).CO.C_{_{0}}\Pi_{_{2}}(CH_{_{3}})(\operatorname{Pr})OH.~~[228^{\circ}].~~Pre$ pared by the action of Br and acetic acid on an alcoholic solution of o eres d-phthalein. Small prisms. By heating with H.SO, to 130° it is readily converted into bromo-oxy-methyl-anthraquinone. Chloride (208') (Fraude, B. 12, 239).
DI-β-BROMO & OXY-α-TOLYL-PROPIONIC

ACID C₁₀H₁₀Br₂C₃ i.e. CHBr₂C(C₇H₃)(OH).CO₂H. Di-bromo-cso-macingl-atrolactic acid. [103°]. Prisms or needles. Prepared by dissolving di-bromo-pyruvic acid and toluene in II₂SO₄at 0°. By hot water it is decomposed into CO2 and tolyl bromo-methyl ketone C,II,.CO.CH,Br. On reduction it gives eso-methyl-atrolactic acid (Böttinger, B. 14, 1597)

BROMO-OXY-VALERIC ACID. Lactone. From βγ-di-bromo-CH, CH.CHBr.CH, CO.O.

valeric acid by boiling with water (Messer-

schmidt, A. 208, 102) Non-volatile oil; converted by boiling baryta-water into di-oxyvalerio acid.

Di-bromo-oxy-valerio acid.

CH, CBr.CHBr.CH_.CO.O. [78°-81°]. (a)-angelico-lactone and bromine. Thick white

CH.Br.CBr.CH..CH..CO.O

hygroscopic needles (from CS₂). Water converts it into 11Br and bromo legulic acid (Wolff, A. 229, 264).

Lactone

DI.BROMO-PALMITIC ACID C₁₆H₂₀Br₂O₂. [29]. From hypogwie acid and Br (Schröder, A. 143, 24). Amorphous and insol. water. Alcoholic KOH converts it into bromo-hypogwic and palmitolic acids. Aqueous alkalis form dioxy palmitic acid.

Di.bromo-palmitic acid C161130Br.O2. From garolic acid and Br. Crystallino; converted by alcoholic KOH into palmitolic acid (S.).

Tri-brome-palmitic acid C₁₆H_{.6}Br₈O₂. [39°]. From brome-hypegoic acid and Br. Amorphons (S.).

Tetra.bromo-palmitic acid. C, II, Br, O2.

Yellow crystals.

BROMO-PALMITOLIC ACID Ct. H., BrO, [31°]. From tri-bromo-palmitic acid (v. sup.) and alcoholic KOH (S.).

BROMO PENTANE v. AMYL BROMIDE.

αβ. Di-bromo-pentane C₃11₁₀Br₂ i.e. CH₃. CH₄.CHBr.CHBr.CH₃. Anylene bromide. (178). S.G. 1 1 6868. From the corresponding amylene and Br (Wagner a. Saytzeff, A. 179,

ωω. Di-bromo.isopentano Pr.C1L.CHBr. Isoamylidene bromide. (170°-180°). From isovaleric aldehydc and PCl₂Br. (Bruylants, B. 8, 406). Alcoholic KO11 gives PrC11:C1113r (1110) and Pr.C:C11.

aa.Di.bromo-pentano Pr.CBr., CH., From methyl propyl ketone and PCl.Br., (R.). Split up by distillation into HBr and Pr.CBr.CH. (123°).

aß. Di-bromo. isopentane (C11_a), CBr.CHBr.CH_a. (170°-175°). S.G. ¹/₁; 1·6370. M.M. 12·947 at 12·6°. From tri-From trimethyl-othylene and Br (Wurtz, A. Ch. [3] 35, 458; Bauer, Dl. 2, 149). Converted by water (20 vols.) and PbO at 150° into methyl isopropyl ketone (Elickoff, J. R. 10, 215).

Isoamylene dibromide, formed by combination of Br with isoamylene from crude fusel oil, is a mixture of several of the preceding dibromo pentanes (Caliours, C. R. 31, 291; Wurtz, A. Ch. [3] 4, 458; A. 123, 202; Reboul, C. R. 58, 1058; A. 133, 84; Baner, Bl. 1860, I48; A. 120, 167; Z. 1861, 590; Golowkinsky, A. III, 252; Olevinsky, Z. 1861, 671).

Valerylene dihydrobromide (170'-175'), formed by the union of HBr with crude valerylene is also a mixture of di-bromo-pentanes.

Tri-bromo-pentane C311, Br3. I'rom bromoisoamylene and Br.

Tetra bromo pentane C. II. Br. Valerylene tetrabromide. [-10°]. From crude valerylene and Br (Reboul, A. 132, 119; 135, 372).

Tetra bromo pentane C.H.Br. Piperylene tetrabronide. [115]. From piperyleno and Br (Hofmann, B. 14, 664). A liquid isomeride is also formed (Magnanimi, G. 16, 390).

Tetra - bromo - pentane C.H.Br. Isoprana tetra bromide. From isoprene (Tilden, C. N. 46, 120).

Penta bromo-pentane C,H,Br,. Two bodies of this composition are formed by the action of

Br on valcrylene in sunlight (R.).

BROMO - PENTENYL ALCOHOL ether C,H,BrO i.e. C,H,Br.OEt. (1770-1800). S.G. 19 1.23. From tri-bromo-pentano and alcoholic KOII (Reboul, A. 133, 84).

BROMO-PENTINENE v. BROMO-VALERYLENE. BROMO-PHENANTHRAQUINONE v. PHEN-ANTHRAQUINONE.

BROMO - PHENANTHRENE v. PHENAN-THRENE.

BROMO-DI-PHENIC ACID

CO_11.C_H_4.C_11_Br.CO_11. [236° uncor.]. Formed, together with its di-bromide, by heating diphenic acid with bromine at 80°-100°. Small white prisms. Sublimes with difficulty. Not volatile with steam. V. sol, alcohol, other, and acctic acid, sl. sol. benzene, chloroform and CS., insol. cold water.

Salts .- A"Na,: white v. sol. amorphous powder.--A"Ba 3aq: sparingly soluble needles.--Λ"Ag₂: white insol. pp. -A"Cu: sl. sol. amorphous green powder.

Di-ethyl ether A"Et,: [65°]; crystals

(Claus a. Erler, B. 19, 3149).

Bromo-di-phonic-acid-di-bromide

C₁₂H,Br₄(CO.11)... [256° uncor.]. Formed in small quantity (15 p.c.) by heating diphenic acid (1 mol.) with bromino (2 mols.) for 8 days at 100°. Glistening colourless needles (from hot alcohol). Sl. sol. ordinary solvents. Its alkaline solution decomposes very easily on heating, forming salts of di-bromo-di-phenie acid. It has a very bitter peculiar taste. - A"Na: solublo silky plates (Claus a. Edler, B. 19, 3152).

Bromo-di-phenic acid [4:1] \mathbf{C}_{0} 11₄(COOH). \mathbf{C}_{0} 11₃Br,COOH [1:4:2]. [208°]. Formed by exidation of the liquid (1,1,1',4',2') mono-brome-ditelyl with CrO, and acetic acid (Carnelley a. Thomson, C. J. 47, 590).

Di-bromo-di-phenic acid C₁₁U_sBr₂O₄, [296°]. From di-bromo-phenanthraquinone, KrCr2O-, and H.SO. Geodes of small crystals, v. sl. sol. hot water, insol. alcohol and other (Ostermayer, B.

7, 1091). Di-bromo-di-phenic acid C12HaBra(CO2H)... [245° uncor.]. Formed by heating di-plication acid (1 mol.) with bromine (2 mols.) at 200°, or by heating aqueous solutions of salts of monobromo-di-phenic-acid-di-bromide. Small glistening needles. Not volatile with steam. Sublimes with difficulty. Sl. sol. benzene, chloroform, acetone, CS,, and hot water, nearly insol. cold water, v. sol. alcohol, ether, and acetic acid.

Salts .- The alkaline salts are very soluble amorphous glassy masses. — A"Ag: white amorphous pp. —A"Ca 3aq: easily soluble plates. -A"Ph: sl. sol. microcrystalline powder.
Diethyl ether A"Et.: [106" uncor.]

crystals (Claus a. Edler, B. 19, 3149).

o.BROMO-PHENOL C. H. BrO i.e. C. H. Br(OH) [1:2]. (195°). From o-bromo-anilino by the diazo reaction (Fittig a. Mager, B. 8, 362; Körner). Formed in small quantity in brominating phonol (Hübner a. Brenken, B. 6, 171). Oil; volatile with steam. Potash-fusion gives

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resorcin. HNO, forms bromo-di-nitro-phenol

Methyl ether C.H.Br.OMe. (223°). From the methyl ether of o-amido-phenol by Sandmeyer's reaction (Wallach a. Hensler, A. 213, 238).

m-Bromo-phenol C.H.Br(OH) [1:3]. (236°). From m-bromo-aniline by the diazoreaction (Körner, G. 4, 389; Wurster a. Nölting, B. 7, 905; F. a. M.). Crystalline. Potash-fusion gives resorcin. IINO3 gives bromo-dinitro-phenol [92°]

p.Bromo-phonol C.H.Br(OH) [1:4]. [64°]. (237°). S.H. (18°-77°) 3157. S. I-422 at 15°.

Formation.- 1. By distilling bromo-o-oxybenzoio acid with BaCO₃ (Calcours, A. Ch. [3] 13, 102) .- 2. By passing air saturated with Br (160 pts.) into cooled phenol (94 pts.) (Körner, A. 137, 197).—3. Bromino (160g.) is dissolved in glacial HOAc (200 g.) and added to phenol (94 g.) dissolved in HOAc (300 g.) (Hübner a. Brenken, B. 6, 171). -1. From p-bromo anilino by tho diazo- reaction (K.; F. a. M.).

Properties.—Large crystals (from chloroform); sl. sol. water, v. sol. alcohol. Dimetric; a:c=1:146. Its heat of neutralisation has been determined by Werner (C. R. 98, 1333; Bl. [2] 46, 281). Nitration gives bromo-di-nitrophenol [76°]. Potash-fusion gives resorein. SCl2 forms S(C, H, Br.OH), [176°] (Tassinari, G. 17, 83).

Methyl ether Call Br.OMe. Bromo-anisol. (223° cor.). S.G. 2 1.191.

Ethyt ether C.H. Br.OEt. (233°).

Isopropyl ether C. H.Br.OPr. (236°). S.G. 2 1 981. μ₀ 1 553. From isopropyl phenol and Br (Silva, Bt. [2] 13, 27).

Benzoyl derivative C. II. Br. OBz. Crys-

Bromo-phenol (Fourth). (236°-2383) (Fittica, J. pr. [2] 28, 176; B. 19, 2632; A. Ch. [6] 4, 561).

Preparation.—Phenol (10 g.), alcohol (10 g.) and amorphous phosphorus (3 g.) are mixed and cooled while bromine (17 g.) is added through a eapillary tube. The product is washed with dilute Na, CO, dried and distilled. It contains di-bromo-phenol and the new body. This can only be distilled when in small quantities, in larger masses it undergoes carbonisation.

Properties. -- Not solid at 10'.

Nitration. -- Bromophenol (1 pt.), glacial acetio acid (3 pts.), and HNO₃ (S.G. 1-1) added gradually form crystals of a molecular compound $C_6H_8Br(NO_2)OH.C_6H_4Br(NO_2)$.OH which crystals lises from alcohol and melts at $[60^{\circ}, 65^{\circ}]$. Fuming HNO₃ converts this into a bromo-dinitro-phenol [1089-1107], isomeric with those known. By the action of baryta on the above molecular compound a second such body (C₄H₃Br(NO₂)OII)₂C₄H₂Br(NO₂)₂OII is got.

The existence of four bromo-phenois would

be contrary to the general rule that only three isomeric di-derivatives of benzenc can be obtained; according to Hand (A. 231, 129) the fourth brome-phenol is merely p-brome-phenol of which the melting point is lowered by a trace of moisture.

Di-bromo phenol $C_0H_4Br_2(OH)$ [1:3:4]. [40°]. S.H. (18°-73°) ·2436. S. ·194 at 15 '(W.).

Formation .- 1. By distilling di-bromo saliorlio acid with baryta (Calcours, A. 52, 329), or ether, nearly insol. water (Daccomo, B. 18, 1168).

by heating with dilute H.SO, at 280° (Peratoner, G. 16. 402) .- 2. By passing bromine vapour (2 mols.) into cold phenol (I mol.) (Körner, A. 137,

Properties. - Crystalline mass, v. sl. sol. water, v. sol. ordinary solvents. HNO3 forms pieric acid. Its heat of neutralisation has been determined by Werner (C. R. 98, 1333). Heated with dilute H.SO, in scaled tubes, it is converted into mono- and, tri-bromo-phenol (Peratoner, G. 16, 403).

Methyt ether C. II, Br. (OMc). [59°]. (272°). From di-bromo phenol, Nathl, and McI. Formed also by brominating anisol (C.).

Nitro-benzoyt derivative C_aH₃Br_aO.CO C_aH₄NO₂ [90] [100]. From ben-zoyl-phenol by bromination followed by ni-

Di-bromo-phenol Call Br (Oll) [6:2:1]. [56°]. Formed by distilling tetra-bromo-phenol-phtha-Icin with conc. H.SO, (Uneyer, A. 202, 138). Also from di-bromo-p-amido-phenol by displacing NH., by H (Möhlau, B. 15, 2191). Mass of thin needles (from water).

Di - bromo - phenol C. H. Br. (OH) [1:3:5]. [76.5"]. Formed, together with its methyl ether, by heating s-tri-bromo-benzene with NaOMe (Blau, M. 7, 621). Converted by potash-fusion into phloroglucin.

Methyt ether C.H.Br. (OMe): [382]

Ethylether C. H. Br (OEt). (268). Formed by boiling di-bromo-o-di-azo-phenetol nitrato, Call Bra(OEt) NaNOa, with water (Möhlau a. Oelminhen, J. pr. 132, 482).

Tri - brome - phenol | CaH Br. (OH) [2:4:6:1] [92] (Wilsing, A. 215, 235); [95] (Korner). S. 007 at 15 (W.).

Formation -1. From phenol and Br Taurent, A. Ch. [3] 3, 211; Körner, A. 137, 298). . . . 2. By distilling tri-bromo-salicylic acid with sand and baryta (C.) .- 3. By treating indigo with bromine-wate .- 1. From potassium phenol disulphorate and Br (Schmidt, B. 11, 852).

Properties. - Long silky needles (from dilute alcohol); may be sublimed. V. sl. sol. water, v. sol. alcohol. Its heat of neutralisation has been examined by Werner and Berthelot (C. R. 98,

1333; A. Ch. [6] 3, 552).

Reactions .- 1. Nitric acid forms di-bromonitro, bromo-di-nitro, and tri-nitro-phenol (Armstrong a. Harrow, C. J. 29, 476). 2. Cro. and HOAs give tetra-bromo-quinone and amorphous insolable C. H. Br. O. (Benedikt, A. 199, 134).-3. Bromine-water forms Call Br.OBr (?) [118] which forms yellow flates, insol. water, ether, and alcohol; it ext note the following reactions: (a) At 120° it splits up into Br and the compound $C_{ij}H_i\text{Br }O_{ij}$ (b) Aniline forms tribromo-phenol and to bromo aniline. (c) Phenol forms tri-bromo-pie col. (d) It is also reduced to tri-bromo-phenol by hoiling alcohol or by Sn and HCl (Benedikt, B. 12, 1005; M. 1, 360; Werner, Bt. [2] 43, 373).

Ethyt ether C.H.Br. OEt. [69]; prisms (Purgotti, G. 16, 526).

Propionyt derivative C.W.O.C.H.Br. [65°] (Guareschi a. Daccomo, B. 18, 1174).

Benzoul derivative C.H.Br.OBz: [82] small colourless prisms; soluble in alcohol and

Tri-bromo-phenol

. Ethyl ether C. H.Br. OEt. [73°]. Prepared from tri-bromo-di-azo phenetol by boiling with water and extracting the product with other (Möhlau a. Ochmichen, J. pr. [2] 24, 484).

Tetra-brome-phenol C_aHBr_aOH [2:3:4:6:1]. [120°]. From tri-brome-phenol and Br at 480° (K.). Formed also by warming C_aH_aBr_aOBr (v.) sup.) with conc. H₂SO₄. Needles (from alcohol); may be sublimed. Bromino converts it into CaHBr4OBr [121°] which separates from chloroform in monoclinic crystals and is reduced by boiling alcohol or by Sn and HCl to tetra-bromo-

phenol (Benedikt. M. 1, 361).

Penta-bromo phenol C. Br. OII. [225°]. Obtained by heating tri- or tetra- bromo-phenol with excess of Br at 220° for some days (Körner, A. 137, 210). Formed also by heating Call Br. OBr with cone. H.SO, (Benedikt, M. I, 360)! Adamantine needles (from CS,); may be sublimed. Conc. HNO, forms bromopicrin and tetra-bromoquinone. PBr, forms CaBr, (Gessner, B. 9, 1505; Ruoff, B. 10, 1234).

Hexa-bromo phonol C.Br.OBr. [128°]. Prepared by heating tri-bromo-phenol with excess of bromine in scaled tubes at 220°. Yellow crystals of trimetric system (a:b:c=I: 82: 114); insol. cold alcohol, but decomposed by boiling into penta-bromo-phenol. On heating with anilino it yields penta-bromo-phenol and tri-bromoaniline (Benedikt, M. I, 363)

TETRA-BROMO-p-DIPHENOL TETRA-BROMO-DI-OXY-DIPHENYL

TETRA-BROMO-PHENOL-PHTHALEÏN

PHENOL-PHTHALKIN.

BROMO-PHENOL o-SULPHONIC ACID CeH3Br(OH).SO3II. The K salt is formed by adding Br (I mol.) to an aqueous solution of potassium phenol o-sulphonate. The free acid is very deliquescent.—KA': pointed needles—

Bromo-phenol p-sulphonic acid ag. in the preceding case (S.), or by passing bromine vapour into a cold cone, solution of phenol p-sulphonie acid (Lo Canu, C. R. 103, 385). Deliquescent needles (containing 2nq).-KA'.

Bromo-phenol sulphonic acid.

Ethyl derivative Callabrio (OFt) (SOall). From potassium phenetol sulphonate and Br (Lippmann, J. 1870, 739). Deliquescent mass (containing 4aq).— KA'.

Bromo-phenol disulphonic acid a

 $\mathbf{C}_{\mathbf{s}}\mathbf{H}_{2}\mathrm{Br}(\mathrm{OH})(\mathrm{SO}_{3}\mathbf{H})_{2}$ [2:1:4:6]. From an aqueous solution of potassium phenol disniphonic acid (I mol.) by adding Br (I mol.) (Armstrong, C. J. 25, 865; Schmidt, B. 11, 852). Crystallino; v. sol. alcohol, m. sol. other. (Fc.Cl., gives a red eolour. IINO3 forms bromo-nitro-phenol sulphonio acid, bromo di-nitro-phonol, and tri-nitro-phenol. — BaA" 2aq. — K₂A". — PbA". — Ag_2A'' .

Di-bromo-phenol o-sulphonic acid $C_0H_2Br_2(OH)(SO_3H)$ [2:4:I:b]. [120°]. Formed

by brominating potassium phenol o-sulphonate or bromo-phenol o-sulphonic acid (Armstrong, C. J. 25, 867; Senhofer, A. 156, 110; Schmidt, B. 11, 855). Concentrio needles; deliquescent. The aqueous solution is coloured violet by Fe₂Cl₈ and is ppd. by Pb(OAc)2.—HNO2 forms dibtomo-o-nitro-phenol[117°]. Ba(CoH2Br2SO4)2.— BaC₀H₂Br₂SO₄. — CdC₀H₂Br₂SO₄ 1½aq. — KC₀H₁Br₂SO₄.—K₁C₀H₂Br₂SO₄.—PbC₀H₂Br₂SO₄. Di-bromo-phenol p-sulphonic acid

C_sH_sbr₂(OH)(SO_sH) [2:6:1:4]. From potassium phenol p-sulphonate (I mol.) and Br (2 mols.) (Schhofor, A. 156, 103; Armstrong a. Brown, C. J. 25, 857); or by passing bromine-vapour into an aqueous solution of phenol p-sulphonic acid (Lo Canu, C. R. 103, 385). Formed also by diazo- reaction from di-bromo-sulphanilic acid (Schmitt, A. 120, 161). Rectangular scales (containing aq). The solution is coloured violet by Fe₂Cl₆ but is not ppd. by Pb(OAc)...

KC_II_Br_SO_aq.—K_C_II_Br_SO_2aq.— Ba(C_II_Br_SO_)_2aq.—BaC_II_Br_SO_4aq. BROMO-PHENOXY-ACETYC ACID v. Bromophenyl derivative of GLYCOLLIC ACID.

BROMO - PHENOXY - PROPIONIC ACID v.

Bromo-phenyl derivative of LACTIC ACID. o-BROMÖ-DIPHENYL C₁₂II,Br i.e.

CaHs.CaHaBr [1:2]. (258°). Formed by decomposing o-diazo diphenyl perbromide with alcohol (Schultz, Schmidt, a. Strasser, A. 207, 353). Oil, smolling of oranges; oxidised by CrO, to o-bromobenzoie acid.

p-Bromo-diphenyl C₀H₅.C₈H₄Br [1:4], [89°] (310° i. V.). Formed by adding Br to a solution of diphenyl in CS2. Laminæ; v. sol, hot alcohol and HOAc, v. e. sol. ether. Smells like oranges. Chromic acid exidises it to p-bromo-benzoic acid

(Schultz, A. 174, 207).

pp-Di-bromo-diphenyl [4:I] C_II_Br.C_II_Br [1:1]. [164°] (G.; F.); [162°] (C. a. T.). (c. 358°). Formed by heating diphenyl dissolved in CS. with bromine at 100° (Fittig, A. 132, 204; Carnelley a. Thomson, C. J. 47, 588). Also from benzidino by the diazo reaction (Griess, Pr. 13, 383). Prisms and octahedra (C. a. T.); v. sl. sol. hot alcohol, slightly volatile with steam. Oxidised by CrO, to Smells like oranges. p-bromo-benzoie acid.

Tri - bromo - diphenyl [4:1] C, H, Br. C, H, Br. Formed by the action of bromine on a mixture of diphenyl and p-tolyl-henzene (Carnelley a. Thomson, C. J. 47, 587). Colourless silky needles, sl. sol. alcohol, not volatile with steam. CrO3 in HOAc gives p-bromo-benzoic acid.

DI-BROMO-DI-PHENYL-ACETAMIDINE C14H12Br2N2 i.e. CH3.C(NHC,H1Br):NC,H4Br. From p-bromo-aniline, HOAc, and PCl₂ (Dennstedt, B. 13, 233).—B'HCl.—B',H,PtCl...
o-BROMO-PHENYL-ACETIC ACID C,H,BrO₂

i.e. [2:I] C₈H₁Br.Cll₂CO₂H. [104°]. From phenyl-acetic acid, bromine and HgO. Separated from the p-compound by its more soluble barium salt (Bedson, C. J. 37, 95). Flat needles (from water). Monoclinic tablets (from glacial acetic acid). $-a:b:c=1:657:I\cdot767$; a=99°44'. KMnO. oxidises it to o-bromo-benzoic acid. - CaA'20-AgA'.

Nitrile Call Br.CH2.CN. Oil (Jackson a. White, Am. 2, 316).

m_Bromo-phenyl-acetic acid

(3:1] C.H.Br.CH.CO.H. [97°] (J. a. W.); [100°] (G.). From (3, 4, 1)-bromo-amido-phenyl-acetic acid by removing NH2 (Gabriel, B. 15, 841).

Nitrile C.H.Br.CH.CN. Oil (Jackson a. White, P. Am. A. 16, 256).

p-Bromo-phenyl-acetic acid [4:1] C,H,Br.CH,CO,H. [115°]. Prepared by treating phenyl-acctio acid with bromine and

HgO (Bedson, C. J. 37, 94) .- Long flat needlee; may be sublimed. Oxidation gives p-bromobenzoic acid.

Salts.-AgA'.-NH,A'.-BaA', - CaA',-

CuA'. Nitrile C.H.Br.CH. CN. [47°]. Prepared by boiling p-bromo-benzyl bromide with alcoholio KCN (Loring Jackson a. Lowery, B. 10, 1210; Am. 3, 248).

a-Bromo-phenyl-acetic acid C, H, CHBr.CO, H. [84°]. From mandelic acid and conc. IlBrAq at 130° (Glaser a. Radziszowski, Z. 1868, 142). Formed also by brominating phenyl-acetic acid at 150° (Radziszewski, B. 2, 208). Monoclinic orystals (from CS₂). Reduced by sodiumorystals (from CS₂). Reduced by sodium-amalgam to phenyl-acetic acid. Boiling NaOHAq gives mandelio acid C_aH_a.CH(OΠ).CO_aH. Alcoholio KOH gives C_aH_a.CH(OH).CO_aH. Tho ethyl ether is converted by alcoholic KCy into CO2Et.CHPh.CHPh.CO2Et, and by Na into CO2Et.CPh:CPh.CO,Et.

Nitrile C.H., CHBr.CN. Is the chief product of the action of bromine on benzyl cyanide at 120°. Sol. alcohol and ether. On heating to 170° or with alcoholic KCN it gives di-cyanostilbeno. With an excess of KCN di-cyano-dibenzyl is also formed. On boiling with alcoholic KOH it gives stilbeno-di-carboxylic acid (Reimer,

B. 14, 1797).

Hydrobromide of the Nitrile

C.H. CHBr.CBr:NII. Phenyl-bromo-acctimidobromide. Formed together with the nitrile by the action of bromine on benzyl evanide at 120' the yield being 15 p.c. Colourless crystals. Bitter taste. Its vapour attack: the eyes. V. sl. sol. all solvents, most easile in boiling acetic acid. Decomposed by was r and alcohol,-HCl at 150° converts it into a womo-phenylacetic acid (?) and NH₂. Dilme HCl at 150° converts it into NH₃, mandelic acid (C_oH₂, CH(OH).CO₂H), and HBr (Reimer, B. 14, 1797).

a-Breme-di-phenyl-acetic acid

(C₆H₃)₂CBr.CO₂H. From diphenyl-acetic acid and Br at 150°. Converted by boiling water into benzilic acid CPh₂(OH).CO₂H (Symons a. Zincke ,A. 171, 131).

Di-bromo-phenyl-acetic acid

C.H.Br., CH., CO.H. [115°]. Formed by action of sunlight on mixture of o- and p- bromo-phenylacetic acids and Br. Purified by conversion into the methyl ether, distillation and saponification (Bodson, C. J. 37, 96). White needles.—AgA'.

BROMO-PHENYL-ACRYLIC ACID v. BROMO-CINNAMIO AOID and BROMO-ATROPIC ACID.

DI - BROMO - DI - PHENYL - ALLOPHANIC ACID C₁,H₁₀N,Br₂O₃ i.e. C₆H₂Br,NH.CO.N(C₆H₃Br).CO₂H.

Ethylether. [153°]. The compound which ie formed from p-bromophenyldicyanate by long boiling with alcohol is probably this ether. Fine white needles. Sol. alcohol and other. By continued action of alcohol it is converted into p-bromo-phenyl-carbamic ether (Dennstedt B. 13, 229).

p.BROMO-PHENYL-AMIDO-ACETIC ACID CH₂(NII.C₄H₄Br).CO₂H. Bromo-phenyl-gtyco-coll. [98°]. Colourless crystals. V. c. sol. aloohol, ether and hot water. Forms a lightgreen copper ealt. Prepared by the action of

obloro-acetic acid on p-oblorosniline. Vol. I.

Ethyl ether A'Et: [96°]; white needien; Incol. water, sol. sloohol and ether. p-Bromoanilide

CH₂(NH.C₆H.Br).CONH(C₆H.Br): [155°-160°]; white microscopic crystals. Sublimes at about 145°. Sl. sol. hot water, v. sel alcohol and ether (Dennstedt, B. 13, 236).

Di-breme-di-phenyl-amide-acetic acid. Nitrile C. H. Br. NH. CHPh. CN. [92°]. From di-phonyl-amido-acctic acid and Br (Tiemann a. Pielist, B. 15, 2032 Yellow rhombolicdra (from alcohol).

Tri-brome-phenyl-amido-acetic acid

CaHaBraNH.CHa.COaH. Formed by adding bromine-water to an aqueous solution of phenylamido-acetic acid. Minute needle. (from HOAc). Insol. water, acids and arkalis (Schwebel, B. 11. 1131)

BROMO-PHENYL-AMIDO-CHLORO-NAPH-THOQUINONE v. CHLORO - NAPHTHOQUINONE -DROMANTLIDE

a-BROMO-PHENYL-S-AMIDO-CROTONIC ACID CH3.C(NHPh):CBr.CO2H. Anil-bromo: aceto-acetic acid. [1380]. Formed by bromination of phenyl-d-amido-crotonic acid (anilacetacotio acid) dissolved in chloroform. Poarly plates, v. sol. hot alcohol, sl. sol. water, chloroform and other. By cold H2SO4 it is condensed to (Py. 2:1:3)-bromo-oxy-methyl-quinolino (Knorr, B. 17, 2874).

BROMO - PHENYL - AMIDO-(a)-NAPHTHO . QUINONE v. (a) NAPHTHOQUINONE BROMANILIDE.

TRI-BROMÓ-PHENYL-AMIDO-PROPIONI-TRILE C₆H₂Br₃NH.CHMe.CN. [130°]. From phenyl-α-amido-propionitrile and Br. Needles (Stephan, C. C. 1886, 470).

p-FROMO-PHENYL-(I'y.3)-AMIDO-QUINOL-

CH:CH C₆H₁ N.C(NII.C₄H₁.NO₂) [I46°]. Sil-

very scales. For ned by heating (Py. 3)-chloroquinoline with p-bromaniline (Friedländer a. Weinberg, B. I., 1533).

DI-BI OMO-DI-PHENYL-AMINE

Il N(C, II, Br), [107]. Prisms. Sol. alcohol and acctic acid. The benzoyl derivative is formed by bromination of benzoyl-di-phenyl-amine.

BzN(C₀H₄Br), Benzoytderivative [142]. Colourless plates. Solublo in alcohol and acetic acid (Lellmann, B. 15, 830).

Tetra-bromo di-phenyl-unine NII(C,H3Br2)2. [182°]. Formed by treating an alcoholic solution of da-phonyl-amino with Br.

(Hofmann, A. 132, 166; Gnellin, B. 8, 925). Acetyt derivative NAc(C, H. Br.)... [158°]. Hoxa-bromo-di-phenyl am ne NH(C, H. Br.)... [218°]. Formed, together with the preceding, by adding Br to a solution of di-phenyl-amine in HOAo (G.). Reduced by sodinm-amalgam to di-phenyl-amine.

Octo-bromo-di-puenyl-amine NH(C.HBr.)2. [c. 304°]. From di-phenyl-amine, Br, and I at 250°. Prisms (from CHCl., (Gessner, B. 9, 1511).

Deca-brome-di-phenyl-amine NII(C₀Br_s),
From di-phenyl-amine, Br, and I at 350°.
Needles (from CHCl₂). Not melted at 310°.
BROMO-PHENYL-BENZENE v. BROMO-DI-

PHENYL.

Bromo-tri-phenyl-benzene C24H1,Br. [104°]. From Br and tri-phenyl-benzene in CS2. Needles (from alcohol) (Berthold a. Engler, B. 7, II23). \mathbf{R} \mathbf{R}

BROMO - PHENYL - BENZOIC ACID

BROMO-DIPHENYL-CARTOXYLIO ACID. BROMO-PHENYL BENZYL OXIDE

C.H.Br.O.CH, C.H. [59°]. From phenyl benzyl oxids, Br, and HgO. Needles (Sintonis, A. 161, 335)

DI-p-BROMO-DI-PHENYL-BIURET

C₁₄**H**₁₁**O**₂**Br**₂**N**₃ *i.e.* (C₆**H**₄**BrNH.CO**)₂**NH.** pared by the action of alcoholio NH, on pbromo-phenyl-dicyanate (Dennstedt, B. 13, 230). Sl. sol. alcohol and other, in cl. water. Begins to sublime at about 240°.

γ-BROMO-γ-PHENYL-BUTYRIC ACID Ph.CHB..CH_x.CH_x.CO_xH. [69°]. From F'rom phsnyl-iso-crotonic acid and cone. HBr at 0'. Crystals (from CS_2) (Jayne, $A.\ 216,\ 102$). By boiling with water or treatment with aqueous Na₂CO₂ it is converted into the lactone of γ-oxy- γ -phenyl-butyrio acid (q.v.).

βy-Di-bromo-y-phsnyl-butyric acid

Ph.CHBr.CHBr.CH₂.CO₂H.[138°]. From phenyliso-crotonio acid in CS, by Br at 0° (Jayne, A. 216, 107). Crusts of small white crystals. Sodium amalgam converts it into sodium γ-oxy-γphonyl-butyrate.

exo-Di-bremo-\beta-pbenyl-isobutyrio acid C.H..CHBr.CBrMe.CO2H(?)[135°]. From phenyl-

methacrylio acid and Br (Conrad a. Hodgkinson, A. 193, 312)

p-BROMO-PHENYL-CARBAMIC ACID

Methyl other C.H.Br.NII.CO2Me. [124°]. From p-bromo-phonyl cyanate and McOII. Needles (Donnstedt, B. 13, 229).

Ethyl ether CeHBr.NH.CO.Et. Bromo-carbanilic ether. [81°] (B.); [85] (D.). From PhNH.CO.Et and bromine-water (Belirend, A. 233, 7) or from p-bromo-aniline and ClCO. Lt. Needles (from benzoline). Boiling alcoholic KOH gives K2CO, and p-bromo-aniline.

Di-bromo-phenyl-carbamic acid. her [4:2:1] C₀H₃Br₂.NH.CO₂Me. cther [97]. Formsd by brominating methyl phenyl-carbamats (Hentschsl, J. pr. [2] 34, 423). Needles (from alcohol). Warm H2SO, gives CO, and

di-bromo aniline.

Hsxa-bromo-di-phsnyl-carbamic sthsr (C_eH₂Br₂)₂N.CO₂Et.

Hexa-bromo-di-phenylamine urethane. [184°]. Formed by bromination of di-phenyl-carbamic ether dissolved in acetic acid (Hager, B. 18, 2577). Long greenish-brown nssdiss. Sol. acetic acid, nearly insol. alcohol.

p.BROMO - PHENYL - CARBAMINE CHLORIDE C.H.Br.NCl:Col. (356). From p-bromo-phsnyl oarbimide and Cl. Yellowish

liquid (Dennstedt, J. 13, 232). DI-BROMO-DI-PHENYL-CARBINOL $C_{13}H_{10}Br_{2}O$. [163°]. Di - bromo - benzhydrol. From di-phenyl-carbinol and Br. Minute needles (from aloohol). Reduced by sodium-amalgam to di-phenyl-oarbinol (Linnemann, A. 133, 6). DÍ-BRŐMO-PHENÝL CARBONATE

(C_sH_sBr₂)₂CO_s. [166°]. Silky needles; forme, by brominating phenyl carbonate (Löwenberg,

C. C. 1886, 390)

BROMO-DIPHENYL CARBOXYLIC ACID. [4:1] C₄H₄Br.C₄H₄.CO₂H [1:4]. [194°). From p-bromo-phenyl-tolueno [30°] and CrO₄ in HOAo (Carnelley a. Thomson, C. J. 51, 88). Y. sol. sther, sl. sol. alcohol.

Dl-brome-diphsnyl carboxylic acid C₁₂H, Br. CO₂H [4':1':1:20r3:4]. Di-bromo- (Claus a. Erler, B. 19, 3155).

p. phenyl-bensois acid. [204°]. From di-bromo-p-tolyl-benzene [115°] by oxidising with OrO, in HOAo. Needle-shaped prisms (from aloohol), sl. sol. alcohol (Carnellsy a. Thomson, C. J. 47,

Di-bromo-diphsnyl p-carboxylic acid

C₆H₄Br.C₆H₃Br.CO₂H [4':1':1:20r3:4]. [232°]. Formed by oxidising di-bromo-tolyl-bsnzene [150°] (C. a. T., C. J. 51, 90).

Di-breme-dipbenyl carboxylic acid [212°] has been obtained from (B)-di-bromo-fluorsns. ketone [197°] by potash-fusion (Holm, B. 16, 1081).-BaA

p-Bromo-Phenyl Cyanate

OC:N.C, H4Br [1:4]. [39°]. (226°). V. solvether. Prepared by distilling brone-phenyl-oarbamic other with P.O. (Dennstedt, B. 13, 228).

p-BROMO-PHENYL DI-CYANATE C₁₁H₈N₂O₂Br₂. [199°]. Small plates. Sl. sol. ether. Prepared by the action of a small quantity of tri-ctlyl-phosphine on bromo-phsnylcyanate heated to 100°. By long boiling with absolute alcohol it gives an acid of meltingpoint [153°] and formula C₁₆H₁₄O₃N₂Br₂ which is probably ethyl dib.omo-phenyl-allophanats (Dennstedt, B. 13, 229).

BROMO-PHENYL-CYSTEINE C.H., BrNSO. C₆H₄Br.S.CMe(NH₂).CO₂H. v-Bromo-aamido-thio-lactic acid. [181°]. Formed by boiling bromo-phenyl-mercapturic acid with conc. HCl. Needles (from dilute alcohol); v. sl. sol. water, v. sl. sol. alcohol, v. sol. diluts HCl. Boiling alkalis slowly separato p-bromo-phonyl mercaptan and form pyruvio acid. Sodiumamalgam forms NH3, lactio acid, and C6H4Br.SH. anlıydrids on warming gives anhydrele $C_6H_1Br.S.CMc < NH < CO > [153°]$

bromo-phenyl-eystoin,' but in presence of benzene it forms bromo-phenyl-mercapturio acid Call Br.S.CMc(NHAc).CO.H. Potassium oyan-

ate, torms C, H, Br. S, CMe(NII. CO.NH.) CO, H. Salts. — CuA'2. — HA'HCl (Baumann a. Preusse, H. 5, 315; B. 18, 258).

BROMO-O-PHENYLENE DIAMINE

 $C_6H_3Br(NH_2)_2$ [4:1:2]. [63°]. From (I,3,4)- or (6,3,4)-bromo-nitro-anilino, tin, and HCl (Hübner, A. 210, 359; Wurster, B. 6, 1544; Remmers, B. 7, 347). Needles; v. sol. water. Sodiumamalgam reduces it to o-phenyleno diamins. Salts .- B"HCl. - B"H SO.

Di-bromo-m-phenylene-diamins

C.H.Br.(NH.). Dark brown pp. formed by adding bromine-water to an aqueous solution of p-phenylone - diamine hydrochloride; may bs crystallised from alcohol (Hollemann, Z. 1865, 555).

BROMO-DIPHENYLENE KETONE

[104°]. From C₁₃H₂BrO i.e. | ℃C6H2Br bromo - fluorene and CrO₃ Matthews, C. J. 43, 165). (Hodgkinson a. Dark yellow needles.

Bromo - diphenylsne kstens C, H, BrO. [122° uncor.]. Formed by distilling bromo-diphenic acid with lime. Yellow plates. V. sol. benzone, ether, and hot alcohol, nearly insol. water. Sublimes readily in feltsd needlss. By distillation with zine-dust it gives fluorene

(a) Di-brome-diphenylene-ketone C18H Br2CO. [148°]. Formed by oxidation of di-bromo-fluorene [166°] with OrO, dissolved in acetic acid (Holm, B. 16, 1081). Long yellow needles. V. sol. ether and benzone.

(β)-Di-bromo-diphsnylsne ketone

C₁₂H₄Br₂CO. [198°]. From (a) di bromo-fluorene [165°] by CrO₃ in slight excess and HOAo (Hodgkinson a. Matthews, C. J. 43, 165; Holm). Yellow needles, sol. alcohol, other, and benzene. Potash-fusion gives rise to di-bromodiphenyl-carboxylic acid.

Di-bromo-diphenylens ketone C, II, Br, CO. [133° uncor.]. Formed by distilling di-bromo-di-phonic acid with lime. Thin yellow plates or long thin needles. Readily sublimable (Claus

a. Erler, B. 19, 3156).

DI - BROMO - DI - PHENYLENE KETONE OXIDE C₁₅H_eBr₂O₂ [210°-213°]. From diphenylono ketone oxido and bromine at 180° (A. G. Perkin, C. J. 43, 193). Long needles (from alcohol). Combines with bromine forming an unstable addition product.

BROMO - DIPHENYI ENE - METHANE v.

BROMO-FLUORENE.

DI-BROMO-PHENYLENE-(a)-NAPHTHYL-ENE-OXIDE C₁₆H₈Br₂O. [284°]. Yellowish white needles. Sl. sol. benzene. Prepared by bromination of phenylene-(a)-naphthylene-oxide (Arx, B. 13, 1727).

BROMO-PHENYLENE OXIDE C.H.BrO. 195°]. From phenylene oxide and Br at 100°. Needles (from alcohol) (Märker, A. 124, 250).

Di-bromo-diphenylone oxide C12H2Br2O. [185°]. From Br and diphanylene oxide in CS... Laminæ (from alcohol) (Hoifmeiste, $A.\,159,211$).

DI - BROMO - DIPHENYLENE - PHENYL . METHANE C₁₉H₁₂Br₂. [182]. From Br and the hydrocarbon in HOAe (Behr, B. 5, 971)

Tri-bromo-diphsnylene-phonyl-methane

C₁₉H₁₁Br₂ [167°-171°] (B.

BROMO-PHENYL-ETHANE v. BROMO-ETHYL-

BENZENE.

p-Bromo-di-phanyI-ethane C₁₄H₁₂Br i.e. C₅H₅CH₂CH₂C₆H₄Br [1:1]. Bromo-dibenzyl. S.G. 21:40. From s-di-phenyl-ethane, Br, and water (Fittig a. Stelling, A. 137, 266). boils above 320°.

pp-Di-bromo-di-phsnyl-ethane C14H12Br2 i.c. [4:1] C₆H₄Br.CH₂.CH₂.C₆H₄Br [1:4]. [115°]. From s-di-phenyl-ethane, water, and Br (F. a. S.). Needles (from alcohol). CrO, gives p-bromobenzoic acid.

Exo-di-bromo-s-di-phsnyl-sthans

C_eH_sCHBr.CHBr.C_eH_s. [237°] (L.); [235°] (K.). Stilbene dibromide. Di-bromo-dibenzyl.

Formation .- 1. From stilbene and Br (Linipricht a. Schwanert, A. 145, 336).-2. From dibenzyl and dry Br (Marquardt, A. 151, 364) .-3. From hydrobenzoin and PBr. (Zincke, A. 198,

Properties.—Silky needles. Decomposes at 285° (Kade, J. pr. 127, 465). V. sl. sol. boiling aloohol, m. sol. boiling xylenc. Alcoholic KOH gives C.H., CH.CBr.C.H., and C.H., Ci.C.c.H., Converted by benzene and Al₂Cl, into s-tetra-phonylathene some tri-phenyl-methane being also formed (Anschütz, A. 235, 207).

Tri - bromo - s - di - phenyl - ethane C, H, Br, From s-di-phenyl-ethane, water, and Br (F. a. S.).

Nacreous laminæ, v. sl. sol. alcohol; decom• poses at 170°.

Tri - bromo - s - di - phenyl - ethane C1.H1.Br. [207°-211°]. From s-di-phenyl-ethane and dry Br (M.).

Tri-bromo-s-di-phenyl-othane

C,H,.CBr,.CHBr.C,H,. Bromo-stilbens dibromide. [100°]. From bromo-s-diphenyl-ethylone and Br (L. a. S.). Needles (from alcohol). Docomposed by distill tion intoll Br, PhCBr:CBr.Ph, and PhC; CPh. / techolic KOH gives PhC:CPh.

Tri-bromo-u-i.-phsayl-ethane (CaH,)2CH.CBr₃. [89']. From bromal (1 mol.), benzene (2 mols.) and conc. H.SO (Goldschmiedt, B. 6, 985). Monoclinic prisms (fro n ether). Alcoholie KOH gives HBr and di-bromo di-phenyl-

cthylene.

Hoxa-bromo-s-di-phenyl-cthylene C₁₁II_sBr_e. From s-di-phenyl-cthane and excess of Br. Hard prisms (from benzene) (F. a. S.).

BROMO-PHENYL-ETHYL-AMINE v. BROMO-

AMIDO-PHENYL-ETHANE.

BROMO-u-DI-PHENYL-ETHYLENE C, II, Br i.e. $(C_0\Pi_s)_2$ C:CHBr. [50°](H.); [40′](A.). (c. 170°) at 11 mm. (A.); (above 300°) [11.). Formed by warming the di-bromide of u-di-phenyl ethylene (Hepp, B. 7, 1410; Anschütz, A. 235, 160). Prisms; sl. sol. cold alcohol.

Bromo-s-di-phenyl-ethylene

C.H. CBr:CH.C.H. Bromo - stilbene. From stilbene di-bromide by distillation or treatment with alcoholic KOH. Prisms. AgOAc gives PhC(OAe):CHPh (Limpricht a. Schwanert, A. 145, 340; 155, 72).

Di-bromo-u-di-phenyl-ethylone(CaHa), C:CBr., [83°]. (above 300°). From CPh_H.CBr, and alcoholic KOII (Goldschmiedt, B. 6, 985). Noodles

(from alcohol-cther)

Di-bromo-s-di-phenyl-ethylene C_oH_{..}CBr:CBr.C, I_s. Tolune di-bromide. [208]. Leaflets. Prepared by the action of bromine on tolane. An ischeride [61'] is also formed in small qu ntity (Limpricht a. Schwanert, A. 145, 318; Liebermann a. Homeyer, B. 12, 1971). Converted by benzene and Al_Cla in presence of CS2 into s-tetra-phenyl-ethane (Anschütz, A. 235, 209)

BROMO-PHENYL-GLYCOCOLL v. BROMO-PHENYL-AMIDO-ACETIC ACID.

DI-BROMO DI-PHENYL-GUANIDINE

C₁₃H₁₁Br₂N₃. From di-phenyl-guanidine hydrochiloride, gater, and Br (Hofmann, A. 67, 148). Seales (from a.coliol).—B'HCl. -B'nII2PtCl.

Tri-brome-tri-phonyl-guanidino
C₁₉H_{1-N-3}Br₂Cl i.e. (O_eH₁PrNH)₂C:N.C_eH₂Br.
White amorphous powder. Prepared by tho action of p-bromaniline on iso-cyan-p bromophenyl-chloride.—B If Cl: white crystals, easily soluble in alcohol and ether.—(B'HCl), Pt Cl.: light yellow plates (Dennstedt, B. 13, 232).

p.BROMO-PHENYL-MERCAPTAN

7. II, Br.SH. [75°]. (231°). From p-bromobenzene sulphochloride, tin, and HCl (Hübner a. Alsberg, A. 156, 327). Formed also by boiling brome-phenyl-cysteine or brome-phenyl-mercapturic acid with NaOHAq (Baumann a. Preusse, B. 12, 806; H. 5, 819). Laminos (from alcohol); volatile with steam; sl. sol. hot water. Cone. H.SO, at 120° forms a green solution, turning blue. Sodium-amalgam forms phenyl-mcreaptan. Chloral forms a compound

[72°]. HCl passed into a mixture of p-bromophenyl mercaptan and benzoic aldehyde forms di-p-bromo-di-phenyl-di-thio-benzaldehydate C.H., CH(S.C.H.Br), [80°] (Baumann, B. 18, 885). HCl and acetone form di-bromo-di-thio-di-methyl ketate (CH₃)₂C(S.C₂H₄Br)₂ [90°]. p-BROMO-PHENYL-MERCAPTURIC ACID

C, H, BrNSO, i.e. C, H, BrS. CMe (NIIAe). CO2H. Acetyl-bromo - phenyl - amido - thio - lactic acid. [153]. S. 1.4 at 100°. Occurs in the urino of animals which have taken brome-benzene. Formed also by treating brome phenyl-cystemo with benzenc and Ac.O (Baumann a. Preusse, B. 12, 806, 77. 5, 309; Baumann, B. 15, 1732; Jaffe, B. 12, 1092). Needles; insol. cold water, sol, hot water and alcohol. Lavorotatory in alcoholic solution, dextrorotatory in alkaline solution. Boiling aqueous NaOII gives p-bromophonyl mercaptan, NH_a, HOAe, and pyruvic acid. Boiling conc. HChAq or dilute H_aSO₁ gives acetio acid and bromo-phenyl-cysteme. Conc. H2SO, gives a blue solution.

Salts.—BaA', 2aq: S. 2 at 15°.—MgA', 9aq.

-NH A': S. 3 at 15°

BROMO-DI-PHENYL-METHANE C13H1,Br i.e. CHPh.Br. [45°]. From di-phenyl-methane (1 mol.) and Br (1 mol.) at 150° (Friedel a. Balsohn, Bl. [2] 33, 339, 587). Crystals, v. c. sol. benzene. Water at 100° forms di-phenylcarbinol and its anhydride. Boiling alcohol forms CHPh₂.OEt. CHPh₂.NH₂. Conc. NH_aAq gives

Di-bromo-di-phenyl-methane CPh.Br.,. From di-phenyl-methane (1 mol.) and Br (2 mols.) at Water at 150° converts it into Decomposed on distillation 150°. Liquid. henzophenone. giving tetra-phenyl-ethyleue. Sodium and water

form tetra-phenyl-ethane.

Bromo-tri-phenyl-methane $\mathbf{C}_{19}\mathbf{H}_{13}\mathbf{Br}$ Ph. CBr. [152°]. Obtained by prominating triphenyl-methane in sunlight or at 150° (Allen a. Köllicker, A.227, 107; Henderson, C. J. 51, 224; Schwarz, B. 14, 1520). Yellow hexagont! rhomboliedra (from CS₂) a: c=1: 784 (Hintze, Z. K. 9, 536). Decomposed above 200° into HBr and phenylene - di - phenyl - methane. Successivo treatment with boiling HOAc and water forms NII, forms Ph3CNH2. tri-phenyl-carbinol. KCy gives Ph, C.Cy. Potassium sulphocyanide gives Ph₃C.SCN.

TRI-BROMO-DI-PHENYL-METHYL-AMINE v. Tri-bromo-methyl-di-phenyl-amineo

BROMO-PHENYL-METHYL-FURFURANE TETRA-BROMIDE Cull, Br.O. [210°]. Bronzy plates. Formed by the action of bromine on HC-CH

phenyl - methyl - furfurane

PhC-O-CMc

(Paal, B. 17, 2760).

BROMO-PHENYL-METHYL-PYRAZOLONE v. Bromo-oxy-phenyl-methyl-pyrazol.

γ-BROMO - γ - PHENYL - DI - METHYL - SUC-CINIC ACID 1'h.CHBr.CH(CO2H).CHMe.CO2H. From the lacton of γ -oxy- γ -phenyl-di-methyl-succinic acid (q, v) and cone. HBr at 0° (Fittig a. Penfiold, A. 216, 123). Small crystals (from benzene). V. sol. alcohol or ether, m. sol. benzone. Warmed with water it gives off CO2 and forms plates of an acid which appears to be Ph.CH:CH.CHMe.CO.H.

TETRA-BROMO-PHENYL . METHYL . THIO. PHENE C11H Br.S. [187°]. Formed by bromination of phenyl-methyl-thiophene. Fine needles or plates. V. sol. benzene, ether, and petroleumspirit, m. sol. alcohol and acetic acid (Paal a. Püschel, B. 20, 2559).

BROMO-PHENYL-METHYL-URETHANE 2. Methyl ether of Bromo-Phenyl-Carbamio acid.

BROMO-PHENYL MUSTARD OIL v. BROMO-PHENYL THIO-CARBIMIDE

DI - BROMO - PHENYL - (a) - NAPHTHYL - AMINE C₁₆H₁₆Br₂NH. [140°]. White needles. Prepared by bromination of phenyl (a)-naphthyl-amine (Stroiff, B. 13, 1853; A. 209, 158).

Totra-bromo-phenyl-(β)-naphthyl-amine C₁₆H₈Br₄NH. [198°]. Sl. sol. alcohol, ether and C₈H₈, m. sol. CS₂ and CHCl₃. Prepared by further bromination of the dibromo-derivative.

Tri-bromo-phenyl-(a)-naphthyl-amineC₁₆H₉Br₃NH. [137°]. Colourless prisms. Soluble in alcohol and benzene. Prepared by bromination of phonyl-(a)-naphthylamine (S.).

Tetra-bromo-phenyl-(3)-naphthyl-amine C₁₈II_{*}Br₄NII. [203°]. Formed by the action of bromine upon an acetic acid solution of benzeneazo-phenyl-(\$)-naphthyl-amine, or by bromination of phenyl- (β) -naphtlyl-amine. Long white silky needles (Zincko a. Lawson, B. 20, 1170).

BROMO-PHENYL-OCTANE v. BROMO-OOTYL-

p-BROMO-PHENYL-OXAMIC ACID

C₆H₄Br.NH.CO.CO₂H. [198°]. S. 194 at 25°. From di-bromo-di-phenyl-oxamide and alcoholio KO11 (Dyer a. Mixter, Am. 8, 355). Crystallino fibres, sol. alcohol and other. KOHAq liberates Salts. - KA': monoclinio p-bromo-aniline. scales.—CaA',—AgA'.—BaA', Ethyl ether EtA'. [156°]. From phenyl-

oxamic ether and Br (Klinger, A. 184, 263).

DI-BROMO-DI-PHENYL-OXAMIDE $(C_a \Pi_i Br. NH)_2 C_2 O_2$, [above 300°]. From Br and di-ph-enyl-oxamido in HOAe (D. a. M.)

DI-BROMO-DI-PHENYL OXIDE (C.11,Br)₂O. [58°]. (above 360°). From Br and di-plicnyl oxido in CS₂. Long plates (from alcohol) (Merz a. Weith, B. 14, 191).

TRI-BROMO-TRI-PHENYL PHOSPHATE C.H.BrO),PO. From tri-phenyl phosphate and Br at 180°. Nacreous scales (Glutz, A. 143, 193).

p-BROMO-PHENYL-PHTHALIMIDE C = 0

>0 C = N.C.H.Cl $[204^{\circ}].$ Fine

needles or seales. Sol. C₈H₆ and acetic acid, less in other. Prepared by heating p-bromaniline with phthalic anhydride (Gabriel, B. 11, 2261)

BROMO-PHENYL-PROPANE v. BROMO. PROPYL-BENZENE.

o-BROMO-8-PHENYL-PROPIONIC ACID [2:1] C₆11, Br.C₂H₄, CO₂H.o-Bromo-hydrocinnamic acid. [99°]. Selles. Formed by reduction of o-bromo-cinnamic acid with HI and P (Gabriel, B. 15, 2295).

m-Bromo-β-phenyl-propionio acid [3:1] C.11, Br.C.H., CO.H. m-Bromo-hydrocinna-mic acid. [75°]. Formed by reduction of mbromo-cinnamic acid with P and HI. Also by eliminating the NH2 group from (3:4:1)-bromoamido-phenyl-propionic soid by diazotising and treatment with alcohol (Gabriel, B. 15, 2294).

Short thick prisms. V. sol. alcohol, ether, bsnzene, ohloroform, and CS2.

p-Bromo-\$-phenyl-propionio acid [4:1] C.H.Br.CH., CH., CO.H. [135°]. From \$-phsnyl-propionic acid and Br in the cold (Göring, C. C. 1877, 793, 808; Gabriel a. Zimmermann, B. 13, 1683). Flat noedles (from CS2). Oxidation gives p-bromo-benzoio acid.

β-Bromo-β-phenyl-propionic acid

C.H., CHBr.CH., CO.H. [137°]. From cinnamic acid and HBr (Fittig a. Binder, B. 10, 518; A. 195, 132; Anschütz a. Kinnicutt, B. 11, 1221). Also from β-bromo-β-oxy-phenyl propionio acid and HBr (Glaser, A. 147, 96). Laminæ. Decomposed by heat into HBr and cinnamic acid. Boiling water forms β-oxy-β-phenyl-propionic acid. Cold NaOHAq gives styrene and CO.,

α-Bromo-α-phenyl-propionio acid CH..CPhBr.CO.H. Bromo-hydratropic acid. [94°]. From atropic or atrolactic acids and cold conc. HBrAq (Fittig a. Wurster, A. 195, 145; Merling, A. 209, 13). Tables, insol. water, sol. ordinary solvents. Boiling Na₂CO₃Aq produces

atrolactio acid.

β-Bromo-α-phenyl-propionic acid CH₂Br.CHPh.CO₂II. [94°]. Formed by heating atropic acid with conc. HBrAq at 100°. Prisms, insol. water. Boiling Na CO Aq produces tropic acid, styrene, and a very little atropic acid. Ammonia forms \$\beta\-amido-a-phenyl-propionic acid [169°] (M.).

αβ-Di-bromo-β-phenyl-propionio acid C₆H₅.CHBr.CHBr.ĈO₂H. [195°] (G.); [201°] (S.). From cinnamic acid and browing-vapour (Schmidt, A. 127, 320; Fittig a. Binder, A. 195, 140). Also from α-bromo-β-oxy-phenyl-propionic acid and HBr (Glaser, A. 147, et). Lamina (from alcohol); v. sol. ether and alcohol, v. sl. sol.

Reactions. - 1. Sodium amalgam forms phenyl-propionic acid .- 2. Boiling water gives cinnamic acid, bromo-oxy-phenyl-propionic acid, phenyl-acetic aldehydo and w-bromo styrene.

3. Alcoholic KOH gives a- and \$\beta\$- bromo-cinnamic acids.

Salts.—NaA'.—BaA'.

Methyl ether MeA'. [HI7°] (Anschütz, B. 12, 538).

Ethyl ether EtA' [69°]. From cinnamio ether and bromino (Perkin, jun., C. J. 45, 172). n. Propyl ether PrA'. [23°].

aβ-Di-bromo-a-phenyl-propionic acid CH_Br.CPhBr.CO.H. [1160]. From Br and atropio acid in CS₂ (Fittig a. Wurster, A. 195, 145). Nesdles (from CS₂). Decomposed by boiling waterinto CO2, HBr, and acetophenone. Sodiumamalgam forms a-phenyl-propionic acid and oxy-phenyl-propionio aoid. Excess of NaOllAq gives atroglycerio acid CaH1aO, (Fittig a. Kast, A. 206, 80).

Tri. brome-phenyl-propionic acid C_cH_s.CHBr.CBr_s.CO_sll. [151°]. From brome-cinnamic acid [120°] and Br (Glaser, A. •143, 335 : Stockmoier, Bn. 2, 872 ; Kinnicutt, Am. Small flat monochinio needles (from diluts alcohol). Boiling water gives CO, dibromo-styrene, bromo-cinnamio acid, and di-

bromo-oxy-phenyl-propionic acid.

Tri-bromo-phsnyl-propionio acid C.H. CBr. CHBr. CO.H. [148°]. Formed by tho combination of the brome-cinnamic acid [159°]

with Br. Triclinic prisms. Sol. hot benzsno, v. e. sol. alcohol and other, sl. sol. cold CS2. It decomposes at its melting-point evolving HBr. By standing for a short time with water it is converted into a neutral oil; hot water decomposes it at once (Michael a. Brown, B. 19, 1380). Tri-bromo-α-phenyl-propionic acid

C,H.Br,O. [150°]. From bromo-atropio acid and Br (F. a. W.). Needles (from ligroin).

αβ-D1-BROMO, FHENYL PROPIONIC ALDE-HYDE Cally CHI'. CHBr. CHO. Cinnamic aldehyde dibromide. [c. 100°]. Small needles. Formed by the direct combination of cinnamio aldehyde and brourine. It readily splits off HBr on heating, giving bromo-cinnamio aldehydo (Zincke a. Hagen, B. 17, 1811).

DI-BROMO-B-PHENYL-PROPYL ALCOHOL

C₉H₁₀Br₂O i.e. C₈H₅.CHBr.CHBr.CH₂OH. Stycerin dibromhydrin. Styrone dibromide. [74°]. From Br and cinnamyl alcohol in CHCl, (Grimanx, BL [2] 20, 120). Tables or needles (from other). Insol. water. Boiling water converts it into C_cH_s.CH(OH).CH(OH).CH₂OH.

Acetyl derivative C.H.CHBr.CHBr.CHL.OAc: [86°]; prisms.

DI - BROMO - PHENYL - PROPYLIDENE -ANILINE CaH, N:CHLCHBr.CHBrPh. [175°]. From cinnamylene-anilino and bromine (Schiff, A. 239, 384). Needles (from alcohol).

DI-BROMO-PHENYL-PYRAZOL DRIDE C_pII_pBr₂N_{...} [98°]. Di-bramo-phenyl-pyrazoline. From phenyl-pyrazoline and Br in chloroform (Fischer a. Knoevenagel, A. 239, 199). Plates (from alcohol); v. sl. sol. water. In diluto acid solution it gives a violet colour with K₂Cr₂O₂. Alcoholic KOH forms C₂H₄Br(OEt)N₂ [60], which crystallises from alcohol in palo yellow prisms. Boiling HClAq gives off EtCl and forms brome-oxy-phenylpyrazol CallaBr OH)N. [214°]; this has acid characters and orms greenish-yellow crystals (from alcohol).

DI-E LOMO-(α)-PHENYL-PYRIDINE CARBOXYLIC ACID C₁₃H, Br. NO₄. [205°]. From [2:I] C₆H₄(CO₂H).C₅H₃N(CO₂H) [3:2] and bromino (Skraup a. Cobenzil, M. 4, 469). Granules, v. sl. sol. water, m. sol. warm alcohol.

DI-BROMO-DI-PHENYL SULPHIDE (C₀H₄Br).S. [110°]. From di-phenyl sulphide and Br, or from (C₀H₄NH₁).S by diazo-reaction. Nacreous laming (Krafft, B. 7, 1165).

Di-p-beomo-di-ph inyl disulphide (C. II Br) S. [94°]. From p-bromo-phenyl mercaptan by atmospheric oxidation. Plates; not volatile with steam (Hübner a. Alsberg, 1, 156, 328; Baumann a. Preusse, H. 5, 329

DI-p-BROMO DI PHENYL SULPHONE

(C₆H₄Br)₂SO₂. [172]. From p-bromo-beuzene and ClSO₃H (Armstrong, G. J. 24, 173) or SO₄ (Nölting, B. 8, 594). Also from brome-benzens, benzeno sulphochloride, and Al.G. (Beckurts a. Otto, B. 11, 2065). Needles, sl. sol, hot alcohol. p. BROMO - PHENYL - THIO - CARBAMIC

ETHER C.H. BrNSO i.e. C.H. Br. NH.CS.OEt. Bromo-phenyl-thio-urethane. [105°]. From p-bromo-phenyl thio-carbinide and alcohol at 120° (Dennstedt, B. 13, 231). Slender nesdles.

p-Bromo-phenyl-di-thio-carbamic sther CaH.Br. NII.CS.SEt. Bromo di-thio carbanilie ether. [89°]. From p-bromo-phenyl-thio-carbimide and mercaptan at 140° (D.).

p.Brome-phenyl-thie-carbinide

O.H.Br.N.CS. p-Brome-phenyl mustard oil.

[61°]. Prepared by heating p-brome-aniline with CS₂ in alcoholic solution with a little aqueons KOH; the resulting thio-urea being distilled with P2Os or heated with cone. HCl at 160° (Dennstedt, B. 13, 230; Weith and Landolt, B. 8,

DI-n-BROMO-DI-PHENYL-DI-THIO-CINNA-MIC ALDEHYDATE C.H.C.H., CH(S.C.H.lir), p-Bromo-phenyl-mercaptal of a mamic aldehyde, [107°]. Formed by passing Hall gas into a mixture of p-bromo-phenyl mercaptan and cinnamio aldohyde (Baumann, B. 18, 885). Long colourless needles. Sl. sol. cold alcohol and ether.

BROMO-PHENYL-THIO-GLYCOLLIC ACID C₈H₁BrSO₂ i.e. C₅H₁Br,S.CH₂CO₂H₃ [112]. From C₆H₂S.CH₂CO₂H and Br in CS₂ (Classon, Bl. [2] 23, 441).

DI-p-BROMO-DI PHENYL-DI-THIO-DI-METHYL - KETATE (CH,)_::C:(SC,H,Br)_:. p-Bromo-phenyl-mercaptol of acetone. [90]. Long transparent prisms. V. sol. hot alcohol, ether, and benzene. 1 ormed by passing HCl gas into a mixture of p-bromo-phenyl-mercaptan and acctone (Baumann, B. 18, 888)

p-BROMO PHENYL-THIO-UREA

NIL CS.NHC H Br. [183"]. From bromophenyl - thio - earbimide and alcoholic NII, (Dennstedt, B. 13, 231). Needles.

p-Bromo-di-phenyl-thio-nrea NHPh.CS.N1I.C,H,Br [158°]. From bromophenyl-thio-earbimide and amiline (D.).

Di - p - bromo - di - phenyla thio - urea CS(NH.C₄II₄Br)... [178°]. From p - bromoaniline, CS., and alcohol in presence of some KOHA4 (D.; Otto, B. 2, 409). Prisms. BROMO - PHENYL - THIO - URETHANE v.

BROMO-PHENYL-PHIO-CARBAMIC ETHER.

TETRA-BROMO-PHENYL-PLOLUIDINE
C₁₃IL_pBr₄N. [156°]. Formed by adding a solution of Br in glacial HOAc to an alcoholic solution. lution of phenyl-p-toluidine (Bonna, A. 239, 58).

Hepta-bromo-pheayl-p-toluidino C13H8Br.N. [185°]. From phenyl-p-toluidine and Br at 280°. Endeca-bromo-phenyl-p-toluidino

C₁₃H.Br₁₁N. [296]. Formed from phenyl-p-toluidine and Br at 310°.

BROMO - p - PHENYL - TOLUENE C₁₃II₁₁Br. 7°-131°]. From p phenyl-toluono and Br. [127°-131°]. From p-phenyl-tolueno and Br. Small tables (from alcohol) (Carnelley a. Thom-

son, C. J. 47, 589). DI-p-BROMO DI PHENYL-UREA

C₁₃H₁₀Br₂N₂O i.e. CO(NH.C₆H₁Br)₂. Di-bromotarbanilide. From di-phonyl-thio-urea and Br (Otto, B. 2, 409). Formed also by decomposition of the product from COCl, and diazobenzene-p-bromanilide. Prepared by the action of earbonyl chlorido on p-bromanilino (Sarauw, B. 15, 45). Whits glistening plates. Sublimes at 225° without melting. Sl. sol. alcohol and benzens.

Tetra-bromo-di-phsnyl-urea

CO(NH.C.H.Br.).. Sublimes in needles (O). BROMO-PHENYL URETHANE v. Buomo-PHENYL-CARDAMIC ETHER.

DI BROMO PHENYL VALERIC ACID C.H. CHBr. CHBr. CH2. CH2. CO2H. [109°]. From styryl propionio acid and Br (Baoyer a. Jackson, B. 13, 122).

BROMO-PHLOBAPHENE v. PHLOBAPHENE. BROMO - PHLORETIC ACID v. PHLORETIC

TRI-BROMO-PHLOROGLUCIN C.Br.(OH), [151°]. Formed by brominating phloroglacia (Hlasiwetz, A. 96, 118; Herzig, M. 6, 885). Long needles (containing 3aq) (from water). Reduced by tin and HCl to phloroglucin. Converted by cold HNO₃ (S.G. 1·4) into tri-bromodi-nitro-propionic acid (Benedikt, A. 184, 255).

Tri-acetyl derivative [183°] (Herzig, M. 6, 887). C,Br,(OAc)

Hexa-bromo-phloroglucia dibromide CoBr, (OBr)3. [1180]. The final product of the bromination of phloroglucin (Hazura a. Sene-dikt, M. 6, 702). Small golden needles (from CHCl₂). At 190° it gives off Br (1 mol.). Aqueous SO₂ reduces it to C_aBr_s(Oli)₃. IICl form C_aH_aCl₃(OII)₃. Tin and

DI - eso - BROMO - v - PHOSPHO - AMIDO-

BENZENE SULPHONIC ACID Tho (HO)₂PO.NH.C₆H₂Br₂.SO₃H. ehlorida Cl2PO.NH.C.H2Br2.SO2Cl is formed by treating di-bronn-anido-benzene sulphonic acid with PCl₂. It is converted by alcohol into the ether-chloride (EtO),PO.NII.C,II_Br₂.SO₂Cl [c. 170] (I.aar. J. pr. [2] 20, 257).

BROMO-PHTHALIC ACID C₂II₁Br₁CO₂El₂Cl [c. 170] (I.aar. J. pr. [2] 20, 257).

[1:3:4]. [140°]. Formed, together with its isomeride, by brominating phthalio acid (Faust, A. 160, 62; Pechmann, B. 12, 2124; cf. Guareselii, A. 222, 295, Stallard, C. J. 49, 187). Powder, v. sol. water, alcohol, and ether.

Salts.—K.A.²2aq: long needles (from alcohol).—BaA" 2aq: crystalline powder; sl. sol. water.—CuA",—Ag.A": cheesy pp., sl. sol. water.

Anhydride C. H. Br(CO). [65°]. (300°—

310°).

Ethyl ether Et, A": (295°); liquid. c-Bromo-phthalic acid C, H, Br (CO, H), [1:2:3].

[176°] (G.); [174°] (M.); [197°] (C. a. T.).

Formation.—1. Together with the preceding. by brominating phthalic acid (Pechmann). 2. By the oxidation of bromo-nitro-naphthalene 122.5°] with KMnO, (Guareschi, A. 222, 292), of bromo-(β)-naphthol with KMnO, (Meldola, C. J. 47, 512), of liquid bromo-ditelyl (Carnelloy a. Thomson, C. J. 47, 591), of di-bromo-naphthalene [130°] with CrO₃ in HOAe (Guareschi, B. 19, 134), of C₁₆H₃Dr₄(OH) [1:x:3:4:2] with KMnO₄ (Smith, C. N. 40, 87), and of (a)-bromenaphthalenc, and bromo-o-tolnic acid [167°] with dilute HNO3 (Raeine, A. 239, 76). The bromoo-toluie acid may be prepared from bromo-otoluidino C,H,MeBr(NH.) [1:5:2]? by Sand-meyer's method; 70 g. of bromo-toluidins gave

53 g. of bromo-phthalic acid. Properties.—White prisms (from water); v. sl. sol. ehloroform, m. sol. water, alcohol, and ether. With resprein it gives a fluorescein (Nourrisson, B. 20, 1016). The salt BaA" forms pearly

plates, sl. sol. water.

Anhydride C,H3Br(CO)2O. [135°] (Msldola); [132°] (G.); [125°] (Smith); [108°] (Nourrisson); [95°] (Racine). Nesdles. Heatod with phenol and H2SO, it forms a body (? bromophthalids) that dissolves in alkalis forming a purple solution.

Di-bromo-phthalic acid C.H.Br.(CO.H). [1:4:2:3]. [135°]. From di-bromo-naphthalene [82°] and HNO. White crystalline powder, sol. boiling water and alcohol (Guareschi, A. 222, 274). On melting it changes to its anhydride.

Salt .- Na A".

Anhydride [208°]. Pearly needles (by sublimation). Heated with phenol and H.SO, it forms a product (dibromophthalein) that dis-

solves in potash forming a purple solution.

Di-bromo-phthalio acid C₆H₂Br₂(CO₂H)₂. [206°]. Formed by oxidation of penta-bromo-(a)-naphthol or of tetra-bromo-(a)-naphthoquinone by means of diluto HNO₃ at 150° (Blümlein, B. 17, 2490). Colourless needles. V. e. sol. alcohol, ether, and hot water.

States.—A"Ag₂: small colourless plates, sparingly soluble.—A"Ca: pp.—A"Ba: pp.

Anhydride C₈H₂Br₂COOO.[208°]. Sub-

limes in long colourless needles. Easily solublo

in alcohol, sparingly in water and ether. Formed by heating the acid. Heated with resorcin it yields a di-bromo-fluoresceïn.

Di-bromo-tetra-hydro-phthalio aoid

CeHeBr2(CO.H)2. Di bromo-tetra-hydro-benzenedi-o-carboxylic acid. Formed by the direct combination of dry di-hydro-phthalio acid with bromine vapour. Rhomboliedra (Bacyer, B. 19, 1810)

Tri-bromo-phthalio acid CaHBr. (CO.H). [191°]. Formed by oxidation of penta-bromo-(β)-naphthol or of tetra-bromo-(β)-naphthoquinone with HNO, (Flessa, B. 17, 1482). Small silvery plates or needles. Nearly insoluble in petroleum ether and in cold water.

Salts.-AgzA".-CaA" 2aq.-BaA" 2aq.

Anhydride C.HBr.(CO).O: [157]; sub-limes in white plates; easily soluble in alcohol and ether, nearly insoluble in cold water.

Tetra-bromo-phthalic acid OaBr (CO2II)2 [266°]. Fermed by oxidation of tetra-bromo oxylene by heating with diluto HNO, and bromine at 170° (Blümlein, B. 17, 2493). Small needles or colourless prisms. V. sl. sol. water. With resoroin it yields a tetra-bromo-fluorescein.

Salts.-A"Ca.-A"Ba.

Anhydride C_eBr₄<CO>O. [259]. Formed by heating the acid. Sublimes in colourless glistening needles. Sl. sol. almost all solvents.

V. also Bromo-Terephthalic ACIDS. $C_{\alpha}H_{a}Br<_{CO}^{CII}$ > 0. BROMO-PHTHALIDE

[sore: 1]. [100°]. Formed in small quantity, together with bromo-o-toluic acid, by the action of bromine-water on o-toluic acid (Racine, A. 239, 76). Needles (from dilute alcohol); sublimes readily. Insol. cold water or Na₂CO₃Aq. Gives brome-e-toluic acid [167°] on oxidation.

(a)-Bromo-phthadide CoH (CO) O. [86°]. From phthalide and bromine at 140° (Racine, A. 239, 79; B. 19, 778). Small cubes or tables (from ether). May be distilled. Hygroscopic. Slowly decomposed by cold, quickly by hot, phthalic forming aldchyde - acid water, C.H.(CO.H)(CHO). Alcohol converts it into

KMnO, oxidises it to

C,H,(CO,Et)(CHO). Ammonia forms amidophthalio acid. phthalide, $C_{\bullet}H_{\bullet} < CO^{OH(NH_2)} > 0$. [167°].

Di-bromo-phthalide C, H, Br, O, i.e.

 $C_4H_2Br_2 < CO^2 > O\begin{bmatrix} 1 & 9 \\ 4 & 3 \end{bmatrix}$. [188°]. S. (94 p.c. alcohol) 87 at 15°. From di-bromo-naphthalene, CrO, and glacial acetic acid (Guareschi, A. 222) 282). Prisms or needles (from alcohol). Neutral reaction. Does not reduce ammoniacal AgNO. With phenol and H.SO, gives no dyo on hoating. DI-BROMO-PICENE v. PICENE.

BROMO - PICE N v. TRI - BROMO - NITRO -METHANE.

BROMO-PIP'A-HYDRONIC ACID v. BROMO-DI-OXY-PHENYL-VALMEIO ACID.

BROMO-PIPERIC ACID Dihydride. C₁₂H₁₁BrO₄ or

 $C\Pi_2 < 0 > C_cH_2Br.CH_2.CH_2CH.CH.CO_2H.$

Bromo-(\$\beta\$)-hydro-piperic acid. [171°]. From bromine and (β)-di-hydro-piperio acid (Fittig a. Buri, A. 216, 177; Weinstein, A. 227, 42). Streaky white plates (from benzene).—Salt.— CaA'2.

Reactions.—1. Not affected by boiling aqueous KOII.—2. KMnO, oxidises it, in neutral solution, to bromo piperonylic acid [204°], bromo-pipero- propionic acid (q. v.) and bromopiperonal; hence Br is in the benzene nucleus .-3. Sodium-amalgam forms piperhydronic or mcthylene-di-oxy-phenyl-valeric acid C12H14O40

DI-BROMO-PIPERINIDE v. BROMO-TRI-OXY-

PHENYL-VALERIO ACID.

BROMO-PIPERONYLIC ACID v. BROMO-DI-Y-BENZOIO ACID.

BROMO-PIPERO-PROPIONIC ACID v. Methylene derivative of Bromo-di-oxy-phenyl-pro-PIONIC ACID.

DI - BROMO - PREHNITOSE v. DI-BROMO-DURENE

BROMO-PROPANE v. PROPYL BROMIT'S.

Di-bromo-propane C₂H₆Br₂ i.e.
CH_aCHBr.CH₁ tr.(141-5'). Propylene bromide.
S.G. ⁹₆ 1 9617 (N nder, A. 214, 175). S.V. 1189
(Z.); 118-4 (Schiff).

Formation .- I. From propylene and Br (Reynolds, A. 77, 120; Cahours, C. R. 31, 291; Wartz, A. 104, 244).-2. From bromo-propylene and HBr .- 3. Together with trimethylene bromide by the union of HBr with allyl bromide (Geromont, Bl. [2] 16, 113, Reboul, Bl. [2] 17, 350). 4. From propyl bromide and Br (Linnemann,

A. 161, 41). Properties.-Liquid with sweet smell.

Reaction . - 1. Alcoholic KOH forms two bromo-propylencs and, finally, allylene (Sawitsch, C. R. 52, 399).—2. AgOAc gives the diacetyl derivative of propylone-glycol (Wurtz, A. Ch. [3] 4, 438). AgOBz gives the corresponding benzoyl derivative (Friedel a. Silva, C. R. 73, 1379).—3. Convert d into propyleno by Zn and HOAc or sodium amalgam in alcohol (Linnemann, B. 10, 1111) .- 4. Aqueous III at 150° gives isopropyl bromids .- 5. Heated to 100° with Ag.O and water it gives propionic aldehyde but no propylene-glycol (Beilstein a. Wiegand, B. 15, 1496).—6. Water (20 vols.) and PbO at 150° gives acetone, propionic aldehyde and propylene-glycol (Eltekoff, J. R. 10, 212).—7. Protracted boiling with water gives propyleneglycol (Niederist, A. 196, 349)

aa Di-brome-propane CH, OBr, CH, Methyl bromacetol. Bromacetol. Acetone bromide. (115°).

S.G. 18 18476; \$5 18314. M. M. 10-137 at 20.7° (Perkin). Formed in small quantity from acetone and PBr, or PCl_Br, (Linnemann, A. 138, 125; Friedel a. Ladenburg, Z. 1868, 48). Also from allylene and conc. HBr (Reboul, C. R. 74, 669); and from a bromo-propylene and HBr.

Reactions .- 1. Water at 160° gives acctono. 2. Zn and HCl give propano (Linnemann, A. 161, 67). - 3. Alcoholic KOH gives a bromopropylene Cll,.CBr:CH,.

ωω-Di.bromo.propane CH₃.CH₂.CHBr₂. (0. 130°). FMm ω-bromo-propyleno CH₂.CHI:CHBr and conc. HBr (Reboul, A. Ch. [5] 14, 467).

ω₁ω₂-Di-bromo-propane CH.Br.CH.CH.Br. Trimethylene bromidc. (165°). S.G. 22.006 (Z.); 2 2·018 (G.); ¹⁷⁻⁶/₄ 1·9228 (F.). S.V. 117·8 (Z.). Formed from ω₁ω₁·di-oxy-propane and HBr (Freund, M. 2, 639).

Preparation.—Allyl bromide is saturated with very nearly dry HBr at -16°, sealed up and left at 30° for 24 hours. The tube is opened and the operation repeated as long as any gas is absorbed (Geromont, A. 158, 370; Reboul, A. Ch. [5] 14, 472; Erlenmeyer, B. 12, 1354; A. 197, 184; Roth, B. 14, 1351; Bogomo litz, Bl. [2] 30, 23).

Reactions .- 1. Alcoholio KOH forms allyl bromide or allyl ethyl oxide. -2. Alcoholic NII, forms some amorphous bases (Niederist, M. 3, 840).-3. Heated to 100° with Ag2O and water it gives tri-methylene-glycol thus differing from the isomeric propylene bromide which gives propionic aldehydo (Beilstein a. Wiegand, B. 15, 1495).—4. Al.Br. converts it into propylene bromide (Gustavson, J. pr. [2] 36, 303).

ωωα-Tri-bromo-propane CH3-CHBr.CHBr2-(201°). From w. bromo - propyleno and Br Reboul, A. Ch. [5] 14, 181).

ωαα.Tri - bromo - propane Ch. CBr. CII.Br. (191°). S.G. ± 2.35. Formed T. the union of bromo-propylene with Br (Reboul, A. Ch. [5] 14, 481; C. Kolbe, J. pr. 133, 393).

s-Tri-bromo-propane CH₂Br.CHBr.CH₂Br. Tri-bromhydrin. [17°]. (220°). S.G. 23 2.44.

Formalion. - 1. From di brombydrin or epibromhydrin and PBr, (Berthelot a. de Luca, A. 101, 76; Henry, A. 154, 369).—2. From allyl bromido and Br (Folleus, A. 156, 168). 8. By bromination of isopropyl bromide (Linnemann, A. 136, 63).—4. From tally liodide and Br (Wurtz, A. 104, 247).

Properties. - Prisms or liquid.

Reactions. — 1, Alcoholic KOII gives CH:C.CH,OEt. — 2. Solid KOII gives two dibromo-propylenes.—3. AgOAc gives triacetin C₃H₃(OAc)₃.—4. KCy gives C₂II₁(Cy)₃.—5. Alcoholio NH₄ forms di-bromo-di-allyl-amine and then methyl-pyridine.

Tetra-bromo-propane CH_a.CBr_a.CHBr_a. Allylene tetra-bromide. (£25°-230°); (110°-130°) at 10 mm. S.G. 2 294. From allylene and Br (Oppenheim, Bl. [2] 2, 6; 4, 434; A. 132, 124). Liquid, decomposed by alcoholic KOH into HBr and tri-bromo propylone (o. 193°) (Pinner, A. 179, 59).

Tetra-bromo propane CH₂Br.CBr₂.CH₂Br. [195°]. From iso-allylene and Br (Hartenstein, J. pr. [2] 7, 817).

The following tetra-bromo-propanes have also been prepared:

(a): (251°); S.G. 2.64. From propylene bromide and Br (Reboul, A. Suppl. 1, 232)

(b): [60°]; (o. 235°). By brominating isopropyl bromido (Linnemann, A. 136, 64).
(c): (226°); S.G. 2·47. From propylene

bromide and Br (Cahours, A. 76, 284).

Penta-bromo-propanes C₄H₂Br₅. The followiug have been described:

(a): (255°); S.G. 2·60. From propylene

bromide and Br (Cahours, C. R. 31, 291).

(b): [173°]. From tri-bromo-propylene and Br (Pinner, A. 179, 60).

(c): CHBr₂.CBr₂.CHBr₂. S.G. 19 3-01. From propargyl bromide and Br (Henry, B. 7, 761).

BROMO-PROPIOLIC ACID CBr C.CO.H. Formed by decomposing nucebromic acid with baryta (Jackson a. Hill, B. 11, 1675; Am. 3, 121). Prisms (from ether). V. e. sol. water (erystallising therefrom with and); may be partially sublimed at 100°. Boiling water liberates bromo-acetylene; boiling baryta forms bromoacetyleno and also malonic acid. The acid gives with di-bromo-acrylio acid a compound C₃HBrO₂C₃Br₂11₂O₂ [105°].

Salts. - Bah' xaq .- AgA'.

a-BROMO PROPIONIC ACID CaHaBro, i.e. CH_a.CHBr.CO_.H. [25°] (W.). (206° cor.). Formed by heating propionic acid (1 mol.) with Br (I mol.) for several days at 150° (Friedel a. Machuea, C. R. 53, 408; A. 120, 286). Formed also from lactic acid and IIBrAq at 100° (Kekulé, A. 130 16).

Reactions .- 1. Sodium-amalgam forms propionio acid .- 2. Boiling water and ZnO givo lactic acid. Tho K salt changes slowly to lactate in cold aqueous solution.—3. Alcoholic NII, forms alanine.—4. Finely divided Ag at 150

gives s-di-methyl-succinic acid.

Ethyl other EtA'. (162); (130°) at 160 mm. S.G. 11 1.10. From the acid (Bischoff, A. 206, 319) Also from lactic ether and PBr, (Henry, A. 156, 176). Preparation.—Propionic acid (300 g.) is converted into the bromide by adding amorphous phosphorns (31 g.) and slowly running in bromine (400 g.). After the evolution of HBr has ceased the mixture is brominated by heating to 40°-50° and slowly running in more bromine (640 g.). When the whole of the bromino has disappeared the bromo-propionyl bromido is converted into the ethyl ethor by the addition of absolute , cohol. It is then treated with water, washed and fractionated. The yield from 300g, of propionia acid amounts to 640 g. of boiling-point 156°-160° (Zelinsky, B. 20, 2026).

Bromide CH₃.CHBr.CO.Br. (155°). From

propionyl bromide and Br; also from propionie acid, P, and Br (Weinig, A. 242, 163). ZuMc₂, followed by water gives methyl isopropyl ketone and di-methyl-isopropyl-carbinol (Kashirski, C. C. 1881, 278).

Imcde (CII, CHBr.CO), NH. [148°]. Formed by the action of water on the compound (C₂H₃NBr₂) of propionitrile with Br (Engler, A. 142, 71). Needles, m. sol. hot water.

B. Bromo-propionic acid CH2BACH2.CO2H. [62°]. Small glistening plates. Formed by heating hydracrylic acid with HBr at 120° (Beckurts a. Otto, B. 18, 227).

aß-Di-bromo-propionic acid

CH.Br.CHBr.CO₂H. [64°]. (227°). S. 1945 (?19°45) at 11°; S. (ethor) 304 (?3°04) at 10°. **Formation.**—1. Got by oxidising di-bromo-

propyl alcohol with HNO₃. The yield is bad (Münder a. Tollens, B. 5, 73; A. 167, 222) —2. Also by union of acrylic acid with Br (Caspary a. Tollens, A. 167, 256),-3. From acrolein dibromide and HNO₃ (Linnemann a. Penl, B. 8, 1097).—4. By the action of HBr upon abromo-acrylic and aa di bromo-propionic acids (Philippi a. Tollens, A. 171, 333)

Properties. - Monoclinic crystals (Haushofer, J. 1881, 687; Zepharovich, J. 1878, 693). It crystallises in two forms: tables [61] and prisms [51°]; the latter slowly change into the former. The salts readily split off bromide,

forming β-bromo acrylic acid.

Reactions. -- 1. Converted into acrylic acid by KI and water, or by Zn and H2SO4 (v. Zotta, A. 192, 102; C. a. 1.) -2 Water at 120° gives bromo-oxy-propionic acid (Melikoff, J. R. 13, 227).

Salts. -AgA'. - NII, A'. - KA'. - CuA', 2aq. -SrA', 6aq. Methyl ether MeA. (205.8° cor.) (Weger,

A. 221, 84).

Ethyl ether Eth. (214 6° cor.). Allyl ether C₃H₄A. (215°-220°). Propyl ether PrA. (233° cor.).

aa-Di-bromo-propionio acid CH, CBr, CO, II. [65°] (F. a. M.); [61°] (P. a. T.). (c. 224°). From a-bromo-propionic acid and Br (Friedel a. Machuca, C. R. 54, 220; Philippi a. Tollens, B. 6, 515). Trimetric tables.

Reactions .- 1. Zn and H.SO, reduce it to propionicacid .- 2. Alcoholic KOII gives a-bromoacrylic acid. -3. Ag O forms pyrav c acid. -4. Finely divided silver in benzene gives di-methylmaleio anhydride. - 5. The salts are more stable than those of the preceding acid; but the silver salt warmed with water changes to pyruvic acid

(Beckurts a. Otto, B. 18, 235).

Salts. -- NH, A' ½aq. -- NaA'. -- KA'aq. -BaA', 9aq. -- CaA', 2aq. -- SrA', 6aq.

Methylether MeA', (c. 177°). S.G. 2 1-901.

Ethyl ether EtA'. (191°). S.G. 6 1.754. Propyl ether PrA'. (c. 202°). S.G. 2 1.684. Isobutyl ether Prell.A. (o. 216°). S.G. 9 1.578.

aaβ-Tri-bromo-propionic acid

CH_Br.CBr...CO.H. [95°].

Formation.—1. By oxidation of acrolein bromido (Linnemann a. Penl, B. 8, 1097).— 2. From a-bromo acrylio acid and Br (Mauthmer a. Suida, M. 2, 99; Michael a. Norton, Am. 2, 17).

Properties.—Monoclinio prisms: a:b:c=1.83:1:315; $\beta=66^{\circ}$; m. sol. water, v. sol. alcohol and ether. On heating the Ba salt CII .: CBr2 is formed. Alcoholio KOII gives as.di.bromoacrylio aoid.

Salt.—BaA', xaq: needles. αββ-Tri-bromo-propionic acA

CHBr. CHBr. CO. H. [118°]. Prepared by heating as-di-bromo-acrylic acid for eight hours with conc. IIBrAq (Hill a. Andrews, Am. 4, 180; P. Am. A. 17, 133). Rectangular plate, v. sol. hot water, ether, and alcohol.—AgA': small rhombio

plates.—CaA'₂ 2aq. D1-bromo-p-d1-pr Tetra-bromo-propionio acid C₂HBr₄.CO₂H *i.e.* [48°]. From di-pr OHBr₂.OBr₂.CO₂H. [120°]. Prepared by tho combination of aβ-di-bromo-acrylic acid with Br. tables (from alcohol).

Triclinic prisms: v. e. sol. aloohol and ether. The Ba salt is decomposed by boiling water with formation of tri-bromo-ethylene. Alcoholic KOH gives tri-bromo-acrylic acid.

Salts. - AgA'. - KA' 2aq. - BaA', 1aq. CaA', aq (Mauthner a. Suida, M. 2, 107; Hill a. Mabery, P. Am. A. 17, 140; Am. 4, 266; 5, 251).

aβ - DI - BROMO - PROPIONIC ALDEHYDE CII2Br.CHBr.CHO. Acrolein dibromide. (c. 82°) at 5 mm. (Grima: x a. Adam, Bl. [2] 36, 136). From aerolein and Br (Aronstein, A. Stuppl. 3, 185; Henry, J. 7, 1112; Linnemann a. Ponl, B. 8, 1097). Pungent oil. Reduces Pehling's solution. Readily polymerises, becoming crystal-line [84°] in presence of HCl. HNG, oxidises it to di- and tri-bromo-propionio acids.

Tri-bromo-propionic aldehyde. A liquid combination of this body with propyl alcohol CBr_s.CII(OH)(OPr) appears to be formed on treating propyl alcohol with Br (Hardy, C. R.

79, 80ďí.

γ - BROMO - n - PROPYL - ACETO - ACETIC ETHER CH₂Br.CH₂.CH₂.CH₃.CO₂Et. Liquid. Insol. water, sol. alcohol and ether. Heavier than water.

Preparation .- 6 grms. of sodium are dissolved in 60 grms. of absolute alcohol and added to 32 grass. of aceto-acetic ether. The sodioaceto-acetic ether is then added to 80 grms. of tri-methyleno bromide and heated on the waterbath for half an hour; the yield is 75 p.c. of the theoretical.

Reactions .- By boiling with dilute acids it yields acetyl-butyl bromide and finally acetylbutyl alcohol Cit, CO.CH, CH, CH, CH, CH, OH. Alcoholic NII, eliminates HBr forming so called 'tri methylone-accto acctic ether' (Lipp, B. 18, 3277. V. also pp. 24 and 40 supra). β-BROMO-PROPYL ALCOHOL C.H. BrO i.e.

CH.Br.CH..CH. CH. Brombydrin of tri methylene glycol. (98°-1127, at 185 mm. S.G. 20 1.537. S. 17 at 15°. From tri-methylene glycol (s-dioxy-proxine) and HBr (Frühling, M. 3, 697).

ω-Bromo-isopropyl alcohol CH₂.CH(OH).CH₂Br (?). Bromhydrin of Propylene glycol. (1455-1486). From propylene oxido and HBr (Markownikoff, Z. 1870, 423).

aβ-Di-bromo-propyl-alcohol CH.Br.CHBr.CH.OH. Dibromide of allyl clocked (219°) (Weger, A. 221, 83). V. p. 134. alcohol. (219°) (Weger, A. 221, 83). V. p. 134. Methytderivative CH_Br.CH_Br.CH_OMo (185°) (Henry, B. 5, 455).

Ethyt dirivative CH2Er.CHBr.CH2OEt (194°).

s-Di-bromo-isopropyl alcahol

CH_Br.CH(OH).CH_Br. Gegeerin di-bromhydrin. (219°). S.G. 12 2.1. From glycerin and PBr. (Berthelot a. de Luca, A. Ch. [3] 48, 313; Reboul, A. Ch. [3] 60, 32). Also from glycerin and Br (Barth, A. 121, 349). V. also GLYCERIN.

DI-BROMO-PROPYL-AMINE Call, Bron i.e CH_Br.CIIBr.CH_NII_ From allylamino hydrochloride and Br. Oil. Salts .- B'HCl: needles. -B'2H2PtCl, (Henry, B. 8, 399).

BROMO - PROPYL - BENZENE v. BROMO-CUMENE.

Di-bromo-p-di-propyl-benzene C.H.Br.Pr. [48°]. From di-propyl-benzeno and Br (H. Körner, A. 216, 227). Needles or rectangular

BROMO-PROPYL-BENZOIC ACID v. BROMO-CUMINIO AOID.

DI-BROMO-PROPYL BORATE

B(O.CH2.CHBr.CH2Br) .. Hexabromide of ALLYL BORATE (q. v.). Oil.

BROMO-PROPYL BROMIDE v. DI-BROMO-PROPANE.

TETRA-BROMO-DI-PROPYL-CARBINOL v.

Tetwi-Bromo-HEPTYL-ALCOHOL.

BROMO - PROPYLENE 1, H, Br i.e. CII, CH:CHBr. (60°). S.G. 20 1.43. Formed, together with CII, CBr:CH, by cating propylene bromide with alcoholic KOII (Reboul, A. Ch. [5] 14, 479). One of the products obtained by boiling eβ-di-bromo-butyric acid with aqueous Na₂CO₂. HBr reunites with it forming chiefly propylene bromide, but the combination is very much slower than with its isomeride (48°). Alcoholio KOII gives allylene. Br forms CH2.CHBr.CHBr2 $(201^{\circ}).$

a - Bromo - propylene CII₂. CBr:CII₂. (48°). S.G. 2 1·39. Formed as above (Reynolds, A. 77, 122). Also by the action of alcoholic KOII on CH₃.CBr₂.CH₃ (Reboul); and by nnion of 11Br with allylene (Reboul, C. R. 74, 669). Oil. Readily combines with IIBr forming CH₂.CBr₂.CH₃. Brforms CH₂.CBr₂.CH₂Br (191°). Mercuric acetate in HOAc at 100 forms acetone.

Bromo-propylene CH.Br.CH.CH., v. ALLYL

вкомира р. 135.

Di - bromo · propylene C.H.Br., i.e.

CH₂:CBr.CH₂Br. (140°-143°). (a)-Epi-dibrom-hydrin. From CH₂Br.CHBr.CH₂Br and solid KOH (Henry, A. 154, 371; B. 14, 401) or sodinm in ether (Tollens, A. 156, 168). Converted by water at 130° into bromo-allyl alcohol.

Di-bromo-propylene CHBr:CII.CH,Br. (152°). S.G. 11 2.06. Formed, together with the preceding and propargyl bromide, from s-tri-bromopropane and KOII (Reboul, A. Suppl. 1, 230; Ilenry, B. 5, 186, 452). Alcohole AgNO, forms CHBr.CH.CH.NO.; AgOAo and potassium sulphocyanido also form \$\beta\$-bronio-allyl salts.

Di-bromo-propylene CH_,CBr;CHBr. \$\beta ttylene

di-bromide. (132"). S.G. 9.205. From allylene and Br (Oppenheim, Bl. [2] 2, 6; 4, 431; A. 132, 126). Also from CH₂, CHBr, CHBr, and AgOAc at 120° (Linnemann, A. 136, 56)

Tri - bromo - propyleno CII₃·CBr:CBr₂ (?). (184°) (O.); (c. 193°) (P.). From allylene tetra-promide and alcoholic KOAe (O.) or aqueous NaOH (Pinner, A. 179, 59).

Tri - bromo - propylene * CH.Br.Chr. CHBr. 12 2.53. Propargyl tribromide. From S.G. 12 2.53. propargyl bromide and Br. Non-volatile oil (Henry, B. 7, 761).

Penta-bromo-propylene C.HBr. From allyl bromide, Br, and I at 210°. Non-volatile oil (Mcrz a. Wcith, B. 11, 2243).

BROMO-PROPYLENE-GLYCOL v. GLYCERIN. BROMO-PROPYLENE-UREA C411,BrN2O i.e.

CO<NH>C₃H₅Br (?). [120°]. Formed by warming an aqueous solution of di-bromo-propylurea (Andreasch, M. 5, 40). Silky needles; sl. sol. cold water.—B'HCl: [143°].—B'HBr: [158°]. B'2H2PtCl

DI BROMO-PROPYL-MALONIC ACID C.H.Br.O. i.e. (CO.H).CH.CH.CH.CHBr.CH.Br. [121°]. From allyi-malonic acid and Br in CS. (Hjelt, A. 216, 58; B. 15, 624). Needles in stars | soluble brown prisms (Gerichten, B. 15, 1258).

(from ether). V. e. sol. ether, m. sol. water. Boiled with baryta, it forms di-oxy-propyl-

malonate of barium (q. v.).
BROMO - PROPYL - PHENOL v. BROMO-CUMENOL

BROMO-PROPYL-THIOPHENE

C,SH₂(C,H,)Br. (189°). Colourless oil. Formed by bromination of n-propyl-thiophone (Rnffi, B. 20, 1741).

Di-bromo-propyl-thiophene C,SH(C,H,)Br, (248°). Oil. Formed by bromination of propylthiophene with 2 mols. of bromine. By further bromination it is converted into tetra-bromothiophene (Ruffi, B. 20, 1741).

DI BROMO PROPYL-URÉA C.H.Br.N.O i.e. NH. CO.NH.CH. CHBr.CH.Br. [109°]. From allyl-urea and Br (Andreasch, M. 5, 38). Needles or leaflets, sl. sol. cold water. Decomposed by heating with water into the isomerio hydrobromide of bromo-propylene-nrca (q. v.).

BROMO-PYRIDINE C, NH, Br [3]. (174°i.V.).

S.G. 9 1.645.

Hormation. — 1. By brominating pyridine (Hofmann, B. 12, 990).—2. By the action of bromoform upon an alcoholic solution of pyrrol and NaOEt or upon potassium pyrrol in ether (Ciamician a. Silber, B. 18, 721; Ciamician a. Dennstedt, G. 12, 211; B. 15, 1173; Dancsi, G. 12, 150).

Properties .- Alkaline liquid, sl. sol. water. Reduced to pyridine by zinc and HCl.

Salts. B'aHaPtCl, 2aq: yellow monoclinic erystals, a:b:c = 1.207:1:1.188. $\beta = 107^{\circ} 7'$. B'HCl.

Di-bromo-pyridine C₅NII₃Br₂ [2:6]. [111°].

 $(222^{\circ}).$

Formation 1. From pyridine and Br (Hofmann, B. 12, 988). 2. Together with pyridine and mono-bromo-pyridine by the action of bromine on acctyl piperidine (Hofmann, B. 16, 587; cf. Schotten, B. 15, 421).-3. From tropidine hydrobromide and bromine at 170° (Ladenburg, A. 217, 148), di-bromo-methyl-pyridine being first formed, and cthyleno bromide being the other product .- 4. Formed by heating the tricarboxylio acid [206].

Properties.-Long flat pearly needles, insol. cold water, sl. sol. cold alcohol. Very weak base (difference from bromo-pyridine).

Salt.—B'₂H₂Cl₂PtCl₄: golden yellow needles (Pfeiffer, B. 20, 1349).

Methylo chloride C,NII,Br,MeCl. Formed by heating di-bromo-apophyllin (q. v.) with HCl (Anderson, A. 94, 358; Hofmann, B. 14, 1498; w. Gerichten, A. 210, 99). Moist Ag₂O liberates an alkaline hydroxido.—B'₂Mo₂PtCl₃.

Di-bromo-pyridine C₂NH₃Br₂. [165°]. Formed

by adding bromine to a boiling aqueous solution of pyridine sulphonic acid. Long white needles, Bogins to sublime at about 80°. Sol. water, alcolol, ether and bonzene.—B'₂H₂Cl₂PtCl₄2aq: large orange needles, sl. sol. water (Fischer a. Reimerschmid, B. 16, 1184; cf. Königs, B. 17, 589).

DI-BROMO-PYRIDINE-BETAÏNE

C,NH,Br, CH2 CO. Anhydride of di-bromeoxy-pyridyl-acetic acid. Formed by heating dibromo pyridine with chloro-acetic acid.—BHCl: colourless needles.—B₂H₂Cl₂PtCl₄: large b. Brome-apophyllenic acid.

Bromo-pyridine di-carboxylic acid U.NH.Br(CO.H). [165°]. Formed, together with oxalyl-authranilic acid, by exidation of (Py)-bromo-quineline with KMnO, Crystals. V. sol. water, alcohol, other, &c. It evolves CO. at its melting-point, yielding brome-pyridine-mono-carboxylio acid (brome-nicetinio acid) [183°] (Claus a. Collischonn, B. 19,2767).

Di.bromo-pyridine-tri-carboxylis acid C₃NBr₂(CO₂H)₃ [2:6:1:3:5]. [206° anhydrous]. Obtained by oxidation of di-bromo-s-tri-methylpyridine [81°] with KMnO. Flat plates (containing 4aq). V. sol. hot water, sl. sol. alcohol, nearly insol. ethe.. FeSO, gives a red coloura-tion. Heated to 165° it gives a sublimate of di-

brome-pyridine [111°]. Salts.—AgA"'aq: crystalline powder .-KII, A", Gaq: glistening colourless needles, sol.

htt, A. 2004; gustering conditions flowers, solution water.—Cu₂A" 2 aq: microcrystallino blue pewder (Pfeiffer, B. 20, 1347).

a.BROMO-a.PYRIDYL. (Py. 2). PROPIONIC ACID C₃II₃BrNO₂ i.e. C₃II₄N.CBrMo.CO₂II. From the corresponding oxy- acid and PBr₃ in

Salt.-(C.H.BrNO.)HAuBr, (Hardy a. Calmels, Bl. [2] 48, 232).

DI-BROMO-PYROCATECHIN C. H.Br. (OII) ... Di-mcthyl-ether CoH2Br. (OMe)... [93°]. Prepared by bromination of the di-methyl-ether of pyrocatechin (Tiemann a. Koppe, B. 14, 2018). Formed also by the action of Br on veratric acid. Colourless prisms. Sol. alcohol, ether and

Tri-bromo-pyrocatochin CaHBr3(OH)2

Mono-methyl-cther CallBr (OMe) (OII). Tri-bromo-guaiacol [102]. White lasted needles. Formed by bromination of guaiacol (Tiemann a.

Koppe, B. 14, 2017).

CaH,Br,(OII)2 Tetra-bromo-pyrocatechiu [1:2:3:4:5:6]. [193°]. Formed by bromination of pyrocatechin in chloroform solution. Colourless prisms (from acetic acid) or long fine needles (from alcohol). On exidation it yields tetra-brome-o-quinone C₆Br₁O₂. Bromine-water at 80° forms C₁₀H₂Br₁₀O [139°] (Zincke, B. 20, 1777; cf. Stenhouse, C. J. 27, 586; 28, 6; Hlasiwetz, A. 142, 251).

TETRA-BROMO.PYROCOLL C10H2Br4O2N2. Formed by heating pyrocoll with bromine at 100°. Small yellow needles. Insol. alcohol, ether, chloroform and toluene, almost insol. acetic acid. By boiling with KOII it yields dibromo-pyrrol-carboxylic acid (Ciamician a. Sil-

ber, B. 16, 2388).

TRI-BROMO PYROGALLOL C,H3Br,O, i.e. C.Br. (OH), Tri-bromo-pyrogallic acid. From tannin, glacial acctic acid and bromine at 100° (Steuhouse, C. J. 27, 586; 28, 7; Webster, C. J. 45, 207). From pyrogallol and Br (Hlasiwetz, A. 142, 250). Flat needles, so, het water. Bromine water converts it into 'xauthogallel' mine water converts it into knowledge and C₁₈H₄Br₁₄O₄ [122°] whence alkalis form C₁₈H₄Br₁₄O₅ [130°] (S.).

β-BROMO-PYROMUCIC ACID C₈H₄Br_{O5}.

αzharnkic acid [129°]. S.

Bromo-furfurane carboxylic acid [129°]. S. 1.26 at 20°. From either di-bromo-pyromucio acid [168°], or [192°] by reduction with zincdust and ammonia (Hill a. Sauger, A. 232, 58) On adding HOl the acid separates in matted

BROMO-PYRIDINE DI-CARBOXYLIC ACID

BROMO-APOPHYLLENIC ACID.

Bromo-pyridine-di-carboxylio acid

NH_Dr(OO_xH)___ [165°]. Formed, together

di-bromo-pyromucio acid [166°]. Dilute H_SO. gives bromo-fumarie acid [177°].
Salts.—BaA', aq. S.2·13 at 20°.—CaA', 3aq.

S. 1.77 at 20°. - AgA'. - NaA'. - KA'.

Ethyl ether EtA' [29°] (235° i.V.).

Amide.—C, II_BrO_NII_. [156]. Silky needles (from water).

(δ)-Brome-pyromucic acid C₃H₃B₁O₃. [184°].

S. 22 at 16°. Formation.—1. By the action of alcoholic KOH on the dibronide of pyroumeic acid (Schiff a. Tassinari, B. 11, 812; G. 8, 297). An isomoride [155°] said to be formed at the same time has not been observed by others. The dibromide of pyromucic other when sapouified also produces di-brome-pyronucio acid (Canzoneri a. Oliveri, G. 14, 172).—2. From pyronneic acid (20 g.) and Br (36 g.) at 100°, the yield being 50 p.o. of the theoretical (Hill a.

Sanger, A. 232, 46; B. 16, 1130).
Properties.—Pearly leaflets (from water); sl. sel. cold water, cold benzeno, and CIICla; m. sel, alcohol and other. Aqueens bromine forms fumaric, and the two di-brome-succinic acids, and di-brome-furfurane tetrabromide C4H2BraO

and di-promo-taritarano tetrapromine C₁H₂Dr₀O [111°]. Dilute IINO₂ produces malore acid. Salts.—KA'.—NaA'.—AgA'.—BaA'₂ 4aq (H. a. S.): pearly plates. S. (of BaA'₂) 3·47 at 18°.—BaA'₂ 2}aq (C. a. O.).—CaA'₂ 8aq: clumps of prisms. S. (of CaA'₂) 1·07 at 20°. Ethyl ether EtA'. [17°]. (235° i. V.). Amide C₃H₂DrO₂(NH₂): [145°]; needles

(from water).

Tetrabromide C.H.Br.O.: [173]; noedles (from HOAc).

Di-bromo-pyromucic acids. Pyromucic acid combines with bromine forming a tetrabromide which when boiled with alcoholic NaOH forms a mixture of two di-brome-pyromueic acids (βγ and \$5', (8)-brome-pyromucic acid and a tribromopyromucio acid (the latter from tetrabromide of (8)-bromopyromucic acid present in the crudo bromide). The calcium salt of the (βδ) acid is ppd. from dilute (1 in 30) solutions of the mixed acids by NH₃ and CaCl₂. Of the remaining acids, the (δ) acid is present in very small quantity, while the tri-bromo- acid is very sparingly soluble in hot water (Hill a. Sanger, A. 232, 67; B. 17, 1759; cf. Tönnies, B. 11, 1085; 12, 1202; Ca..zoneri a. Oliveri, G. 14, 177).

Constitution .- (Hill a. Sanger, A. 232, 97). Baeyer has proposed for py mucic acid

(I)
$$HC = C - CO_2H$$
 (II) $HC - C - CO_2H$ $HC - CH$

The two di-bromo-pyromucic acids would then be

$$(Ia) \begin{array}{c} \operatorname{BrC} = \operatorname{C-CO}_2H & \operatorname{BrC} - \operatorname{C-CO}_2H \\ | > O & (\operatorname{II}a) & | > O \\ \operatorname{BrC} = \operatorname{CH} & \operatorname{HC-OBr} \\ \end{array}$$

$$(\operatorname{III}a) \begin{array}{c} \operatorname{BrC} - \operatorname{C-CO}_2H \\ | > O \\ \operatorname{BrC} - \operatorname{C-CO}_2H \\ | > O \end{array}$$

$$(\operatorname{III}a) \begin{array}{c} \operatorname{BrC-O-CO}_2H \\ | > O \\ \operatorname{BrC-O-CH} \end{array}$$

(IIIb)

inasmuch as they are formed from the tetrabromide

HCBr-CBr.CO2H HCBr-CHBr

by removing 2HBr. One of these acids produces, on oxidation with IINO3, di-bromomaleïc acid, the other gives mono-bromo-maleïc acid. Neither of the di-bromo pyromucic acids derived from (II) could produce dibromo malcic acid, hence that formula is disproved. Formula (III) is unusual in form. Assuming formula (1) $(\beta \gamma)$ -di-brome-pyromucic acid is represented by (1a) while its $(\beta \delta)$ -isomeride is (Ib), and (δ) bromo pyromucic acid is CH = C—CO₂II

>0 , since on oxidation it gives CH = CBr maleïc acid, while (B)-bromo-pyronmeic acid is

 $CH = C - CO_2II$

since it may be got by reducing

CBr=CH either of the di-brome-pyromucic acids. It will be noticed that in the preceding argument it has been assumed that the bromo-pyremucic acids are similar in constitution to pyromucic acid itself.

(βγ)·di-bromo-pyromucio • acid C₃H₂Br₂O₃ [192°]. S. 21 at 20°. From the tetrabromide of pyromucio acid and alcoholic NaOII (see above). Short prisms grouped concentrically (from benzene) or bulky feathery crystals (from water). Sol. alcohol or other, en. sol. chleroform, sl. sol. CS., beiling water or light petro-leum (Hill a. Sanger, A. 232, 82,

Reactions .- 1. Dry bromine forms tribromopyromucic acid.—2. Bromine-vapour passed into an aqueous solution of the acid forms tetrabromo-furfurane, C, Br,O, [65°] and a little of a body C.IL, Br.O., [89°] (di-bromo-maleïcaldchyde).
3. Dilute HNO₃ on warming gives mucobromic and di-bromo-maleic acids .- 4. Zinc-dust and NH, from bromo-pyromucic acid [129°].

Salts.—AgA'.—NaA'2aq.—KA'.—BaA'23aq. S. 35 at 20°.—CaA'25aq. S. 117 at 20°. Ethyl ether.—EtA'. [68°]. Mosol. alcohol.

Amide C,HBr,O2NH., [196°]. Slender needles (from dilute alcohol). Iusol. CS₂ or light petroloum. Sl. sol. boiling water.

(\$\beta\$\begin{array}{c} (\beta\$\beta\$\beta\$\end{array}\rightarrow\beta\$\b

[168°]. S. 28 at 20°. From pyromucic acid and bromine at 100° (Hill a. Sanger, A. 232, 73). Also from the tetrabromide of pyromucic acid and alcoholic NaOH (see above). Small prisms, often in twins (from water). Very soluble in often in twins (from Auter). Very soluble in alcohol, ether, and boiling chloroform, less in benzene, sparingly in CS, or light petroloum.

Reactions .- 1. Aqueous bromine in the cold forms bromo-malcyl bromide.-2. Dilute HNO, forms bromo-fumaric acid, bromo-maleïc acid probably being an intermediate product of the

reaction.

Salts.—BaA'2aq. S. ·10 at 16°.—CaA'23aq. S. ·30 at 17°.—AgA'.—NaA'2aq.—KA'.

Ethyl ether EtA'. [58°]. (271° i.V. with decomposition). V. sol. ether, ohloroform, benzone, and boiling alcohol, m. sol. cold alcohol or CS2.

Bromide.-C, HBr, O2.Br. [46°]. (c. 154°) at 24 mm. One of the products of the action of

bromine upon pyromucio acid.

Amide C,HBr,O,NIL, [176°]. Silky necdles.

Tri-brome-pyromucio acid C,HBr,O,. [219°].
S. 072 at 10°. From the tetrabromide of (8)breme-pyror neic acid and alcoholic NaOH (Hill a. Sanger, ... 232, 91). Some tri-bromo-fur-furanc is also formed. Small needles united in clumps. V. sol. alcohol and ether, sl. sol. chloroform or benzene, nearly insol. CS, light petroleum or cold water.

Reaction: -1. Water and bromine form tetrabromo-furfurane [64°], thus: C₃HBr₃O₃ + Br₂ = C₄Br₄O + CO₂ + HBr.—2. Dilute HNO₃ forms di bromo maleie acid.

Salts.—BaA'₂3aq. S. (of BaA'₂) ·20 at 20°.—CaA'₂4aq. S. (of CaA'₂) ·56 at 20°.—AgA'.
—NaA'aq.—KA'aq.
Ethyl ether EtA'. [I04°]. Rectangular

prisms (from alcohol).

Amide C.Br.O.NH. [223°]. Slender needles.

Almost insol. CS., light petroleum or water, m. sol. ether, chloroform or benzeue, v. sol. alcohol.

 $Ita ext{-}BROMO ext{-}PYROTARTARIC ACID$

C₅H₇BrO₄ i.e. CII₂Br.CH(CO₂H).CII₂.CO₂H. Bromo methyl-succinic acid. [137°]. (c. 250°). From itaconic acid and cone. HBrAq at 0° (Beer, 1. 216, 79; cf. Fittig, A. 188, 73; Swarts, Z. Monoclinic crystals, v. sol. hot 1866, 722). water. Boiling Na, CO, Aq gives itaconie and itamalic acids; boiling water produces paraconic

Ethyl ether Et,A". (270"-275")

Citra · bromo - pyrotartario acid C.H.BrO. [148°]. From citraconic anhydride and conc. HBrAq at 0°. Also from mesaconic acid and fumisg IIBrAq at 140° (F.). Monoclinic crystals. Decomposed by heating alone or with Na₂CO₃Aq into methacrylic acid, CO,, and HBr. The silver salt on heating with water at 130° gives off allylene CH:C.CH₃ (Bourgoin, Bl. [2] 28, 459).

Bromo-pyrotartaric acid C.H.BrO, [204°]. White prisms. Formed together with bromocretonic acid by the action of Br on propanetricarboxylic acid CH3.CII(CO2H).CII(CO2H) (q. v.) (Bischoff a. Guthzeit, B. 14, 616).

Ita-di-bromo-pyrotartaric acid C, II, Br.O. From itaconio acid, Br, and water (Kekulé, A. Suppl. 1, 339). Crystals, v. sol. water, alcohol,

and ether.

Reactions .- 1. Sodium-amalgam reduces it to pyrotartaric acid. - 2. Moist Ag.O forms dioxy-pyrotartario acid. — 3. Boiling aqueous Na₂CO₃ forms aconic acid.

Anhydride C₂H₄Br₂O₃* [50°]. Formed by adding Br to a solution of itaconic acid in chloroform (Petri, B. 14, 1637).

Cira-di bromo-pyrotartaric acid CO₂H.CBr₂.CHMc.CO₂H. [150°]. S. 133 at 13°. From citraconic acid and Br (Kekulé, A. Suppl. 2,86; Krusemark, A. 206,1). Groups of neodles; v. e. sol. water, alcohol, and ether. Heated with water or aqueous Na₂CO₂ it yields propionic aldehyde, bromo propionio aldehyde, bromomethacrylic acid, and HBr. - CaA".

Anhydride C.H.Br.O. From citraconic

anhydride and Br; formed also by heating the following acid with water.

Mesa-di-bromo-pyrotartaric acid

CO2H.CHBr.CBrMe.CO2H. [194°] and [204°]. S. 81.5 at 13°. From mesaconic acid and Br on warming (Kekulé, A. Suppl. 2, 102; Fittig, A. 188, 86; 206, 1). Nodules. Heated with Na₂CO₃Aq it gives propionio aldehyde, two bromo-methaorylic acids, CO2, and HBr. Heated with water it gives propionic all chyde and bromo-oitraconic anhydride.

Di-bromo-pyrotartaric acid [102]. Formed by brominating pyrotartaric neid (Reboul a. Bourgoin, Bl. [2] 27, 348).

Di-bromo-pyrotartaric acid [128°]. Front propano tri-carboxylic acid and Br (Bischoff a. Emmert, B. 15, 1107).

Tri-bromo-pyrotartaric acid C₃H₃Br₃O₄. From pyrotartaric acid, Br, and water at 120° (Lagermark, Z. 1870, 299). Hexagonal prisms; sublimes above 240°.—Ag₂A".

TETRA-BROMO-PYROTRITARIC ACID

C,H,Br,O3. Tetra-bromo-uvic acid. Obtained by exposing powdered dry pyrotritarie acid to the vapour of drs bromme at the ordinary temporature. Large colourless crystals. V.sol. alcohol, ether, acctone, acctic acid, chloroform, benzene, and CS2, insol. water and petroleumother. The bromine is removed by alkalis and by aniline. By sodium amalgam in slightly acid solution it is reduced back to pyrotritarie acid. By the action of an excess of bromine at 100° it yields penta-bromo-pyrotritaric acid.

Tetra - bromide CiliBr,O3: Formed by dissolving tetra-bromo-pyrotritaric acid in an excess of dry bromine. Small prisms; v. sol. acetic acid, sl. sol. benzene, chloroform, and CS2, insol. water and ligroin. Decomposed by alkalis. Reduced to pyrotritaric acid by sodium amalgam (Dietrich a. Paal, B. 20, 1078).

Penta-bromo-pyrotritaric acid C_bH₂Br₅O(CO₂H). [c. 197°]. Obtained by heating the tetra-brome-derivative with excess of bromine at 100°. White glistening crystals. V. sol. ordinary solvents except water and ligroin (Dietrich a. Paal, B. 20, 1082).

TRI - BROMO - PYRROL - (a) - CARBOXYLIC ACID C₄NHBr₃(CO₂H). Tri - bromo - (a) - carbopyrrolic acid. Long colourless needles. V. sol. alcohol, ether, acetone, sl. sol. hot water, insol. petroleum-ether. The methyl ether is formed by bromination of the methyl ether of pyrrol-(a)carboxylic acid.

Methyl ether A'Me: [210]. Longslender needles, v. sol. ether and hot alcohol, sl. sof. benzene and petroleum-ether, insol. water (Cia-

mician a. Silber, B. 17, 1153).

Di-bromo-pyrrol di-carboxylic acid. Methyl ether C₄NHBr₂(CO₂Me)₂. [222°]. From dimethyl pyrrol di-carboxylato and Br (Giamician a. Silber, G. 17, 249). Long white needles; insol. water, sol. ether and hot alcohol. Converted by cold furning HNO₃ into C₄H₄BrNO₄ [c. 171°

BROMO-PYRRYL METHYL KETONE CH3.CO.C3H3BrNH. Pseudo - acetyl - bromo -

pyrrol. [108°]. Long colourless accdles; obtained by bromination of pyrryl methyl ketono (Ciamician a. Dennstedt, B. 16, 2354).

Di-bromo-pyrryl methyl ketone CH. CO.C. HBr. NH. [144°], white needles.

Formed by bromination of pyrryl methyl ke-

Tri-bromo-pyrryl methyl ketone C.H.Br.ON. [179°]. White silky needles. Sol. hot alcohol, ether, and aqueons alkalis, insol. water. Formed by the action of bromino upon pyrryl methyl ketone in aqueous solution (Ciamician a. Silber. B. 18, 1765).

Ponta bromo pyrryl methyl ketone C.H.Br.ON. [200]. Small white needles. Formed by bromination of the tri-bromo-derivative dissolved in acctic acid (C. a. S., B. 18, 1765) or of pyrref methyl ketone (C. a. D.).

BROMO PYRUVIC ACID C3H,BrO, i.e.

CH2Br.CO.CO2H. From pyruvic acid, Br, and water at 100° (Wichelhaus, B. 1, 265). Syrup.

Di-bromo-pyruvic acid CHBr. CO.CO.4H. [91°] (W.); [93°] (C.). From pyruvio acid (15g.), water (10g.), and Br (45g.) at 100° (Böttinger, B. 14, 1236; cf. Grimaux, Bl. [2] 21, 231; Clermont, Bl. [2] 19, 103; Wislicenus, A. 148, 208). Monoclinio efflorescent tables (containing 2aq); sol. water and other. Baryta converts it into tartronic acid. Benzeue and conc. H2SO4 form CHBr2.C(OH)Ph.CO2H (Böttinger, B. 14, 1235). - Di-bromo-pyruvic acid (1 mol.), mea (1 mol.) and conc. H₂SO₁ form di-bromopyrnvnreide C1H2Br2N2O2, whence brominewater forms tri-bromo-pyruvurin C.Br. N.O. II., a body which is decomposed by cold ammonia into bromoform and ammonic oxalurate. Ammonia converts di-bromo-pyruvureïdo into dibromo-pyruvuramide C₁H₂Br₂N₃O₂ which is docomposed by boiling baryta-water into NH, urea, HBr, tartronic acid, and amido-nraoil $C_4H_2N_3O_2$ (E. Fischier, A. 239, 185).

Tri - bromo - pyruvic acid CBra CO.CO2H. [90°], [101°, hydrated]. Formed, together with the preceding, by brominating pyruvic acid (Grimaux, Bl. [2] 21, 390). Also from lactic acid and Br. Lamida resembling naphthalene (containing 2aq); si sol. cold water. Decomposed by boiling water into bromoform and oxalio

Ethyl ether EtA'. [97]. Formed by adding Br to a solution of lactic acid in ether (Klimenko, J. R. 8, 125; Wislicenus, A. 143, 10).

DI-BROMO-PYRUVURAMIDE C,11, N, Br, O,. Di-bromo-pynuramide. [170°-180]. From di-bromo-pyruvureide and conc. NH₃Aq in the cold (Fischer, A. 239, 191). "Stender needles (from alcohol). V. sol. warm water, but slowly decomposed by boiling water. Decomposed by boiling baryta-water Into NII, nrea, HBr and tartronic acid, another portion forming amido-pracil.

DI-BROMO-PYRUVURFIDE C. 11 Br. N.O. acid (q. v.), urea, and conc. H.SO. (Fischer, A. 239, 188). Granular crystals (from HOAc), v. sl. sel. alcohol, water, and acids; sel. dilute alkalis. Decomposed by holing alkalis. Decomposed by heat above 280°. Its ammonium and guanidine salts are sl. sol. water.

TRI-BROMO-PYRUVURIL ANHYDRIDE Tri-bromo-anhydro-pyvuril. C,H3Br3N4O2. [180°]. Formed by heating tri-bromo-pyruvic acid and urea at 100° (Grimanx, A. Ch. [5] 11, 373). Light needles (from water).

TRI BROMO PYRUVURINE C'H,Br,N,O, Ureide of trii.e. CBr₂.CO.CO.NH.CO.NII₂. bromo-pyruvic acid. Tri-bromo-pyvurine. [247°]

From di-bromo-pyruvuroide and excess of bromine-water at 100° or HNO, (S.G. 1.4) (Fischer, A. 239, 189). Glittering platee, m. eol. boiling water and alcohol, v. sl. eol. ether. Decomposed even by cold alkalie into bromoform and ammonium oxalurate.

(B. 2)-BROMO - QUINOLINE C.H., BrN i.e. C.H., Br(C.H., N). Benz-bromo-quinoline. (278'). Liquid. Volatile with etect. Prepared by heating p-bromo-aniline with glycerin, nitrobenzene and H.SO.; the yield is 80 p.o.

benzeno and H.SO.; the yield is 80 p.o.
Salts.—B'HCl: small white needles.—
(B'HCl)_PtCl.: microscopic needlee (La Coste,

B. 15,558].

Bromo-quino'ine C.H.BrN. (270°). Yellowish oil. Prepared by bromination of quinoline. Perhape identical with the preceding.

Salts. — B'HCl: monoclinio prisms. (B'HCl)₂PtCl₄: fine orango-red needles.

Methylo-iodide C₀H₀BrNMe1. By the action of AgO on an aqueous solution of the iodide, a strongly alkaline solution of the hydrate is produced (C₀H₀BrNMeOH); this is transformed on etanding or warming, by splitting off H₂O, into the much more stable methylo-oxide.

Methylo-oxide (C_uH_cBrNMe)₂O. [147°]. This is also formed by the action of KOH on tho iodide. Colourless needles. Solublo in hot alcohol, sparingly in cold, very slightly coluble in water and other. Combines with acide very slowly (La Costo, B. 14, 915; 15, 188).

(Py. 1 or 2)-Bromo-quineline C.H. N: CH

or C.H. CII:CBr . (274° nncor.). Formed, to-

gether with propyl bromide, propylene bromido, quinoline hydrobromide, &c., by heating the propyle-bromide of quinoline-li-bromide to 170°-190°. Prepared by heating to 180° the hydrobromide of quinoline di-bromido: $C_0H_0Br_0N_1HBr_1HBr_1thequino-line-di-bromido is formed by the action of bromino on an ethereal solution of quinoline. Oil of aromatio smell resembling quinoline. On oxidation with KMnO, it yields oxaloxyl-anthranilic aoid <math>C_0H_1(CO_2H)_0NH_1CO_0C_0H$ and bromo-pyridine-di-carboxylic acid $C_0H_2Br_0N(CO_2H)_2$.

Salts.—B'HCl: needles or tablos; sublimes without melting.—B'HBr: foursided tables or prisms; sublimes at c. 190° without melting; sol. alcohol, sl. sol. cold water.—B'HNO₄*: [180° uncor.]; small concentric prisme.—B'₂H₂SO₄*: [183° uncor.]; small needles; diseociated by water.—B'₂H₃Cr₂O₃: [145°]; sparingly soluble flat yellow prisms (from hot water).—B'₂H₂Cl.PtCl₄: small orange-yellow needles.—B'₄AgNO₃: [173°]; needles (Claue a. Collisohonn, B. 19, 2763).

(B.1:4)-Di-bromo quinoline

CH:CBr C_bH₃N. [128°]. (a)-Di-bromo-quinoline.

Formation.—1. By bromination of quinoline by heating the hydrochloride with bromine at 180° (La Coste, B. 14, 917; 15, 191).—2. By heating di-bromo-aniline [1:4:5] with a mixture of glycerine, nitrobenzene and H₂SO₄ (Metzger, B. 17, 186).

Properties.—Dietils without decomposition. Volatile with eteam. Long white needlee. Almost

insol. water, v. sol. alcohol, ether, benzene and aqueous acids.

Salte. — B'HCl: small needles. — B'₂H₂Cl₂PtCl₂: fine yellow needles. — B'₂H₂Cr₂O₂: crange-red microoryetalline powder, decomposed by water into the base and CrO₂. The piorate forms long yellow needlee, decomposed by water.

Methylo-iodide B'MeI; Slonder red needlee. So, hot water, insol. ether and cold

alcohol.

Methyld-oxide B'2Me2O. Formed by the action of NaOH on the iodide. Microecopia needles.

(B. 2, 4) Di-brome-quinoline C₈H₂Br₂(C₈H₂N)₂ [101^o]. Slender colourloss needles. Volatifises undecomposed. Formed by heating di-bromo-aniline with glycerin, nitrobenzene and H₂SO₄. (B'HCl)₂PtCl₄ (La Coste, B. 15, 559).

Di-brome-quineline (probably B. 2: Py. 1) C_pII₁Br₂N. [124° uncor.] Formed by the action of bromine (2 mols.) upon quineline-(B. 2)- eulphonic acid (1 mol.) in cold aqueous solution. Long colourless needles (from ether). Sublimable. It is oxidised by KMnO, to brome-pyridine-di-carboxylic acid [165°] (Claus a. Küttner, B. 19, 2884).

Di-bromo-quincline tetrahydride C_bH_bBr₂N. [66° uncor.]. Formed by reduction of tetrabromo-quincline with sodium amalgam. Colourless tables. Volatile with steam. Sol. alcohol

and other, insol. water.

Salts.—B'HCl: [75°], acicular crystale.—
(B'HCl), PtCl, 2aq: yellow crystalline powder.—
B'HNO₃: [189°]: prisms.—B'H₂SO₄: white
plates, decomposes at 246° uncor.—B'H₂C₂O₄:
colourless tables, decomposes at 171° uncor.
(Claus a. Istel, 73. 15, 822).

Tri-bromo-quincline $C_uH_1Bc_3N$. [170° unoor.]. Formed by the action of bromine (3 mols.) upon an aqueous solution of quinclino-(B. 2)-sulphonic acid (1 mol.) at 100° . Long silky needles. Sl. sol. cold ether (Claus a. Küttner, B. 19, 2885).

Tri-brome quiuoline C_sH₄Br₃N. [175°]. From quinoline and Br (Lubavin, A. 155, 318). Silky needles; v. sol. hot alcohol. Possibly identical with the preceding.

Tri-bromo-quinoline C_sII₁Br₃N. [198° uncor.]. Formed by the action of bromine upon an aqueous solution of quinoline-(B. 4)-sulphonic acid at 100°. White felted silky needles. V. sol. ether and lot alcohol. Sublimable (Claue a. Eüttner, B. 19, 2882).

Tetra-bromo-quinoline c,H,Br,N [119° uncor.]. Long colourless needles or thick prisms. Insoluble in water. Formed by bromination of quinoline in CS₂ (Claus a. Istel, B. 15,820).

Hexa-bromo-quinoline C. HBr. N. [90°]. From pyridino (2, 3)-df-carboxylic acid, Br, and water (Weidel, A. 173, 55). Needles (from alcohol). Reduced to quinoline by sodium amalesm

(Wetter, 2. 173, 33). Reedies (from alconol). Reduced to quinoline by sodium amalgam.

(B. 4)-BROMO-QUINOLINE (B. 1)-GAP.
BOXYLIC ACID C_sH₁Br₁GCO_sH₁ [275°]. From bromo-amido-benzoie acid C_sH₂Br₁(NH₂).CO_sH [1:2:4] (10g.), glycerin (22·5g.), o-nitro-phenol (6g.), and H₂SO₄ (20g.) by heating for 5 houre at 160° (Lellmann a. Alt. A. 237, 313). White powder, v. sl. eol. water and ether, sl. sol. hot alcohol. Salt.—(HA'₁)₂H₂PtCl₄4aq.

(a) BROMO QUINOLINE JULPHONIC ACID C.H.(Br)N(SO.H). S. 08 at 22°; 9 at 100°. Short thin needles. Sl. sol. alcohol. Formed together with the B-acid by sulphonating bromo-

Salts.-A'2Mn4aq: small yellow needles.-A'Ag: spangles or needles.—A'K: short prisms, S. 1.37 at 17°.—A'NH,*: felted needles.—A'2Ba: nearly insoluble crystallino pp.—A'_Mg 10aq: colourloss plates.—A'_Zn 4aq: thir colourless needles (La Coste, B. 15, 1910).

(B) Bromo quinoline sulphonio a id C, H, N(Br)(SO, H). S. 15 at 22°; 2.75 at 100°. Short thick needles (containing aq). Formed together with the (a)-acid by sulphonation of

bromo-quinoline.

Salts.-A'K12aq: large tables, S. 17.25 at 22°.—A'Ag: colourless needles.—A'.Ba 2aq: sparingly soluble needles.—A',Mg 9aq: small needles.—A',Zn 9aq:six-sidedtables.—A',Mn Gaq: colourless tables (La Coste, B. 15, 1915).

BROMO QUINONE C. II. BrO2. [56°]. Formed by oxidising brome-hydroquinene with FegCla (Sarauw, A. 209, 106). Groups of needles, v. sol. alcohol, ether, and benzene, sl. sol. hot water. Ammonia gives a green colouration, turning

black on warming.

Di-bromo-quinone C₆H₂Br₂O₂. [188°]. Formed by oxidation of di-bromo-hydroquinono (S.; Benedikt, M. 1, 346). Small golden crystals, insol. water, sol. alcohol, ether, and benzene. Boiling KOIJAq gives di-bromo-di-oxy-quinone (dibromanilio acid).

Di-bromo-quinone C₆H₂Br₂O₂. [76°].

Formation. - Di - bromo - p - diazo - plienol, $C_4H_2Br_2< 0$ is converted by a boiling solution of calcium chloride into di-bromo-liydroquinone, C₆H₂Br₂(OH)₂. This solution is mixed with Fe₂Cl₄ and distilled, when the quinone passes over. The yield is small.

Properties.-Long, extremely slender needles, sol. in alcohol, ether, CHCl3, CS2, benzene, and alkalis. Pungent. May be sublimed (Böhmer.

J. pr. 132, 465).

Di-bromo-quinone C₆H_aBr_aO₂. [122°]. From tri-bromo-phenol and fuming HNO₄ at 0° (Levy a. Schultz, A. 210, 158). Yollow lamina (from dilute alcohol).

Di-bromo-quinone ? [88°]. From quereite and HBrAq at 160° (Prunier, A. Ch. [5] 15, 67). Three di-bromo-quinones are indicated by theory.

Tri-bromo-quinone C. HBr. O2. [147°]. Formed by oxidising tri-bromo hydroquinone in diluto' alcoholic solution (Sarauw, A. 209, 120). Golden leaflets (from alcohol); sol. alcohol, ether, and benzone. Alkalis give a green colouration, followed by separation of red prisms. Boiling cono. NaOH gives di-bromo-di-oxy-quinone and tri-bromo-hydroquincne. A tri-bromo-quinone [108°] is got by heating queroite with HBr (P.). A tri-bromo-quinone is also formed by reduction of tetra-bromo-quinono (Stenhouse, A. Suppl. 8, 20; cf. Horrmann, B. 10, 110).

Tetra-bromo-quinone C,Br,O2. Bromanil. Formation.—1. By treating phenol with Br and I (Stenhouse, C. J. 23, 10).—2. By boiling piorio acid with Br and water (Stenhouse, A. 91, 307).-3. From quinone and Br (Sarauw, B. 12, 680, A.209, 126). -4. A product of the action of Br | alcohol and HOAo; sl. sol. water.

and water on benzoic acid (Hübner, A. 143, 255), and on proteids (Hlasiwetz a. Habermann, A. 159, 820).—5. From tri-brano-phenol and HNO. (Losanitsch, B. 15, 474).—6. From di-oxy-dihydro-terephthalio (succinylo-succinic) acid and Br (Herrmann, A. 211, 341).-7. From (1,3,5,4)bromo-di-nitro-phenol by heating with Br (Ling, C. J. 51, 147).

Properties .- Golden lamine (from HOAc); sublimes as sulplar yellow erystals. Insol. water, m. sol. boiling aloohol, sl. sol. ether. HIAq reduces it to tetra-bronio-hydroquinone. Potash forms a greenish black solution turning

purplo.

Tetra-bromo ortho-quinone C.Br.O. [1:2:3:4:5:6]. [151°]. Obtained by oxidation of tetra-bromo-pyrocatechin in acetic acid solution with IINOs. It can also be prepared directly from pyrocatechin by adding bromine (10 to 12 pts.) to a boiling solution of the latter (I pt.) in acctio acid (20 pts.). Dark red thick prisms, tables, or transparent plates. V. sol. alcohol. ether, acetic acid, and benzeno, sl. sol. petroleum spirit. It is a powerful oxidising agent, being readily reduced to tetra-bromo-pyrocatechin. With aniline it gives a compound which crystallises in bluish black glistening plates or thick needles [173°] (Zineke, B. 20, 1776).

DI BROMO QUINONE CHLORIMIDE

NCl CaH2Br2 2:6:4 [8**0**°] Long vellow prisms. Prepared by adding a solution of chlorido of lime to an acidified solution of dibromo-amido-phenol [2:6:4:1] (Möhlau, B. 16,

2845) DI-BROMO-QUINONE-PHENOL-IMIDE

 $N < C_6H_4OH \\ C_6H_2Br_2O [4:2:6:1].$

Formation. By adding di-bromo-quinone chlorimido to an a kaline solution of phenol.-2. By oxidising an alkaline solution of di-bromoamido-pi-mol [2:6:4:1] and phenol with K2Cr2O,.

Properties .- Dark red prisins with metallie reflection. Sol. alcohol, ether and acctic acid with a magenta-red colour; insol. water.

Reactions .- On heating with HCl it is split up into quinone and di-bromo-amido-phenol. On reduction it yields di-bromo-di-oxy-di-phenyl-

amino HN C.H.Br.OH

Sodir m salt C.H.Br.ON(ONa): long blue prisms with golden-green reflection. Soluble in water and alcohol with a blue colour. Heated with an oxcess of aqueous NoOII the blue colour changes to red, but reappe its on cooling (Möhlau. B. 16, 2845).

BROMO-RESORGIN C.H.Br(OII)2. Formed by boiling bromo-di-oxy-benzoic acid with water for some hours (Zehenter, M. 8, 293). Groups of needles; v. sol. water and ether, m. sol. alcohol. Fc.Cl, colours the aqueous solu-tion bluish-violet, a red pp. being subsequently formed. Heated with water, K,CO, and SnCl. it gives resorcin and di-oxy-benzoio acid.

Bromo resorcin. Di-propyl derivative C₅H₂Br(OPr)₂. [71°]. Formed by brominating di-propyl-resorcin (Kariof, B. 13, 1679). Colourless silky needles; may be sublimed.

Di-bromo-resoroin C₆H₂Br₂(OH)₂. [93°]. Formed, together with 'di-bromo-mono-resor-[93°]. oin phthalein,' by heating tetra-bromo-fluorescein (eosin) with dilute NaOHAq at 140° (Baeyer, A. 183, 57; Hofmann, B. 8, 64). Formed also by boiling di-bromo-(1,3,2)-di-oxybonzoic acid with water (Zehenter, M. 2, 478; 8, 293). Needles (from water); m. sol. hot water, v. c. sol. alcohol and ether. Fe₂Cl₆ gives a transient violet colour.

Di-methyl ether C.H.Br.(OMe). [141°]. Slender needles. Insol. water, sol. alcohol and ether. Prepared by broamation of the dimethyl-other of resorein (Tiemann a. Parrisius, B. 13, 2365; cf. Honig, B. 11, 1041).

Di-bromo-resorcin (C_b|1_bBr₂(OH)₂, [112]. From Br and resorcin in CS₂ (Zehenter, M. 8) 293). Colourless needles (containing aq) (from water); m. sol. hot water. Fe Cl. gives a blno colour followed by a dark pp.

Tri-bromo-resorcin C.HBr3(OH)2-(Typke, B. 10, 1578). From resordin, Br, and water (Hlasiwetz a. Barth, A. 130, 357), or Br, and HOAc (Benedikt, M. 4, 227). Formed also by heating penta-bromo-resorcin with aldchyde or formio acid (Claassen, B. 11, 1439). Small needles; sl. sol. water, v. sol. alcohol.

Mono-acetyl derivative

C. IIBr. (OH)(OAc). [114°]. From mono-acetyl-

resorein and Br (C.); sol. hot water. Di-acetyl derivative C.J CaHBr3(OAc)2. [108°]. From ponta-bromo-resorcin and Ao₂O. Sol. hot water.

Mono-methyl ether C.HBr. (OH) (OMe). [104°]. From mono-metbyl-resordin and Br. Slender white needles, sol. alcohol and other, insol. water (Tiemann a. Parrisius, B. 13, 2364).

Tetra-bromo-resorcin C.Br. (OH), (C.); [167°] (B.). Formed by treating pentabromo - resorcin with H.SO, (Claassen, B. 11, 1440; Benedikt, M. 1, 366). Small needles Small Leedles (from alcohol).

Di - acetylderivative C6Br4(OAc). [169°]; v. sol. hot water.

Penta - bromo - resorcin C.Br.(OH)(OBr)? [114°]. Formed by adding an aqueous solution of resorcin to a cooled mixture of Br and water (Stenhouse, A. 163, 184). Dimetric crystals, a:c = 6076:1. V. sl. sol. water. Alcoholic AgNO, pps. more than two-thirds of its Br. At 160° it splits up into bromine and tri-bromo-resoquinono Callbr₃O₂ (Liebermann a. Dittler, B. 5, 1090; A. 169, 256). Converted into tribromo-resorcin by conc. HI, H.S. SnCl., warm alcohol, aldehyde, or formic acid (Benedikt, M. 1, 351; Claussen, B. 11, 1433). Boiling Ac.O gives di - acetyl - tri - bromo - resorcin. Anilino forms tri-bromo-aniline and tri-bromo-resorcin; phonol acts similarly (Benedikt, B. 11, 2168). Boiling dilute KOH produces bromoform.

Hexa-bromo-resorcin C₆Br₄(OBr)₂? [136°]. S.G. 165 3 188. Prepared by heating tetra-bromoresorcinol with excess of bromine. Monoclinic erystals: a:b:c = .983:1:1.687; $\beta = .85^{\circ}36'$. Decomposed by alcohol forming tetra-bromo-resorcin (Benedikt, M. 1, 365)

TRI-BROMO-RESOQUINONE Call BraO2 or C12H2Br6O4. Formed by heating penta-bromorosoroin at 160° (Liebermann a. Dittler, B. 5, 1090). Orange needles; insol. water, v. sol. alcohol and other. At 230° it gives off Br ETHILENE.

leaving amorphouf C₁,H₂Br₂O₄. Reduced by Sn and HCl to tetra-bromo-tetra-oxy-diphenyl (Benedikt, M. 1, 850; B. 11, 2170).

DI - BROMO - RESORCIN - PHTHALEIN SOoalled $C_{14}H_8Br_2O_5$ i.e. $CO_2H.C_6H_4.CO.C_6HBr_2(OH)_2$ [220°]. Di-bromo-di-oxy-benzoyl-benzoic acid. Formed, together with di-bromo-resoroin, by heating tetra-bromo-fluorescoin with diluto NaOHΛq (βaeyer, A. 183, 56). Plates, v. sl. sol. water.

BROMO RETENE v. RETENE.

BROMORICINELAIDIC ACID C18H33BrO. From the dibromide of ricinelaidic acid and alcoholic KOH. Oil. Alcoholio KOH forms an aoid [71°] (Ulrich, Z. 1867, 549).

BROMO - RICINOLEIC ACID C18H33BrO3. From ricincleic acid by successive treatment with Br and alcoholio KOH (Ulrich, Z. 1867, 546). Oil; converted by alcoholic KOH into ricinstearolic acid C18H32O3.

Di-bromo-ricinoleio acid C18H32Br2O3. From

ricinstearolic acid and Br. Oil.

BROMO-ROSANILINE v. ROSANILINE,

TETRA-BROMO-ROSOLIC ACID C20 H12Br4O30 From Br and rosolic acid in HOAc (Grache a. Caro, A. 179, 201). Lustrous green plates, insol. water. Its alkaline solutions are violet .- A"Ag2: dark violet pp.

Ethyl ether A"Et,: [110°-115°], soluble in alcohol, ether, and benzeno, insoluble in water (Ackermann, B. 17, 1627)

BROMO-ROSOQUINONE C,2H,Br,O2 i.e.

C₆II₂Br₂—O (?). Red and steel-blue crystals. C.H.Br.-O Prepared by the oxidation of tetra-bromo-phenolphthalcin (5 pts.) dissolved in H.SO. (250 pts.) with a mixtero of IJNO, (5 pts.) and H₂SO, (50 pts.).

Bromo-rosohydroquinone C12II6Br4O2 i.c.

C₆I1₂Dr₂OII

(?). Tetra-bromo-di-oxy-diphenyl.

[264°]. Sublimable. Prepared by the reduction of the corresponding quinono (Baeyer a. Schraube, B. 11, 1301).

BROMO-SALICYLIC ACID v. Baomo-o-oxynenzoic acid.

BROMO-SALICYLIC ALDEHYDE v. BROMO-O-OXY-BENZOIC ALDENYDE.

BROMO-STEARIC ACID C₁₈II₃₈BrO_{2.} [41°]. S.G. ²² 1 0053. From steario acid (7 pts.), bromine (4 pts.), and water at 135° (Ondemans, J. pr. 89, 195). Crystalline mass, insol. water, v. sol. alcohol and other. The silver salt heated with water forms stearidic acid C18H31O2.

Di-bromo-stearic acid C18H31Br.O2 oleio acid and Br (Overbeck, A. 140, 42). Oil. Alcoholic KOH forms bromo oleic and stearolic acids. Moist Ag₂O gives oxy-oleïe acid C₁₆H₃₄O₂

and di-oxy-stearic acid C_{1,} H_{3,}O₄.

Di-bromo-stearic acid C_{1,} H_{3,}D₄.

From claidic acid and Br. Reduced to claidio acidely sodium amalgam.

Tri-bromo-stearic aoid C18H33Br3O2. From

bromo-oleio acid and Br. Oil. Tetra-bromo-stearic acid C16H82Br4O2. [70°] From stearolic acid and Br. Laminæ (from

alcohol) BROMO STILBENE v. BROMO-DI PHENYL- BROMO-STRYCHNINE . STRYCHNINE. . C.H.Br C₈H₇Br

O.H. CHBr. Bromo-phenyl-cthylene. Formed by boiling styrene dibromide with alcoholio KOH or by heating it with water at 190° (Glaser, A. 154, 168; Radziszewski, B. 6, 493). Heavy pungent oil; decomposed by distillation. Converted by heating with water into phenyl-acetic aldehyde (Erlenmeyor, B. 14, 323)

a - Bromo - styrone C.II .CBr CH.

(220° i.V.).

Formation .- 1. From styrene dibromide and alcoholic KOAc at 160° (Zincke, A. 216, 290). - By boiling αβ-di-bromo-phenyl-propionic acid with water (Barisch, J. pr. [2] 20, 179; Fittig a. Binder, A. 195, 141).-3. From bromo-oxyphonyl-propionic acid and water at 200° ((1.).

Properties. - Oil, with pleasant odour of hyacinths. May be distilled. Does not readily give up its Br. Converted into acctophenoue by heating with water at 180° (Friedel a. Balsohn,

Bl. [2] 32, 614).

Di-bromo-styrene C₈H_aBr₂. (254°). From tri-exo-bromo-β-phenyl-propionic acid and water at 100° (Kinnicutt a. Palmer, Am. 5, 381). Oil.

Tri-bromo-styreno C,H,Br, From the preoeding and Br. Oil (K. a. P.).

BROMO-STYRENE DIBROMIDE v. Di-BROMO-ETHYL-BENZENE.

BROMO - SUBERIC ACID C.II. Br(CO.II). [103°]. Prepared, together with di-bromosuberio acid, by the action of bromine and phosphorus on suberic acid. Crystallino powder. Sol. alcohol and ether. By alcoholic KOH it gives suberconic acid (Gantiner a. Hell, B. 15, 142).

Di - bromo - suboric acid U.H., Br. (CO.H). [173°]. Formed by bromination of n-suberic acid. Glistening needles. V. sol. alcohol, ether, and hot water, v. sl. sol. benzene, chloroform, ligroin, and cold water. By heating with alcoholio KOH it gives di-ethoxy-subcric acid together with a small quantity of subercolic acid C₆H₆(CO₂H), (Hell a. Rempel, B. 18, 813). BROMO - SUCCINIC ACID C₄H₅BrO

CO₂H.CH₂.CHBr.CO₂H. [160°]. S. 192 at

15.5°.

Formation .- 1. By heating succinic acid (5g.) with Br (2½ c.c.) and water (40 c.c.) at 120° (Kekulé, A. 117, 125; Carins, A. 129, 6; Hell, B. 14, 892).—2. From succinic acid (5 g.), Br (2½ c.o.) and chloroform (5 c.c.) at 1600 (Orlowsky, J. R. 9, 277).—3. From succinic other and Br (Schacherl, B. 14, 637).—4. By the action of HBr on fumarie, tartaric, malic, and racemio acids (Kekulé, A. 130, 21; Fittig, A. 188, 88; Anschütz a. Bennert, B. 15, 613). 5. By decomposing its bromide with water (Volhard, A. 242, 153).

Properties .- Small prisms, v. sol. water. Its silver salt rapidly decomposes Moist Ag₀O gives malio acid. Sodium-amalgam produces succinio acid. Boiling water slowly forms fumario acid.

Anhydride C₄H₃BrO₃. [31°]. (137°) at 11 mm. From the acid and AcCl at 100° (A. a. B.). Decomposed by heat into HBr and maleïo anhydrido.

Methyl ether Me.A". (c. 134°) at 30 mm. Ethyl ether Et.A". (226°). Inflames the Cold aqueous or alcoholio NH, convert Vор. Į.

it into fumaramide. Aquoous NH, at 110° gives asparagine (Körner a. Menozzi, G. 17, 171).

Bromide C,H,Br(CD,Br)₂. Formed by adding Br (1100 g.) gradually to a mixture of succinic anhydride (300 g.) and amorphous P (36 g.) (Volhard, A. 242, 151). s-Di-bromo-succinic acid

CO.H.CHBr.CHBr.CO.H. S. 2-04 at 100°.

Formation. By heating succinic acid (42 g.) with Br (11 c.c.) and water (12 c.c.) at 180° (Kekulé, A. 117, 123; Suppl. 1, 131; Bourgoin, B. [2] 19, 148) — 2. From fumerio acid and Br (K.; Baeyer, 13, 18, 676).

Properties. - Opaque prisms, sl. sol. cold

water, v. sol. alcohol and other.

Reactions .- 1. Sodium amalgam reduces it to succinic acid.—2. Boiling water converts the sodium salt into hydro-sodic bromo-malate, the Isa salt into hydro-bario bromo-maleato and barium racemate, the silver salt into inactive tartaric acid, and the acid itself into IIBr and bromo-maleïe acid.—3. Water at 140° gives iso-bromo-maleïe acid.—4. NII₃ gives brome-amido-succinic acid.—5. Reduction in acid solution gives fumaric acid (Ossipoff, Bl. [2] 34, 346) .-6. Heating with thio-urea gives fumario acid (Neucki a. Sieber, J. pr. [2] 25, 79). Salts.—(NII₄)₂A".—Na₂A" 4aq.—Ag₂A".—

CaA" 2aq.

Mono-methylether MoliA": decomposes about 215° .-- NaMeA" 4aq (Claus, B. 15, 1814).

Mono-ethyl ether EtilA" [275°].

KEtA" 1;aq. 'NaEtA' 2aq.—AgEtA' 1;aq (C),

Mathyl ether McA". [62']. Mono-symmetrical crystch... Prepared by the action of

bromine on methyl fumarate.

Ethyl ether Et, A". [58°]. (K.; A.); [68°] (Lehrfeld, B. 14, 1820). Rhombio crystals. Prepared by the action of bromine on ethyl fumarate. On heating to 170° it decomposes into bromo a deie ether and HBr (Anschitz, B. 12, 2281) Aniline converts it into C₂H₂/NPhH₂(CO₂Et), [145°] (Lopatino, C. R. 105, 250).

Methyl-ethyl other McEtA" [63°] (C.). Chloride C.H.Br.2O.Cl. [63]. From Br and succinyl chloride or funnaryl chloride

(Perkin a Doppa, C. J. 13, 102; K.).

Amic acid CO.H.C.H.Br. CO.NH. Unstable crystals (C.; Michael a. Wing, Am. 6, 421).

AniI:de(?)

NHPh. 50 CHBr CHBr.CO.NHPh. From the anilide of fumaric acid and bromine (Anschütz a. Wirtz, A. 239, 138; Am. 9, 240). White powder, does not melt below 300°.

Phanyl-imide 1 ?h:(C,O,):C, II,Br, [159°]. From the pheny! imide of malere aoid (maleanil) in chloroform by adding Br (A. a. W.)

Iso- (or allo)di-bromo-succinic aoid

CHBr—C(OH), CHBr - CO CO2H.CH2.CBr2.CO2W or <

(ef. Anschütz, A. 239, 181). [160].

Formation.-1. From maleic acid and Br (Kekulé, A. Suppl. 2, 89) .- 2. Together with its isomerido, by heating bromo-maleio anhydride with HBr, or succinio acid with water and Brat 140° (Franchimont, B. 6, 199; Bourgoin, B. 6. 624). -3. From (8)-brome-pyromucio acid, Br. and water (Hill a. Sanger, A. 232, 53).

Preparation.—By dissolving its anhydride in water (Piotet, B. 13, 1670).

Properties.—Large crystals; more soluble in water than its isomeride. At 180° it gives off HBr, bromo-fumario acid being formed.

Reactions .- 1. Boiling water converts tho acid and its Ba salt into bromo-maleïe acid, but the Ag salt into racemio actd.—2. Moist Ag₂O gives pyruvic acid (Beilstein'a. Wiegand, B. 15, 1499).—3. Sodium amalyam produces sucoinio

The Di-methyl ether M"Me, and the Di-ethyl-ether A"Et, are oily liquids, insol. water (Piotet, B. 13, 1670).

Anhydride C2H2Br2.C2O3 [32°]. Propared by heating maleic anhydride with bromine at 100° (Pietet, B. 13, 1669). Colourless tables. Has a great affinity for water, with which it forms

isc-dibromo-succinio soid. On heating to 100° it evolves HBr forming bromo-maleic anhydrido. Tri-bromo-succinic acid

CO2H.CBr2.CHBr.CO2H. [137°]. From bromomaloïo or bromo-fumario acid and Br (Petri, A. 195, 69). Deliquescont needles; boil converts it into di-bromo-acrylic acid. Deliquescont needles; boiling water

BROMO-SULPHI-BENZOIC ACID $C_eH_3Br(SO_2H)(CO_2H)$ [4: 2 or 3:1]. [238°-245°]. From $C_eH_3Br(SO_2CI)CO_2H$ by treatment with alcohol and zine-dust (C. Böttinger, A. 191, 24) .--BaA".—BaH,A", 2aq.—CaH,A", 8aq. BROMO-SULPHI-BENZOIC ALDEHYDE

CoH2Br(SO2H)CHO. [131°]. One of the products got by reducing, by zinc-dust and alcohol, the mixture of chlorides got by acting on C₆H₂Br(SO₃Na)(CO₂Nc) by PCl₃. It is formed from C.H.Br(SO₂Cl)(COCI) present in the mix-ture. Salt.—BaA', 5aq. BROMO-SULPHŌ-BENZOIC ACID

C₆H₃Br(SO₃H)(CO₂H) [2:3 or 5:1]. From tho corresponding bromo-tolueno sulphonio acid by chromio mixture (Retsohy, A. 169, 45). KHA" ½aq.—BaA" 2aq.—PbA" 2aq.

Bromo-sulpho-benzoic acid C₆H₃Br(SO₃H)(CO₂H) [1:3:5]. From m-bromobenzoio acid and SO; (Hübner a. Upmann, Z.

[2] 6, 295; Rocters van Lennen, Z. [2] 7, 67; Böttinger, B. 7, 1779). Delicate deliquescent needles. Potash-fusion converts it into s-dioxy-benzoic acid.

Salts .- NaHA". - Ag.A". - CaA" 1]aq. BaA" 2 aq.—BaHA" aq.—CuA".

Brome-sulpho-benzoio acid C.H.Br(SO.H)(CO.H) [4:2:1]. From bromotoluene o-sulphonic soid by ehromic mixture (Weiss, A. 169, 26).—KHA".—CaA".—BaA": V. sol. water.

Imide C_eH_sBr < SO₂ NF. Bromo-benzoic sulphinide. [217°]. From (4, 1, 2)- bromotoluene sulphamide and KMnO. Also from tho acid K salt by successive treatment with PCls and NH_a (Remsen a. Bayley, Am. 8, 229). Long needles (from water); v. sol. alcohol and hot water, v. sl. sol. HClAq. Sublimes at 200°. Its taste is extremely sweet at first and then extremely bittor. — Ba(C,H₃BrNSO₃)₂7½aq. — Ca(C,H,BrNSO₂), 71aq. — AgC,H,BrNSO₃. — C,H,(C,H,BrNSO₂): [199°]; formed by successive treatment with PCl, and alcohol.

Bromo-sulpho-bensoio acid C_eH_eBr(SO₂H)(CO₂H) [4:3:1]. Formed by oxi- | C_eH_eBr(CO₂H)₂ [2:1:4]. [306° cor.],

dation of the corresponding brome-toluene sulphonic acid (Hässelbarth, A. 169, 12).—KHA" aq.—BaA" aq.—PbA" 2aq.

Brome-sulphe-benzeic acid

 $C_0H_3Br(SO_3H)(CO_2H)$ [4: 2 or 3:1]. Probably identical with the preceding. From p-bromo-benzoio acic and fuming H.SO, hoated for 8 hours at 13 to (Böttinger, A. 191, 13). Matted

Holdis at 13 (Bottinger, A. 191, 15). Matted needles, v. s. l. water.

Salts.—FaltA" 2aq.—Ag.A"3aq.—BaA"8aq.

—BaILA" 24 q.—CuA" 3aq.—PbA" 7aq.

Chloride C.H.Br(SO,Cl)CO,H. [197°] (with decomposition). Needlos (from ether). M. sol. cold ether, which separates it from another eliloride.

Acid other C,H,Br(SO,Et)(CO,H). From the chlorido and alcohol.

Amie aeid C_BH₃Br(SO₂NH₂)CO₂H. [230°]. BaA', 12aq.

Amic acid CaH3Br(SO3H)CONH2. Amie ether C.H.Br(SO,Et)(CONIL) [128°].

Bromo-di-sulpho-benzoio acid C,H,Br(SO,H),CO,H. From p-bromo-toluene disulphonio acid and boiling fuming HNO, Kornatzki, A. 221, 196). KaA'"aq.-Ba₃A‴₂12aq.

Chloride. [151°]. Trimetrio tables from ether).

Amide. [above 260°]. Small prisms in stars. BROMO - SULPHO - PHENYL - PROPIONIC

ACID C.H.BrSO, i.e. [4:3:1] C₆H₃Br(SO₃H).CH₂.CH₂.CO₂H. Propared by the action of fuming sulphurie acid on pbromo-phenyl-propionic acid (Göring, C. C. 1877, 793, 808). Non-deliquescent rhombic plates (containing 2 laq). a:b:e = 1:3013:1:0.7831.

Salts. — NaIIA" 3aq. — BaA" 2aq.—

monoclinio crystals: a:b:e=0.7062:1:0.9774: $\beta = 86^{\circ} 45'$.

DI-BROMO-SULPHO-PYROMUCIC ACID

 $\begin{array}{c} CBr = C - CO_2H \\ | > 0 \end{array}$ Di-bromo-sulpho-furfurane- $CBr = C - SO_3H$

carboxylic acid. Formod by sulphonation of di-bromo-pyroniucio acid [192°] with fuming H.SO.. By the action of bromine upon its barium salt, di-bromo-maleïo acid is formed. By zino-dust and aqueous NH, it is debrominated, yielding sulpho-pyromueio acid. A"Ba 5aq: easily soluble long fino necdles (Hill a. Palmer, B. 18, 2096).

BROMO-SULPHYDRO-BENZOIC ACID

C₆H₃Br(SH)CO_.H(?) [256°] (U.); [243°] (L.). From the chloride of sulphonated m-bromebenzoic acid by tin and HCl (Upmann, Z. 1870, 295; Van Lennen, Z. 1871, 67). Needles, insol. water. Reduced by sodium-amalgam to CoH,(SH)CO2H. 10dine converts its Na salt into sn acid [130°].
Salts.—ZnA'₂.—PbA'₂.—BaA'₂.
Bromo-sulphydro-benzoio acid

C₅H₃Br(SH)CO₂H. [194°]. From the chloride of (1, 3, 5)-bromo-sulpho-benzoic acid by tin HCl (Frorichs, B. 7, 795). PbA', 3aq.

BROMO-TEREPHTHALIO AOID Needles

containing aq (Fis.) or anh drous (Fil.). Prepared by exidation of bromo-toluio acid with KMnO₄ (Fischli, B. 12, 619), by exidation of bromo-oymene (Fileti, G. 16, 286), or of p-phenyl-toluene [129°] (Carnelley a. Thomson, C. J. 51, 88). It gives a sublimate anhydride) [245°].

Salts.-K2A": needlos.-Ag2A aq. White

insoluble flocculent pp.—A"Cu: light blue pp.

Chloride C₆H₃Br(COCl)₂. (34) ° cor.).

Amids C₆H₄Br(CONH₂)₂. [270°]: insoluble needles.

CaHaBr(CO.OEt)2. Methyl ether (Fis.); [52°] (Fil.). (above 300°). Needles.

Di-bromo-terephthalic acid C.H.Br. (CO.H), [6:3:4:1]. Formed by oxidation of di-bromo-ptoluio acid [195°] with KMnO, (Schultz, B. 18, 1762) or of di-bromo-cymeno with dilute IINO, (Claus a. Wimmel, B. 13, 902). Laminæ (from HOAc); does not melt below 320°.

Salts .- A"Ca 4aq: easily soluble microscopio needles. - A'Ba 2aq and A'Ba 5aq: mioroscopio needles.

Ethyl ether A"Et2. [121°]. (c. 335°). Pearly plates.

o-Di-bromo-terophthalic acid.

Hexa - hydride C₆ $H_8Br_2(CO_2H)_2$ [2:3:1:4]. Di-o-bromo-hexa - hydro - benzene - di-p - carboxylic acid. Formed by direct combination of tetra-hydro-terephthalic acid with Br in the cold. Granular orystals (containing aq). Nearly insol. cold water, sparingly in hot. By Ag₂O it is converted into an acid (probably C.H. (OH) (CO.H).) which by treatment with bromine yields tetrabromo-pyrocatechin (Baeyer, B. 19, 1808).

DI - BROMO - TETRADECANF, Tetradecylene bromide. [0°]. Colourless liquid. Formed by addition of Br to tetradecylene (Krafft, B. 17, 1372).

 C_4Br_3S HEXA-BROMO-DITHIËNYL r2550 C₄Br₃S

Formed by heating an acetic acid solution of dithienyl with an excess of bromino (Nahnsen, B. 17, 2198). Small needles. V. sol. hot benzene, v. sl. sol. cold benzene and hot alcohol.

TRI-ω-BROMO-DI-THIËNYL-ETHANE

CBr₂.OII(C₄II₃S)₂. [102°]. Obtained by adding H₂SO₄ to a mixture of thiophene and bromal dissolved in acctic acid. Small pyramids. sol. ether, CS2, and hot alcohol. With isatin and H2SO4 it gives a violet-red colour (Peter, IL 17, 1844).

DI-ω-BROMO-DI-THIËNYL-ETHYLENE CBr₂:C(C₄H₂S)₂. Formed by boiling tri-bromodi thlënyl-ethane with alcoholio KOII, or better KCN (Peter, B. 17, 1344). Colourless oil.
Volatile with steam Gives a violet-red colour
with isatin and H.SO.
BROMO-THIENYL METHYL KETONE

C.SH.Br.CO.CH. Bromo-acetothienone. [94°]. Formed by the action of acetyl chlorido upon mono- or di-bromo-thiophene in presence of Stout colourless needles. alcohol, less in cold. Very volatile with steam. By alkaline KMnO, it is oxidised to bromo-thio-

phene-carboxylio acid [140°].

Phenyl hydrazide
C₄SH_xBr.C(N_xHPh).CH₃: [122°]; tables; sl. sol. alcohol (Gattermann a. Römer, B. 19, 689).

BROMO-THIO-CRESOL v. BROMO-TOLYL MER-OAPTAN.

DI-BROMO-THIOHYDANTOÏN C.H.ON.SBr. Formed by the action of bromine on a solution of thiohydantoin in aqueous HCl (Muldor, B. 8, 1263; Kramps, B. 13, 789). Colourless crystals. Insol. oold water, sol. alcohol and ethor. Decomposed by he, water.

*BROMO - THIO - OXY - BENZOIC ACID v.

BROMO-SULPHYDRO-BENZOIC ACID.

(a) - BROMO THIOPHENE C.SH.Br. 'B'-Bromo-thiophene. (150°). S.G. 2 1 652. Colourless liquid. Formed by bromination of thiophene (Meyer, B. 16, 1472). Isolated from the ernde di-bromo thiophene obtained by fractional bromination of benzene that contains thiophene. By EtBr and Na it is converted into 'B'-ethylthiopheno Schleicher, B. 18, 3015).

Di-bromo-thiopheno C.SH.Br.. (211° cor.). S.G. $\frac{23}{23}$ 2.147. Colourless oil. Formed by dropping bromino into thiophene cooled with water. Prepared by fractional bromination of benzene that contains thiophene. With isatin and H.SO, it gives a deep-blue colour (Moyer, B. 16, 1469; Meyer a. Stadler, B. 18, 1488).

Tri-bromo-thiopheno C.SIIBr., [29°]. (260° Formed by further bromination of dicor.). bromothiophene. Long white glistening crystals. V. sol. hot alcohol and other, sl. sol. cold alcohol. Gives the indophenine reaction. By sulphonation and debromination it yields thiophono-(3). sulphonic acid (Rosenberg, B. 18, 1773).

Tetra-bromo-thiophene C₁SBr₁. [112°]. (326° cor.). Long white needles. Formed by further bromination of di-brome-thiophene (Meyer a.

Kreis, B. 16, 2172).

BROMO-THIOPHENE-CARBOXYLIC ACID C,SH₂Br(CO₂II). [140°]. Formed by exidation of bromo-thienyl methyl ketone with alkaline KMnO. Colour oss glistening needles (from water). Sublimes in pearly spikes. M. sol. hot water, learly insol. cold water (Gattermann a. Römer, B. 19, 690).

Di-bromo-thiophone-(a)-carboxylic acid

C4SIIBr2.CO2H. Di - bromo - thiophenic acid. [222]. Obtained by bromination of (a)-thiophene-carboxylic acid (q,v,). White monoclinic needles. Sublimes on hating. V. sol. alcohol and ether, sl. sol. hot water, insel. cold water. Sparingly volatile with steam. Isatin and ILSO yield a dity-green colouration quickly becoming brown. A solution of the ammonium salt gives white pps. with AgNO₃, Pb(OAc)₂, H₅NO₃, and SnCl₂; yellow pp. with Fe Cl₆; and greenishwhite pp. with CuSO.

Salts.—A'Ag: white early pp. becoming crystallino. — A'K: easily soluble crystals. — A'2Ba 32aq: white needles, v. sol. hot water, sl.

sol. cold.

Chloride C.SIIB .COCl: (250°-270°); ^esilky needles.

Amide CaSHBrg. CONHg: [167°]; fine white felted needles; v. sol. alcohol and other, sparingly in hot water.

Methyl cther CaSHBr. CO.Me: [80°]; white needles (Peter, B. 18, 543; Bonz, B. 18, 2308)

DI-BROMO-THIOPHENE-SULPHONIC ACID C.HBr.S.SO.H. Formed by sulphonating dibromo-thiophene. By sodium-amalgam it is

g s 2

reduced to thiophene - 'B'-sulphonic acid.-PbA', 5 inq: small crystals, sol. hot water. Chloride C,HBF,S.SO,Cl: [33°]. Amide C,HBF,S.SO,NII,: [147°]; felted

needles; sparingly soluble in water (Langer, B. 17, 1566; 18, 553; Rosenberg, B. 18, 3030).

Di-brome-thiophene-di-sulphonic acid C₄SBr₂(SO₄H)₂. Obtained he boiling the anhydride with alkalis. It very readily splits off H₂O with conversion into the anhydride.

Salts. - Na2A" 3aq: very solublo ncedles.—(NII₁)₂A" aq.—PbA": plates, sol. hot water.—BaA" aq: spavingly soluble white glistoning spikes.

Anhydride C,SEc, SO2 O: white glistening plates, v. sol. alcohol and benzene, insol. water and ligroin. Obtained by the action of fuming sulphuric acid (4 vols.) on di-bromothiophene (1 vol.). Chloride C.SBr₂(SO₂Cl)₂: [220°]; glisten-

ing white needles; sol. ether.

Amide C₄SBr₂(SO₂NH₂)₂: [o. 270°]; crystalline powder; nearly insol. water (Langer, B. 17, 1569; 18,554; Rosenberg, B. 18, 3030).

Tri-brome thiophene-sulphonic acid C4SBr3(SO4II). Formed together with the anhydrido by sulphonation of tri-bromo-thiophene [29] -BaA'2aq: sparingly soluble white warty crystals.

Anhydride (C,SBr2.SO2)2O: [116°]; volatile with steam; white solid; v. sol. alcohol and

ether, v. sl. sol. water.

Chloride C₃SBr₄(SO₂Cl): [126°]; needles. Amide C₄SBr₄(SO₂NIL): needles (from water) (Rosenberg, B. 18, 1774, 3028).

DI-BROMO-THIOPHENIC ACID v. DI-nROMO-THIOPHENE-CARBOXYLIC ACID.

BROMO - THIO - PHENOL v. BROMO-PHENYL

TETRA BROMO THIOPHTHENE C.S.Br. i.e. CBr:CBr

C = C . [172°]. Formed by bromination

CBr:CBr of thiophthene. Long white needles (from benzene). V. sol. hot benzene, sl. sol. alcohol (Biedermann a. Jacobsen, B. 19, 2117).

BROMO - THIOTOLENE v. BROMO-METHYL-THIOPHENE.

BROMO-THIOXENE v., Bromo-dy-Methyl-THIOPHENE.

m-BROMO-THYMOHYDROQUINONE

. C₆HMcPrBr(OH)₂ [1:4:5:3:0]. [53°]. Formed by the action of cone. HEr upon thymogninon at a low temperature. Colouriess needles (Schniter, B. 20, 1318). Formed also by reduction of the corresponding quinone with SO2 (Mazzara a. Discalzo, G. 16, 195). Changes spontaneously to a substance melting at 37° (M. a. D.).

Di-acetyl derivative C, HMcPrBr(OAc), 4-1 [91°]. From thymoquinono and AcBr (Schulz,

B. 15, 657).

Di-breme-thymo-hydroquinene

Di-acetyl derivative C.McPrBr. (OAc)2. From di-acetyl bromo-thymo-hydroquinono and Br (S.).

BROMO-THYMOLS and their ethyl derivatives appear to have been obtained by Paterno a. Canzoneri, G. 10, 233, Armstrong a. Thorpe, Brit. Assoc. Reports, 1875, 112; and Lallemand, A. Ch. [3] 49, 148.

BROMO-THYMOL SULPHONIC ACID C₁₀H₁₃BrSO₄ i.e. O₆HMePrBr(OH)(SO₃H). From potassium thymol (a)-sulphonic acid and Br.— KA' 12aq.—BaA'2 (Engelhardt a. Latschinoff, Z. 1871, 261).

1871, 261, 6 m-BROM D-THYMOQUINONE C₆HMePrBrO₇ [1:4:5:3:6]. [3:9] (M. a. D.); [45:9] (S.). Formed by oxidation of the corresponding bromo-thymo-hydroquinons with Fe₂Cl₈. (Histening yollow plates (Schitter, B. 20, 1318). From bromo-mide thymological properties and Mazzara a amido thymol and nitrous acid (Mazzara a. Discalzo, G. 16, 195). A crystalline bromotliymoquinone was got by Corstanjen (J. pr. [2] 3,55) in Frominating thymoquinone. Andresen (J. pr. [2] 23, 184) obtained a liquid isomeride by brominating thymoquinone chloro-imide.

Di-breme-thyme-quinene C₆Br₂McPrO₂ [74°]. The ethoreal extract from the product of the action of HBr on thymo-quinone-chloroimido (q. v.) is evaporated and the residue distilled with steam (Andresen, J. pr. [2] 23, 184). From thymoquinone and Br (C.). Lemonyellow plates (by adding water to the alcoholic solution).

o-BROMO-TOLUENE C,H,Br i.e. C,H,MeBr

[1:2]. (182°). S.G. 1°2 1·2031. S.V. 141·95 (Schiff, B. 19, 564).

Formation.—1. Together with p-bromotoluene, by brominating cold toluene, in the dark, in daylight, or with addition of iodine (Hibner a. Wallach, Z. [2] 5, 22, 138, 499; A. 154, 293; Dmochowsky, B. 5, 333; Kckulé, A. 137, 192; Belistein, A. 143, 369; Cannizzaro, A. 141, 198; Glinzer a. Fittig, A. 133, 47; 136, 301; Fittig, A. 145, 39; Rosenstichl a. Nikiforoff, Z. [2] 5, 635; Hübner a. Retschy, Z. [2] 7, 618; Lauth a. Grimaux, Bl. 1866, i, 347; 1867, i, 108; Körner, G. 4; Hübner a. Januasch, A. 170, 117; Longuinine, B. 4, 514; Reyman, Bl. [2] 26, 533; Schramm, B. 18, 607).—2. From o-toluidine by the diazo-reaction (Wroblewsky, A. 168, 171; Jackson, Am. 1, 93).—3. Together with naphthaleno, by the action of (a)-hrome-naphthalene on toluene in presence of Al₂Cl_e (Roux, Bl. [2] 45, 520).

Properties .- Oil.

Reactions .- 1. Converted by dilute HNO. into o-bromo-benzoic acid (Zincke, B. 7, 1502).-2. Sodium has no action at 15°.—3. Sodium and McI form o-xylene.—4. The copper-zinc couple has no action (Gladstone a. Tribe, C. J. 47, 448).

m-Bromo-toluene Collimor [1:3]. (184°). S.G. 24 1.401 (W.).

Formation. -1. From C6H3MeBr(NH2) [1:3:4] by the diazo- reaction (Wroblewsky, Z. [2] 7, 609; A. 168, 155; Grete, A. 177, 231).-2.
From the same brome p-to-uidine by successive conversion into C₆H₂(NO₂)MeBr(NH₂) [5:1:3:4], C₅H₃(NO₂)MoBr [5:1:3], C₅H₄(NH₂)Me [5:1], and C. IlcBrMo [5:1] (Wroblewsky, A. 192, 206).

Properties.-Liquid, even at -20°. Oxidised by chromic mixture to m-bromo-benzoic acid.

p-Breme-teluene C.H.MoBr [1:4]. [28:5°]. (185° i. V.) (Hübner a. Post, A. 169, 6). S.G. 🛂 1 411 (Kekulé, 🖊. 137, 192)

Formation.-Togother with o-bromo-toluene (q. 1.) by brominating toluene.

Properties .- Trimetric crystals.

Reactions.—1. OrO, rosms p-bromo-benzoio acid.—2. Sodium even at 15° forms ditolyl (Zinoke, B. 4, 396; Louguinine, B. 4, 514).-8. Sodium and MeI gives p-xylene.-4. The copper-sinc couple has no action (G. a. T.) .-5. Taken internally, it is excreted as p-broincbenzoio and p-bromo-hippuric acids (Preusse, H. 5, 63).—6. CrO₂Cl₂ forms C₆H₄Br CH(O.CrOCl)₂ (Etard, A. Ch. [5] 22, 241).

ω-Bromo-toluene v. BENZYL BREMIDE.

Di-bromo-toluene CoH, MeBr 12:2:3]. [28°]. From C₀H₂(CH₂)(NH₂)Br₂ [86°] by displacing NH₂ by H (Nevile a. Winther, C. J. 37, 431). Gives on exidation with HNO₃ di-brome-benzoic actl [146°-148°]. On nitration it gives a mono-nitro- derivativo [57°] which reduces to an amido- compound [53°].

Di-breme-teluene C.H.MeBr. [1:2:4]. From di-bromo-m-toluidino [75°] by eliminating NH., Also from C₆H₂(CII₄)(NII₂)Br [1:2:4] by diazo-perbromido reaction. Oil. On nitration gives di-bromo-nitro-tolucno [80°]. On oxidation with HNO, gives di-bromo-benzoic acid [169].

Di-bromo-tolueno C₅H₃MeBr₂ [1:2:5]. (236°). S.G. ¹⁴ 1813. Formed by brominating toluene (Wroblewsky, Z. [2] 6, 239). From acctyl-otoluidino by brominating, saponifying, and displacing NH₂ by Br by the diazo- reaction (N. u. W.). Also from acetyl-m-toluidino in the same Thus Coll3(CH3)(NH2)Br [1:2:5] and wav. $C_8H_3(CH_3)Br(NH_2)$ [1:2:5] give the same $C_8H_3(CH_3)Br_2$. Oil. On nitration it gives a nitro- compound [88°], which reduces to an amido- compound [85°]. Oxidised by diluto IINO₃, it gives di-bromo-benzoic acid [149°-153°]; this distilled with lime gives a solid, [86°], probably p-di-bronio-benzence.

Di-bremo-toluene C₆H₃MoBr₂ [1:2:6]. (216°). **S.G.** ²² 1:812. From di-bromo-*m*-toluidine [35°] (Wr.; N. a. W.) or from di-bromo-p-toluidino [88°] by diazo- reaction. Oil. On nitration it gives two nitro-compounds, [161°], and [2.80°]. On oxidation it gives di-bromo-benzoic acid

[150°-167°].

Di-bromo-toluene CaHaMeBr, [1:3:4]. (241°). S.G. 19 1.812. Formed in brominating tolueno containing iodino in sunlight (Jannasch, A. 176, 286). Also from bromo-p-toluidine by the diazoperbromido reaction (N. a. W.; Wr.). Oil. Gives a nitro- derivativo, [87°], reducing to an amidoderivative [98°]. By oxidation with CrO, it gives di-bromo-benzoic acid [233°]

Di-bromo-toluene CaHaMcBr. [1:3:5]. [39°]. (246°). From C₆H₂(CII₃)(NH₂)Br₂ [73°] and 4ts isomeride [46°] (N. a. W.; Wr.). With HNO₃ it gives two di-nitro- compounds, [158°] and [105°]. On oxidation with CrO, it gives di-bromo-benzoio

acid [208°-210"].

Di-bromo-tolnene C₆H₃MeBr₂. [108°]. existence of this body, said to be formed in brominating toluene (littig, A. 147, 39), is called in question by Novilo and Winther.

Other Di-bromo-toluenes are described as BENZYLIDENE BROMIDE and BROMO-BENZYL BRO-

Tri-bromo-toluene C_sH₂MeBr, [1:2:3:4]. [44°]. From C_eH(CH_a)(NH₂)Br₃, [97°], by nitrous gas (Nevile a. Winther, C. J. 37, 447). On nitration It gives a nitro-compound [107°].

Tri-brome-toluene C.H.MeBr, [1:2:3:5]. [53°]. From di-bromo-o-toluidine [40] by diazo-

perbromide reaction (N. a. W.). Long flat

Tri-bromo-toluene CaH2MeBra [1:2:5:6]. [59°]. From [1:3:2:5:6] CaH(CHa)(NH2)Br, [94°] by nitrous gas. On nitration it gives a nitro-compound [91°].

Tri-bromo-tolnene Call, McBr, [1:2:4:6]. [66°]. (290°). From tr-brono-m-toluidine [101°] by nitrous gas (N. a. W.; Wroblewsky, A. 168, 194). On nitration it gives a di-nitro-compound [c. 220°].

Tri-bromo loluene C. II. McBr. [1:3:4:5]. [89°]. From di-bromo-p-toluidine by diazo-perbromido reaction (N. a. W.).

Tri-bromo-tolucne C_11_MeBr_a[1:3:4:6].[112°]. From di-bromo-m-toluidine [75°] and from dibromo-toluidine [85°] by diazo perbromide reaction (Noa. W.).

Tri-bremo-teluene? [1503]. Formed by heating potassium tri-bromo-phenol with KOAo

(Pfankuch, J. pr. [2] 6, 103).

Tetra-brome-telucno C. HMeBr. [1:2:3:4:6]. [105°-108°]. From tetra bromo-m-toluidino [224°] by alcohol and nitrous gas. Also from tri-bromo-m-toluidine [100°] by diazo-perbromide reaction (Nevile a. Winther, C. J. 37. 449). Furning HNO, forms the nitro-compound [216°].

Tetra-brome-teluene C. HMcBr. [1:3:4:5:6]. [111°]. From tri-bronio-m-toluidine [96°], by diazo-perbromide reaction (N. a. W.). Its nitro-

compound melts at [212°].

Tetra-breme-telucne C.HMcBr, [1:2:3:5:6]. [117°]. From tai-bromo-m-toluidine [94°] by diazo-perbromido reaction. Its nitro-derivativo melts at [213"].

Penta-brome-telucne CaMeBr., [285°]. From tetra-bromo-m-toluidine by diazo-perbromido reaction (No ile a. Winther, C. J. 37, 450). From tolume Al₂Br₄, and Br at 0° (Gustavson, Bl. [2] 28, 347, B. 10, 971). Long needles (from benzene).

O-DROMO-TOLUENE SULPHONIC C.H. BrSO, i.e. C.H. McBr(SO, 11) [1:2:5?]. From o-bromo-tolueno by sulphonation (Hübner a. Post, A. 169, 31; cf. Dmochowsky, B. 5, 333). Also from brominated (1, 1, 3) p-toluidine sulphonic acid by displacement of NII, by H (Pechmann, A. 173, 212). Reduced by sodiumamalgam to toluene m-sulphonic acid.

Salts.— RA' laq.—NaA' laq.—BaA'₂ 2aq.—BaA'₂ 5.ja₁.—S. (of BaA'₂) :55 at 14°.—PbA'₂ 2aq.—S. (of PbA'₂) :52 at 18°.—PbA'₂ 3aq.

Chloride C.H.MeBr(SO.CI). [539]. Amide Calla MeBe (50) N 112). [1340]. o-Breme-telucne salphenic acid

CaHaMeBr(SOalt) [1:2:5]. From o-tolnidine by sulphonating and displacing NH, by Br (Pagel, A. 176, 294; Nevde a. Winther, B. 13, 1943; cf. Gerver, A. 109, 384). Reduced by sodiumamalgam to tolucue meulphonic acid.

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Salts.—KA'aq.—CaA'₂aq.—BaA'₂½aq.

1·2 at 25° (P.).—BaA'₂2aq.—BaA'₂9aq. S. 8·9

at 17·5° (G.).—PbA'₂2aq.—CuA'₂3aq.

Chloride [53°] (P.); [56°] (N. a. W.).

Amide [c. 136°] (P.); [147°] (N. a. W.).

This acid is probably identical with the preceding. o-Bromo-toluene sulphonic acid

C₈H₃MeBr(SO₃H) [1:2:4]. From o-toluidine psulphonic acid by the diazo- reaction (Hayduck, A.172, 206).— KA': nodules of minute needles.— BaA', 2aq.—PbA', 2 aq. Chloride C,H,MeBr(SO,Cl). [54°]. Amide C,H,MeBr(SO,NH,). [161°].

Bromo-toluene sulphonio acid

O.H. MeBrSO.H [1:3:5]. From bromo-o-toluidino sulphonio acid or from bromo-p-toluidine sulphonio aoid by displacement of NH₂ by H. Potash-lusion gives orein (Nyvile a. Winther, B. 13, 1944; C. J. 41, 420).

Chloride C, H, MeBr.SO, Cl. [52°]. Amide C.H. MoBr.SO.Nil. [1390].

m-Bromo-tolueno sulphonic acid $\mathbf{C}_{\mathbf{d}}\mathbf{H}_{\mathbf{d}}\mathbf{MeBr}(\mathbf{SO}_{\mathbf{d}}\mathbf{\Pi})$. Formed by Sulphonating mbrono-toluena (Grete, B. 7, 795; 8, 565; A. 177, 233).—Bo A', 2 aq.—SrA', 2 aq.—CaA', 2 aq.—MgA', 2 6aq.—CuA', 2 4aq.—PbA', 3 aq. According to Wroblewsky (A. 168, 166; Z. [2] 7, 6) three bromo-toluene sulphonic acids are formed by sulphonating m-bromo-toluenc, the Ba salts boing BaA' aq. S. 528, BaA' 3aq. S. 1.452, and BaA', 2 aq. S. 5.248 at 19°; Greto, however, could only obtain the acid just described.

Bromo-tolucno o-sulphonio acid C,H,MoBr(SO,H). From (1, 4, 2)-toluidino sulphonic acid by bromination and climination of NH₂ (Weekwarth, A. 172, 196).—NaA' laq.— $KA'aq.-BaA'_22_3^1aq.-SrA'_22_3^1aq.-PbA'_23_2^1aq. CuA'_22_3^1aq.$

Chloride C, H, MeBr(SO, Cl): erystalline at -- 20°.

Amido Calla McBr(SO2NH2): [0. 164°]; ncodles.

Bromo-toluene sulphonic acid

C.H.McBr(SO.II). From toluene by sulphonation, nitration, reduction, and displacement of NH, by Br (Weckwarth, A. 172, 193; Hayduck, A. 177, 57). —BaA'"aq.

Chloride. Oil, slowly solidifying. Amide. Does not melt below 230.

 $p extbf{-Bromo-toluene}$ $m extbf{-sulphonic}$ acid C.H.MeBr(SO.H) [1:4:3]. [c. 108°]. From p-toluidino m-sulphonic acid le exchange of NII. for Br. Formed also in sulphonating p-bromotolueno (E. Richter, A. 230, 319; Retschy, A. 169, 7; v. Pechmann, A. 173, 208; Nevile a. Winther, C. J. 37, 631). Lamine (containing aq). -- BaA'₂7aq. -- SrA'₂7aq. -PbA'. 3aq.

Chloride C,H,MeBr(SO,Cl). [61°]. Amide Calla McBr(SO, NH.). [152]. p-Bromo-tolueue sulphonic acid

C_sH_sMeBr(SO₂H) [1:4:2]. The chief product of the sniphonation of p-brome-tola no (Hübner, A. 169, 6; Z. [2] 7, 618). Formed also from p-toluidino o-sulphonic acid by exchange of NII. for Br (Jenssen, A. 172, 237). Reduced by sodium-amalgam to tolaene o-sulphonie acid. NaA'2aq.—CaA'24aq.—BaA'2aq. S. 53 at 8°.—SrA'2aq.—PbA'23aq.—CuA'27aq.

Chloride CaHaMeBr(SO.Cl). [35°]. Amide C. H. MeBr(SO2NH2). [1679 p-Browo-toluene ex-sulphonio acid

C, H, Br.CH, SO, H [1:4]. p-Bromo-benzyl sulphonic acid. From p-bromo-benzyl bromide and K,SO, (Jackson a Hartshorn, Am. 5, 264). Also from C₆H₄(NH₂), CH₂SO₄II by diazo- reaction (Mohr, A. 221, 222).—KA'. S. 6·2 at 18°.
—CaA'₂.—BaA'₂1¹₂aq.—BaA'₂aq. S. (of BaA'₂)
67 at 18°.—PbA'₂. S. 2 at 18°.

Chicaida. (10.721/Al.). (115°1/H a. H.)

Chloride. [107°] (M.); [115°] (J. a. H.).

Brome-toluene disulphonic acid

C.H.MeBr(SO.H), [1:2:3:5]. From

C.H.Me(NH.)(SO.H), by diazo-reaction (Limpricht, B. 18, 2177; Hasse, A. 230, 294).—

K.A. 4aq.—BaA' 1aq.

Chloride [90°].

Amide [238°].

p-Bromo oluene disulphonic acid
C₄H₂BrMc(SO₃H)₂. From p-bromo-toluene,
H₂SO₄ and SO₅ (Kornatzki, A. 221, 192). Cauliflower-like deliquescent crystals. Boiled for sixteen houls with cone. nitric acid it forms sulphurio acid Callabr(Co.H)(SO,H),, with Call(NO.)Br.McSO,H, and CaH.(NO.)Mc(SO,H),.

Salts. KA" aq.—BaA" 5aq.—PbA" 2aq. Chloride. [99]. Trimetric plates (f.em

 $A'mide C_0\Pi_2McBr(SO_2NH_2)_2$. [above 260°].

Bromo-toluene disulphonic acid C₆ILMoBr(SO_3H)₂ [1:4:3:x]. Prepared from C₆ILMo(NH_2)(SO_3H)₂ by diazor reaction (Limpricht, B. 18, 2179; E. Richter, A. 230, 324).— BaA" 6aq.—K.A" aq.

Chtoride C.H.MeBr(SO,Cl)2. [129°-133°]. Amide C.H.MeBr(SO.NH2). [above 240°].

Di bron.o-toluene sulphonio acid Call MeBr2(SO3H). From o-bromo-toluene msulphonic acid by nitration, and displacement of NO2 by Br (Schäfer, A. 174, 365). -NaA' 2aq.-BaA', 2, aq.

Tri-bromo-toluene sulphonic acid

C.HMeBr3(SO4H). From o-toluidine p-sulphonic acid by bromination and exchange of NII2 for Br (Hayduck, A. 174, 354). - KA'. - BaA', 13aq. The chloride is a syrup, the amide an amorphous powder.

BROMO-(a)-TOLUIC ACID v. BROMO-PHENYL-ACETIO ACID.

p-Brome-o-toluic acid C₆II₄MeBr(CO₂H) [1:f:2]. [118° uneor.]. Fine white needles. Formed by existation of brome-o-ethyl-toluene with dilute HNO, (1.1) at 200° (Claus a. Picszcek, B. 19, 3088).

Bromo-toluic acid CaH, MoBr. COaH [1:30r4:2]. [167°]. From o-tolnie acid and bromino in the cold (Jacobsen a. Wierss, B, 16, 1959; Racino, A. 239, 74). Needles; volatile with steam. On exidation it gives brome-phthalic acid [157°].

Salt .- BaA'2 5aq.

Methyl other MeA'. [16°]. Bromo-o-toluic acid C, H, MeBrCO₂H

[1:4or5:2]. [176°]. Glistening needles. V.sl. sol. hot water. Formed by oxidation of bromo oyylone with dilute HNO3. - CaA'2 2aq (Jacobsen, This acid is perhaps identical B. 17, 2375). with the following.

Bromo-o-toluio acid Call, MeBr(CO2H) [1:5:2]. [187°]. Formed by saponification of the nitrile, Sublimes in needles. V. sol. alcohol, v. sl. sol. water. By alkaline KMnO, it is oxidised to bromo-phthalio acid [168°] (Nourrisson, B. 20, 1016).

Amido C.H.MeBr(CONH.): [182°]; pearly plates (from alcohol); sublimes in needles. Nitrile C.H.MeBr(CN) [1:5:2]. Formed from bromo-o-toluidino by diazotisation and treatment with cuprous cyanide. Long needles. Easily volatile with steam.

Bromo-m-toluio acid C,H,MeBr(CO,H)

[1:4:3]. [1ŏ3°].

Formation .- 1. From bromo-isocymene, C.H.MePrBr [1:3:4] by treatment with dilute HNO. (Kelbe a. Czarnomski, A. 235, 296).— 2. Together with the following acid, by brominating m-toluio acid in the cold (Jacobson, B. 14, 2351).-3. From bromo-nitro-toluene [45°] by treatment with KCN and alcohol at 220°, and saponification of the resulting nitrile (Richter, B. 5, 425).

Properties.—Slender needles; sl. sol. cold HOAc. Oxidation gives (1, 1, 3)-bromo-isophthalic aoid.

Bromo- m-toluio acid CaHaMeBCOaH [1:6:3]. [209° cor.].

Formation.—1. By oxidation of bromo-m-tene (Fittig, A. 147, 32; Jacobsen, B. 14, 2352).—2. Together with the preceding by brominating m-toluic acid (J.).—3. By oxidising the corresponding bromo-m-isocymene (Kelbe, B. 15, 41). -4. From the corresponding amidotoluio acid by exchange of NH2 for Br (Remsen a. Kuhara, Am. 3, 431)

Properties.-Crystallino powder, sol. hot alcohol, insol. water. - CaA', 3aq. -BaA', 4aq. -

Ethyl ether EtA'. [o. -5°]; (270°-275°). Bromo-p-toluio acid C, H, MeBr(CO, II) [1:2:4].

Formation .- 1. By the exidation of bromecymene CoII, MePrBr [1:4:2] (Landolph, B. 5, 268), bromo p-xyleno (Jannasch a. Dieckmann, A. 171, 83), bromo-p-ethyl-toluene (Reiusen a. Morse, Am. 1, 138). -2. By brominating p-toluic acid in the cold (Brückner, B. 9, 407).

Properties .- Thin needles or laminæ (from

water); ni. sol. hot water.

Salts. -BaA', 4aq. -CaA', 3\aq. -CaA', 3aq. Bromo-p-toluicacid CaHa. Mell (COaH) [1:3:4]. [196°]. Formed by oxidising bromo-p-cymene CuH3MePrBr [1:4:3] (Kelbe a. Koschmitzky, B. 19, 1731).

Di-bromo-m-toluic acid C. II. MeBr. (CO.II). [186°]. Formed by oxidising crudo di-bromoxylene with CrO, (Fittig, Alirens, a. Mattheides, A. 147, 36). Minute needles (from alcohol) .-

BaA', 9aq. - AgA'.

Di-bromo-p-toluio acid C₆H₂MeBr₂.CO₂H [4:6:3:1]. [195°]. Needles (from alcohol). V. sl. sol. hot water. Formed by oxidation of dibromo-r-xyleno C. H. (CH₃). Br. [1:4:3:6] in accide acid solution with CrO₂. By further oxidation with KMnO₄, it gives di-bromo-terephthalic acid

With Akino 10 5100 at 2000.

C.H.Br. (CO₂H)₂ [6:3:4:1].

Salts.—CaA'₂&aq: S. 1 at 20°.—BaA'₂ 2aq.

Ethyl ether A'Et: [49°]; (c. 310°); long
needles (Schultz, B. 18, 1762).

C.H.Me(NIL)Br

BROMO o TOLUIDINE C.H.Me(NIL.)Br [1:2:3]. Formed by reducing bromo-nitro-toluene, itself got by the diazo-reaction from bromonitro-toluidine [88°]. Oil. Gives with brominewater di-bromo-o-toluidino, C_aH₂(CH₂)(NH₂)Br₂, [47°]. Heated with conc. HCf at 160° it forms the above di-bronno-toluidine and a crystalline and bromno (Nevne de man, strong acids. 87, 630).

Bromo - toluidine CgH3Me(NH2)Br [1:2:4]. [32°]. (0. 255°). Formed by reducing the corresponding bromo-nitro-toluene, [45°] (Hübner a. Wallach, A. 154, 298; Körnor, Z. 1869, 636; Hübner a. Roos, B. 6, 799; Wroblewsky, A. 168, 177; Heynemann, Z. [2] 6, 402; A. 158, 340; Nevile a. Winther, C. J. 87, 442). Lamine.-B'HCl: six-sided trimetric tables, S. 827 at

11.5°.-B'2H2SO

Bromo-o-toluidine C.H.Mc(NH.)Br [1:2:5]. [58°]. Formed by brominating acetyl-o-tolu-idine (Wroblewsky, A. 168, 162; Z. [2] 7, 135). Also from brome-nitro-toluene [55°] by tin and HCl (Grete, B. 8, 565; A. 177, 249). Rhombohedra (from alcohe!). Its constitution is known because it gives the same di-brome-toluene (q. v.) as bromo-m-toluidine. By displacement of NH. by II it yields m-bromp-toluene.—B'2II2SO4.- $B'_{2}H_{2}SO_{4}I_{4}^{\dagger}aq. - B'HCl. - B'HNO_{3}: [183°]; S.$ 4.92 at 17°

Acetyl derivative Coll3Me(NHAc)Br.

Bromo-toluidine $C_{\rm s}H_{\rm s}Me(NH_{\rm s})Br_{\rm s}[1:3:4].$ [32°] (N. a. W.); [67°] (Wr.); [75°] (H. a. R.). By reducing the corresponding brome-nitro-toluene (q.b.) (Nevile a. Winther, C. J. 37, 442; Wroblewsky, A. 168, 177; Hübner a. Roos, B. 6, 800).

Acetyl derivative C. H. Me(NHAc)Br.

[114°] (N. a. W.).

Bromo toluidine C₀II₃Me(NII₂)Br [1:3:5]. [36°]. (c. 258°). S.G. ¹⁰ 1·1442. Formed by reducing bromo-nitro-toluene, [81°]. Crystallises with difficulty (N. a. W.; Wroblewsky, A. 10.2 (20.2). 192, 203). Reduced by sodium-amalgam to m. toluidiné.—B'HCI.—B'HNO. S. 25 at 13°.— B',H,SO,.

Acetyl derivative Calla Me(NIIAo)Br.

[168°].

Bromo-m-tofnidine C, ff, Me(Nff,)Br [1:3:6] [78°]. (240°). Formed by brominating acetylm-toluidino and Loiling the product with alooholic potash (N. a. W.). Formod also by reducing (1,2,5)-C₆H₃(CfI₃)Br(NO₂). It gives the samo di-bromo-toluene (q. v.) as bromo-o-tolnidine.-B'HNO3: prisms.

Bromo-tolu dine Calla Me(NILa) Br. From obromo-toluene by nitration and reduction (Hübner a. Roos, B. 6, 801). Oil.—*B'HCl: S. 3·1 at 16', .—*B'HNO₃: 1·25 at 19°. Perhaps iden-

tical with the preceding.

Bromo-p-toluidine C. II. Me (NH.) Br [1:4:3]. [26°] (Claus a. Steinberg, B. 16, 914). (240°). S.G. 22 1 50. From acetyl-p-toluidino by bromination and saponification (Wroblewsky, A. 168, 153). Elimination of NII₂ gives m-bromotolucno.—B'HNO₃: [182°]; S. 2·533 at 19°.—B'H.SO, aq.—B'H.C.O.

Active derivative C. H. Me(NHAc)Br.

[117·5°].

Bromo-p-toluidine CaHaMe(NH2)Br [1:4:2]. [26°]. Formed by reducing the corresponding bromo-nitro-tolucno (Nevile a. Winther, C. J. 39, 85).—B'HBr.—B'HCI (Wallach, A. 235, 255).

Di-bromo-o-toluidine C,H,(CH,)(NH,)Br, [1:20r63:5]. [40] (N. a. W.); [50°] (Wroblewsky, A. 168, 187; Z. [2] 7, 210). From o-toluidine and bromino (Nevile a. Winther, C. J. 37, 436).

Di - bromo - toluidine CaH2(CH3)(NH2)Br2 [1:x:3:4]. [98°] (N. a. W.); [85°] (Wr.). By reducing the corresponding nitro-compound (Nevile a. Winther, C. J. 37, 439; Wroblewsky, A. 168, 184). Does not combine with acids.

Caff Me(NH2)Br. Di - bromo - m - toluidine [1:5:3:4]. [59°]. Formod by reducing the corre-

spending di-bromo-nitro-toluene (Nevile a. Winthor, C. J. 37, 447).

Acetyl derivative CaH2(CH3)(NHAc)Br2.

[163°].

Di-bromo - m - toluidine C₆H₂Me(NH₂)Br₂ [1:3:2:5]. [73°]. From the corresponding dibromo-nitro toluene [70°] by reduction. V. sol.

alcohok Nevile a. Winther, C. J. 37, 448). Acetyl derivative C. H. (CH.) (NII Ac) Br..

[145°].

Di-brome-m-toluidine C₆H₂Me(NH₂)Br₂
[1:3:4:6]. [75°]. Formation. -1. Acetyl-m-toluing the minimated. The Moduct is boiled with akoholio KOII and then distilled with dilute acid. This retains brome-toluidine. By fractionally distilling the rest with steam, two di-bromo-m-toluidines [75°] and [35°], and one tri - bromo - toluidino [101°] may be isolated (Nevile a. Win'her, C. J. 37, 410). \$\times 2\$. By brominating the acetyl derivative of bromo-toluidine [32°], and removing acetyl by heating with H2SO4 (2 vols.) and water (1 vol.).

Acetyl derivative C, 112 (CH2) (NHAc) Br...

[168°]. Di - bromo -m - toluidine Call_Me(NH_)Br.

[1:3:2:6]. [35°]. Prepared as above.

Di - bromo - m - toluidino C₆H₂Me(NH₂)Br₂ [1:3:5:6]. [86°]. Prepared by reducing di-bromonitro-toluene [105°] (Nevile a. Winther, C. J. 37, 434). Formed also by heating the acetyl derivative with equal volumes of 11,80, and water .-B'HCl.

Acctyl derivative CaH (CH a) (NHAc) Br. [205°]. Formed by acetylation of the base; also from Br and bromo-acetyl-tolnidine [168°].

 \mathbf{Di} - bromo - p - toluidine CaH, Mc(NIL)Br. [1:4:3:5]. [73°] (N. a. W.); [76°] (Wroblewsky). From p toluidine and bromine (Wroblewsky, A. 168, 188; Nevile a. Winther, C. J. 37, 136). From p-toluidine m-sulphonic acid and Br (Pechmann, A. 173, 216). Converted by N₂O₃ into di-bromo-toluene [39°].

Di - bromo - p - toluidine $C_qH_qMe(NH_s)Br_2$ [1:4:2:5]. [85°]. Formed by reducing the corresponding nitro- compound [87°] (Nevile a. Winther, C. J. 37, 415; Wroblewsky, A. 168, 185). Yields tri-bromo-toluene [111°

Di-bromo p-toluidino C.H.Me(NH.)Br. [1:4:2:6]. [88°]. Formed by reduction of the

corresponding nitro- compound [57°] Di-bromo-toluidine C.H.Me(NH.)Br. [1:4or6:2:3]. [53°]. From the corresponding di-bromo-nitro-tolucne [57°] (N. a. ¿V.)

Tri - bromo - o - toluidine C.HMe(NII.) Br3. [106°]. Described by Gerver (A. 169, 379) as formed by brominating o-toluidine. Nevile a. Winther (C. J. 37, 438) say that no such body is so formed.

Tri - bromo - m - toluidine C.IIMe(NIL)Br3 [1:3:2:5:6]. [94°]. From the acetyl derivative by boiling with alcoholic potash.

Acetyl derivatives C₀H₂(CH₃)(NHAc)Br₂, [181°]. From C₀H₂(CH₃)(NHAc)Br₂ [1:3:2:5] [144°] and Br (Nevilo a. Winther, C. J. 37,

448).
Tri - bromo - m - tolnidine C.HMe(NH.)Br. [1:5:2:3:4]. [96°]. From its acetyl derivative by alcoholio KOH (N. a. W.).

Acetyl derivative C.H(CHs)(NIIAc)Brs [1:5:3:4:2]. [173°]. Formed by brominating C.H. (CH.) (NHAc)Br.

O.HMe(NH2)Br. 'Tri - bromo - m - toluidine [1:5:2:4:6]. [101°] (N. a. W.); [97°] (Wr.). Formed by brominating di-bromo-m-toluidine (Nevile a. Winther, C. J. 37, 448; Wroblewsky, A. 168, 195).

Tri-brome-p-toluidine C₀HMe(NH₂)R₁ [1:4:2:3:5], [33°], From the hydro-chloride of C₁H₂(CH₂)D²(NI₁) [1:2:4] and bromine-water (N. a. W.), t Needles (from alcohol), Tri-broke-p-toluidine C₄HMe(NH₂)Br₂ [1:4:5:6:2], [118°]. Formed by redneing the corresponding nitro-compound [106°] by iron and against abid (Naville a Winther C. J. 39 85). and acctic abid (Nevile a. Winther, C. J. 39, 85).

Tri-bromo-toluidino CaHMe(NHa)Bry. [1130]. From p-tolnidine m-sulphonio acid and Br

(Pechmann, A. 173, 217). Tri-bromo-toluidine CallMc(NH2)Br3. [82°]. From o-bromo-toluene m-sulphonic acid by nitration, reduction and brominution (Schäfer,

A. 174, 362; B. 7, 1355). Tri-bromo-toluidino C.HMe(NII2)Br3. [72°]. From p-bromo-toluene o-sulphonio acid by

similar treatment (S.).

Tetra-bromo-m-toluidine C.Me(NIL)Br. [1:3:2:4:5:6]. [224°]. From bromo-m-toluidine. [37°], aqueous HCl, and bromine (Nevile a. Winther, C. J. 37, 419). White needles (from alcohel).

Tetra-bromo-p-toluidine C.Me(NIL,)Br. [1:4:2:3:5:6], [227°], From $C_0H_2^2Me(NH_2^2)Br_2^2$ [1:4:2:6] dissolved in dilute HCl and treated with bromine-water (Nevile a. Winther, C. J. 39, 85). Also from p-nitro-toluene, FeBr, and Br at 90° (Schenfelen, A. 231, 179).

BROMO-TOLUIDINE SULPHONIC ACID v.

Bromo-amido-toluene sulphonic acid DI - BROMO - TOLUQUINONE C, HMeBr.O. [85°]. Formal together with the tri-brome-derivative by the action of bromine on tolu-quinoue, and separated from it by crystallisation from dilute acetic acid in which it is the more soluble. Yellow needles; in. sol. water and alcohol (Canroneri a. G. Spica, G. 12, 472).

Tri - bromo - taluquinone C.MeBr.O.

[1:3:4:6:2:5]. [235°].

Formation .-- 1. From toluquinone and Br .-2. From tri-bromo-hydrotoluquinone and Fe_Cl 3. In small quantities, by heating cresol with H.SO, MnO, and KBr (C. a. S.).

Properties. — Pale yellow faminæ; insol. water, sl. sol. alcohol. Aniline forms black crystalline C₆MeBr(NPhII).O₂. Aqueous KOH forms C₆MeBr₂(OH)O₂ [197°] (Spica a. Magnanimi, G. 13, 312).

BROMO - op - DITOLYL

[4:30r2:1] C₀H₃MeBr.C₀H₄Me [1:2]. [95° cor.]. From di-tolyl and Br (Carnelley a. Thomson, C. J. 47, 590). Purified by crystallisation from alcohol, from which an oily isomeride first separates. Oxidation gives bromo-terephthalio acid [309° cor.].

Bromo-op-ditolyl [4:1] C_aH₁Me.C_aH₃MeBr [1:2:3or6]. Oil prepared as above. Oxidation gives bromo-diphenic acid [208°] and c-

brome-plithalio acid [197°].

Di-bromo-ditolyl C₁₁H₁₂Br₂ [152°]. From Br and ditolyl in CS₂ (Carnelley a. Thomson, C. J. 47, 591). Long hair-like needles; lese soluble in alcohol than the preceding compound. CrO. in HOAc gives C14HaBr.O2 [166°] and C14HaBr.O (?) [198°], neither of which compounds dissolves in KOHAq.

BROMO - TOLYL - AMIDO - CHLORO - NAPH-THOQUINONE v. CHLORO-N. PHTHOQUINONE-BBO-MO-TOLUIDE.

DI-BROMO - o - TOLYL - AMIDO - PROPIO-NITRILE C.H.McBr..CHMc.CN. [105°]. From o-tolyl-amido-propionitrilo and Fr (Stephan, O. O. 1886, 470).

Di-bromo-p-tolyl amido-propion krile C₄H₂MeBr₂.CHMe.CN. [117°]. (S.). TETRA-BROMO-DI-TOLYL-AMINE

(C₆H₂MeBr₂)₂NH. [162°]. From Br and ditolyl-nitrosamine in alcohol (Lehn § B. 13, 1544). Small needles.

BROMO-p-TOLYL-BENZENE C₁₃H₁₁Br i.e. [4:1] Bromo-phanyl-toluene. [c, 30°]. A product of the bromination of p-tolyl-benzene. Oxidation gives bromodiphenyl-carboxylio acid [191°] and p-bromobenzoio acid (Carnelley a. Thomson, C. J. 51, 88).

Bromo-p-tolyl-bonzene C_aH_aC_aH_AMeBr [1:20r3:4]. [129°]. From tolyl-benzene in CS₂ by Br (Carnelley a. Thomson, C. J. 47, 586; 51, 87). Pearly plates, sl sol. hot alcohol, v. e. sol. bonzene. Oxidises to bromo-terephthalic acid [306° cor.].

Di-bromo-p-tolyl-benzene

[4:1] C₆H₁Br.C₆H₂BrMe [1:2or3:4]. [115°]. From p-tolyl-benzeno in CS₂ and Br (Carnelloy n. Thomson, C. J. 51, 89). Oxidation gives dibromo-diphenyl-carboxylic acid [204°] and p-bromo-benzoic acid.

Di-bromo-tolyl-benzene

[4:1] C₄H₄Br.C₄H₄BrMe [1:30r2:4]. [150]. Formed, together with the isomeride [115°] in brominating p-tolyl-benzene. Oxidation gives di-bromo-diphenyl-earboxylie act [232°] and p-bromo-benzoie acid.

BROMO-TOLYLENE-m-DIAMINE

C_aH_a(CH_a)(NH_a), (Br) [1:2:4:x]. (104°). Colourloss plates. Sol. alcohol, ether, and CS_a. Prepared by bromination of the di-benzoyl-deriv tive of (1:2:4)-tolylene-diamine and subsequent saponification.

Di-benzoyl-derivative [214°]. White needles (Ruhemann, B. 14, 2658).

Bromo-tolylone diamine C_aH₂(CH₄)(NH₂), Br. [107°]. Formed by reducing bromo-di-nitro-tolueno [104°] (Grete, A. 177, 252).—B"21KCl.—B"21KCl.—B"H₂SO₄.—B"H₂SO₄.—B"H₂C₂O₄. This body is perhaps identical with the preceding.

Bromo-tolylone-o-diamine

C₈H₂(CH₃)(NH₂)₂Br [1:2:3:4]. [59°]. Obtained by nitration and reduction of p-brono-o-tolaidine (Hübner a. Schüppbaus, B. 17, 775). Small colourless needles. V. sol. water, alcohol, and benzene.

Salts.—B'HCl: very soluble colourless needles.—B'₂H₂SO₄: colourless tables.

Anhydro-formyl desivative v. Methenyl bromo-tolyleni-o-diamine.

BROMO-TOLYL-ETHANE . BROMO-ETHYL-

a-BROMO-a-m-TOLYL-ETHYLENE

C.H.,CBr:CII₂. Formed by boiling m-tolyl dibromo-etbano C.H.,CHBr.GII₂Br with alcoholio KOH. Vory unstable body: begins to blacken syen at 100° (Müller, B. 20, 1216).

@-Bromo-a-m-tolyl-ethylene C.H.,CH:CHBr.
m-Methyl-bromo-styrene. (242°). Oil. Formed
by adding bromine to a warm solution of sodium

m-tolyl-acrylate (methyl-cinnamate) (Müller, B. 20, 1216).

BROMO-TOLYL MERCÄPTAN C₆H₃MoBr.SH. [c. 7^o]. (c. 246^o). From p-bromo-toluene sulphoethoride, tin, and HCl (Hübner a. Wallach, Z. [2] 5, 500).

Bromo-tolyl mercaptan C_oH_sMeBr.SW. (246°). From o-bromo-tolatine m-sulphochloride (53°] by Zn and H₂SO₁ (Hübner, A. 169, 41). Oil.

DI-BROMO-DI-"OLYL-METHANE C₁₃H₁₄Br₂, [115°]. From the hydrocarbon and Br (Weiler, B. 7, 1181).

BROMO-p-TOLYL-p-METHYL-IMESATIN v. p-METHYL-ISATIN-BROMO-p-TOLUDE.

TETRA-BROMO-p-TOLYL-(8)-NAPHTHYL-AMINE C_{1:} H₁₁Br₄N. [169²]. Formed by bromination of p-tolyl-(B)-maphthyl-amine (Priedländer, B. 16, 2086). White silky needles. Sol. alcohol and other.

BROMO - DI - α - TOLYL - PROPIONIC ACID C_aH_AMeBr.C(C_aH_AMe)Me.CO_aH. [144°], Colourless crystals. Sol. alcohol, ether, &c. Formed by bromination of di-α-tolyl-propionic acid (Haiss, B. 15, 1178).

αβ-DI-BRÓMO-m-TOLYL PROPIONIC ACID C₁₀ II₁₀Br₂O₂ i.e. [3:1] C₁ II₁ Me₂CHBr.CHBr.CO₂H₂ [167]. From m-methyl-cinnamic acid and Br (Müller, B. 20, 1215).

DI - BROMO - DÍ - TOLYL - DI - SULPHIDE (C.11, MeBr), S.,... [78°]. From o-bromo-tolyl mercaptan and dilute HNO₃ (Hübner a. Post, A. 169, 42).

BROMO-UMPELLIFERON

[154°]. Formed by the action of bromine upon a solution of unbelliferon-methyl ether m CS₂, thoug white needles; m. sol. hot alcohol and ether, sl. sol. coll alcohol and ether, insol. water; its dilute alcoholic solution has a green fluorescence. By alcoholic KOH it is converted into methoxy-counterlike acid

silvery tables; sol. boiling alcohol and ether, sl. sol. cold acchol; its dilute alcoholic solution has a violet fluorescence. Analogous to the methyl-other in its formation and properties (Will a. Beck, B. 19, 1782).

Di-bromo-umbellifc...on

[2510]; formed by bromination of numbelliferonmethyl-ether dissolved in acetic acid; white glistening needles; sl. sol. alcohol.

[216°]; formed by bromination of umbelliferonethyl-ether in acetic acid (Will a. Beck, B. 19, 1786).

Tri-brome-umbelliferon C.H.Br.O. [194°]. From umbelliferon and bromine-water (Posen, B. 14, 2746; Mössmer, A. 119, 261). alcoholic solution shows greenish-yellow fluor-

BROMO-UVITIC ACID CaH2Br(CH3)(CO2H)2 [x:1:3:5]. Prepared by oxidising bromo-ω₁ω₂-dioxy-mesitylene with KMnO, (Colson, A. Ch. [6] 6, 102). White crystale, carbonisos at 285°

without melting. Sol. alcohol.—Na.A": tables.
a.BROMO-n-VALERIC CID C.II.BRO. i.e. C.H., CHBr.CO.H. Formed by bromination of valerio acid (propyl-acetic acid) (Juslin, B. 17, 2504).

Ethyl ether A'Et: (191°); S.G. $\frac{18}{4}$ =

1.226; colourtess fluid:

γ-Bremo-n-valeric acid CH₃.CHBr.CH₂.CH₂.CO₂H. From allvl-acetio acid and conc. HBrAq at 0° (Messeaschmidt, A. 208, 94). Boiling water or cold Na CO, Aq convert it into the lactone of γ-oxy-valeric acid

a-Brome-isevaleric acid

(CH₃)₂Cl1.ClIBr.CO₂H (chiefly). [40°]. (230°) From ordinary valeric acid and Br (Callours, A. Suppl. 2, 74; Borodine, A. 119, 121; Fittig a. Clark, A. 139, 199; Ley a. Popoff, A. 174, 63; Schmidt, A. 193, 104). Formed also by the action of water on its bromide which is formed by treating isovaleric acid with Br and P (Volhard, A. 212, 163). Oil.

Ethyl ether EtA'. (186°). Is best purified by distillation with steam, the liquid is collected as soon as the oily drops sink under

water (Lovén, J. pr. [2] 33, 112).

Brome-valeric acid CMeEtBr.CO2H or, possibly, CHa CHBr. CHMc. CO2II Bromo - methyl ethyl-acetic acid. Bromo-hydro-tiglic acid. [66°]. From tiglic and angelic acids with conc. IIBrAq at 0° (Fittig a. Pagenstecher, 4. 195, 108, 128; ef. p. 267). Monoclinio tables (from CS₂); insol. cold water. Boiling water wrms tiglic acid and somo pseudo-butylene (s-di-methyl-ethylene). Na₂CO₃Aq produces chiefly pseudo - tutylene. Sodium - amalgam forms methyl - ethyl - acetio acid.

Bromo - valeric acid C₃H₉BrO₂ i.e. CMeEtBr.CO.H? From methyl - ethyl - acetio acid and Br at 160° (Böcking, A. 204, 23). Liquid. Should be identical with the preceding.

Ethyl ether EtA'. (185°). S.G. ¹⁸ 1·2275. Decomposed by boiling Na CO₃Aq into α-methyl-

α-oxy-butyric ether CMeEt(OH).CO.H.

γδ-Di-bromo-n-valeric acid CH2Br.CHBr.CH2.CH2.CO2H. From allyl-acetic acid and Br in CS2 (Messerschmidt, A. 208, 100). Thin lamine. Converted by sodium-amalgam into allyl-rectic acid. Boiling water forms the lactone of bromo-oxy-valerio acid, and finally C,H,O,.

Di brome-valsrio acid CMe,Br.CHBr.CO,II. [106°]. Solidifies at 76°. From CMe, CH.CO,II. and bromine (Ustinoff, J. pr. [2] 34, 483).

Di-breme-valeric acid CH2.CHBr.CBrMe.CO2II. [86°]. From tiglio acid and Br (Schmidt a. Berendes, A. 191, 119); also from angelio acid and Br (Jaffé, A. 135, 293; Pagenstecher, A. 195, 123). Triolinic 293; Pagenstecher, A. 195, 123). erystals (from CS_2); insol. cold water. Converted by distillation or by sodium-amalgam B. 18, 1277). Mol and Na give ψ -oumene.

into tiglic seid (Demarcay, B. 8, 830). Beiling water decomposes its salte forming bromo-buti-

nene (87°).—KA'. insol. conc. KOHAq.
Ethyl ether Eth'. (185°). (J.).
DI-BROMO-VALERIC ALDEHYDE C.H.Br.O i.e. CH3.CHBr.CBrMo.CHO. From tiglio alde-

hyde and Br (Lieben a. Zeisel, M. 7, 55).
BROMO (ALERO-LACTONE v. Brome-oxy-

VALERIC ACID.

BROMO-VALERYLENE C.H.Br. 130°). From valerylene dibromide and alcoholic KOH (Reboul, A. 135, 372). Forms a yellow pp. of C₃H₃Cu with ammoniacal cuprous ehlorido.

BROMO-VANILLIC ACID v. Methyl deriva-

tive of Bromo-di-oxy-benzoio cid.

BROMO-VERATRIC ACID v. Methyl deriva-

tive of Bromo-di-oxy-benzoic acid.

BROMO-VINYL ACETATE C.H.Bro. i.e. CHBr:CH.OAc. From acetylene di-bromide and KOAc at 160° for 2 days (Sabanejeff, A. 216, 272). Crystals. Explode when quickly heated. Forms with bromino a compound C4H4Br2O2, which solidifies at 0°.

ω-BROMO-p-VINYJ -- PHENOL, Methul ether C.H.(OMe).CH:CHBr. [55°]. From the di-methyl-ether of aB-di-bromo-p-oxy-phenylpropionio aoid C₆∏₄(OMe).CHBr.CHBr.CO₂Me by boiling with aqueous KOH (30 p.c.). Platos, of pleasant smell and tasto (Eigol, B. 20,

eso-Breme-o-vinyl-phenel

CaHa(OII)Br.CII;CH2. Bronto-oxy-styrene. (265°). Formed by distilling di-bromo-othyl-phonol, C.H. (OH)Br.C.H.Br obtained by brominating o-othyl-phenol (Suida a. Plohn, M. 1, 180). Liquid, sl. sol. water. Gives a reddish-brown pp. with FogCls.

Di-breine-o-vinyl-phenel. Methyl ether C_vH_sBr₂O i.e. C₆H₃(OMe)Br.C₂H₂Br. From tho methyl derivative of tri-bromo-oxy-phenyl-propionic acid CoH2(OMe)Br.CHBr.CHBr.CO.H and Na₂€O₃Aq (Perkin, C. J. 39, 418). Oil; slightly

volatile with steam.

BROMO-VINYL-PIPERIDINE

 $C_3H_{10}N(C_2H_2Br)$. Piper-ethyl-alkine-bromide. Formed by heating the hydrobromido of piperethyl-alkino (oxethyl-piperidine) with 1 mol. of bromine at 100° 120°. On reduction with sodium-amalgam it yields ethyl-piperidine.-B'HBr: thin colourless prisms, sol. water, v. sl. sol. cold alcohol. -B'HCl. -B',H,Cl,PtCl,-

St. 801. Solid alcohol.—Br. 17. 154).

BROMO - o - XYLENE C₅H₅Br t.e.

C₅H₅(CH₃)₂Br [1:2:4]. [0°]. (214° i. V.). Mol.

w. 185°. S.G. ½ 137. Formed by the action of bromine in presence of iodine upon o-xylene in the cold and in the dark (Jacobson, B. 17, 2372; Schramm, B. 18, 1278).

ω-Brome-o-xylene C₀H₄(CH₂),CH₂Br. o-xylyl bromide. [21°]. (217¿) at 742 mm. S.G. 23 1·381. Colourless liquid. Prepared by the action of bromine-vapour upon boiling o-xylene; or in the cold upon o-xylene exposed to direct sunshino (Radziszowski a. Wispek, B. 15, 1747; 18, 1281; Schramm, B. 18 1278; Colson, A. Ch.

[6] 6, 115).

Bromo-m-xylene $C_aH_3(CH_3)_2Br[1:3:4]$. (206°). Formed by the action of bromine upon cold m.

(B) - Bromo - m - xylene O,H,Me,Br [1:3:2]. (o. 206°). Liquid at -10°. Obtained by adding HCl solution of broming to a solution of sodium m-xylene (β) -sulphonate, which is prepared by debrominating di-bromo-m-xylene sulphonic acid C_cHMe₂Br₂(SO₃H) [1:3:4:6:2] with zinc-dust and aqueous amnonia. By methyl iodide and sodium it give hemimellithene C_cH₃Mo₃ [1:2:3] (Jacobsen a. Deike, B. 20, 903).

Bromo-xylene C, II, Mc, Br [1:3:5]. (201°). 8.G. 20 1.362. Oil. From C.H.Me.(NH.) [1:3:4] vid Calla Moull HAc, Calla Bran HAc and C.H.BrMc2(NH2) (Wroblewsky, A. 192, 215; B.

Promo-m-xylene Call (CH3).CH Br. m-Xylyl bromide. (215°) at 735 mm. (R. a. W.); (c. 218°) (C.). S.G. ²³ 1·371. Colourless liquid, with pungent vapour. Prepared by the action of the vapour of bromine on boiling m-xylene, or from Br (1 mol.) and cold m-xylene exposed to direct sunlight (Radziszowski a. Wispek, B. 15, 1745; 18, 1282; Schramm, B. 18, 1277; Colson, A. Ch. [6] 6, 117).

Brome-p-xylone C. 14, Br(CH,), [2:1:4]. [10°]. (206° i.V.). Formed by the action of bronnine upon cold p-xylene in the dark (Fittig a. Jannasch, A. 151, 283; 171, 82; B. 17, 2709; Jacobsen, B. 18, 356; Schramm, B. 18, 1276).

Large plates.

w-Bremo-p-xylene C_aH₁(CH_a).CH_Br. p-Nylyl bromids. [36°]. (219°) at 740 mm. Long colourless needles. Sol. ether and chloroform. Pungent smell. Prepared by the action of bromine vapour on boiling p-xylene, or of Br (1 mol.) upon cold p-xylene exposed to direct sunshine (Radziszewski a. Wispel, B. 15, 1743; B. 18, 1279; Schramm, B. 18, 1: 4).

Di-bromo-o-xylene C₆H₂(CH₂)₂Br₂ [1:2:3:4?]. [7°]. (277°). S.G. ¹⁵/₁₅ 1.7842. Liquid at ordinary temperatures. Formed together with the isomeride [88°] by the action of bromine (2 mols.) in presence of iodine upon o-xyleno (1 mo'.) in

the oold (Jacobsen, B. 17, 2376).

Di-bromo-o-xylene $C_e H_2(CH_3)_2 Br_2$ [1:2:4:5]. [88°]. (278°). Large trimetric plates, or long needles. V. sol. hot alcohol; v. sl. sol. alcohol at 0°. Formed as abovo (Jacobsen, B. 17,

2376).

o-Di-\(\omega\)-bromo-xylone C₆H (CH_{\omega\}Br)₂ [1:2], o-Xylylene bromide. [95°], S.G. \(^0\) 1-934, S. (ether) 20. S.H. (15°-40°) 183. Splendid large trimetric crystals. Prepared by heating o-xylene with bromino (2 mols.) at 130°-155". Also by the action of Br (2 mols.) in the cold upon oxylene (1 mol.) exposed to direct sunshine. Trimetric orystals; a:b:c=8581:1:5014; v. sol. ether and chloroform. By boiling with a solution of Na₂CO₂ for 3 hours it yields phthalylalcohol (Bacyer a. Perkin, B. 17, 123; Radziszewski a. Wispek, B. 18, 1281; Schramm, B. 18, 1278; Colson, A. Ch. [6] 6, 705; C. R. 101, 428; Perkin, jun., C. J. 53, 5).

Di-bremo-m-xylene C.11. (Cll.) Br. (256°). From m-xylene and Br (Fittig, A. 147, 25; 156, 236).

Di-bromo-m-xylene C6H2(CH3)2Br2. Liquid. From brominated m-xylidine by the diazo- reaction (Wroblewsky, A. 192, 216). ω₁ω₂-Di-bromo-m-xylene C₂H₄(CH₂Br)₂ [1:3].

Kylylene dibromide. [77°]. S.G. 2 1.734; 22 | 19, 2137).

(liquid) 1.615. S. (ligroin) 83. S.H. (15°-40°) 184. Formed by the action of bromine (2 mols.) in the cold upon m-xylene exposed to direct sunshine; also by adding Br (1 mols.) to m-xylene at 130²–180' (Schramm, B. 18, 1277; Radziszewski a. Wispek, B. 13, 128; Colson, A. Ch. [6] 6, 109, C. R. 101, 428, Kipping, C. J. 53, 26). Prismatic needles (from CllCl₂). Attacks the eyes. Converted or alcohol or boiling water into C₂H₂(CH₂OH). Alkalino permanganate forms isoplithalic a rid.

Di-brome-p-xy: ne C. II. (CII.). Br. [1:4:2:5]. Ji-promo-p-xy. ne Collatori, in Landon [76°]. (261°). From p-xylene, br, and L. A small quantity of a liquid isomeride, C₆H₄(CH₂)₂Br₂ [1:4:2:6] (?) is formed at the same time. Large triclinic crystals, plates, or flat needles (Jacobsen, B. 18, 35s; Fittig, Almens a. Mattheides, A. 147, 26; Januasch, B. 10,

1357).

 $\omega_1\omega_2$ -Di-bromo-p-xylene $C_nH_1(CH_nBr)_2$ [1:4]. p-Xylylene bromide. [114°]. (c. 245°). S. (ether) 2·65 at 20°. S.11. (15°-40°) ·180. Formed by the action of bromine (2 mols.) in the cold upon p-xylene exposed to direct sunshine; or by the action of bromine vapour on boiling p-xylene (Grimaux, Z. 1870, 394; Sehraum, B. 18, 1277; Radziszewski a. Wispek, B. 15, 1744; 18, 1279; Löw, A. 231, 362; B. 18, 2072; Colson, C. R. 104, 128; A. Ch. [6] 6, 119; Kipping, C. J. 53, 34). Plates; sol. CHCl, Boiled with water (20 pts.) and lead nitrato (1 pt.) it forms terephthalic ablehyde and some terephthalic acid and p-aldehyde-benzoic acid. Fuming HNO, acts similarly. Alcehol converts it into C.H.(CH.OH); the rate of this saponification is less than with the o-, and still less than with the m-isomeride.

e.vo-Tri-bromo-p-xylene C_uII₄(CHBr₂)(CII₂Br). [106°]. Formed from p-xylene and impure bromine-vapour (Löw, A. 231, 363). Philing water gives w-oxy-toluic aldehyde.

Tetra-bromo-o-xylene $C_b((\Pi_s)_2Br_s, [262^\circ]$ (J.); [55°] (B.). (375°). From o-xylene and Br (Jacobsen, B. 17, 2378) in presence of Al Br_s (Blümlein, B. 17, 2492). Long glistening needles. V. sl. sol. hot alcohol; v. sol. hot

Tetra-bremo-p-xylcne $C_a(C11_3)_2Br_4$. [253]. (355°). From p-xyleno and Br (Jacobsen, B. 18,

BROMO-o-XYLENE-SULPHONIC ACID C₆H₂(CH₃)₂(J)(SO₃H) [1:2:4:5]. Formed by sulphonation of bromo-o-xylene. Crystals (containing aq). Very soluble in water, sparingly in cold dilute H SO

Salts .- NaA'laq: long fine needles. v. sol. hot water. - KA' aq. - BaA', 3aq: long thick prisms, sol. hot water.

Amide C₀II₂(CII₃)₂(Br)(SO₂NII₂): [213°]; leng fine needles, v. sl. sol. water, sl. sol. cold alcohol (Jacobsen, B. 17, 2373).

Bromo-o-xylene-sulphenie acid

C. H2(CH3)2Br(SO311). Formed by the action of bromino upon an aqueous solution of o-xylenesulphonic acid. - BaA' 4aq : sparingly soluble necdles.

Amide C.H.Mc.Br.SO.NH.: [1870]; thick needles; v. sol. hot alcohol (Kelbo a. Stein, B.

Bromo-m-xylene sul ihonio gold

Bromo-m-xylene sul honio acid

O_eH₂(CH₂)₂Br(SO_eH) [1:3:2:4]. From di-bromom-xylene sulphancide by sodium amalgam
(Jacobsen a. Woinborg, [B. 11, 1636).

Amide C_eH₂Mo_eBr(SO₂NH₂. [161° cor.].

Bromo-m-xylene sulthonic acid

C_eH₂Mo_eBr(SO₂H) [1:3:6:1]. From

C_eH₃M·₂(SO₂H) [1:3:4] by fir or from C_eH₃Me₂Br

[1:3:6] by funning H₂SO₁(Neinberg, B. 11, 1062).

From C_eH₂Mo_e(NH₂)(SO_e) by diazo- reaction
(Sartig, A. 230, 335; Nochung a. Kohn, B. 19, 139; Limpricht, B. 18, 2188.

V. e. sol. water. v. e. sol. water.

Salts. -- BaA' aq. -- NaA' aq. -- ZnA' 29aq. --CuA', 7aq.

Chloride [61°]: large prisms.

Amide [194°] : small trimetric prisms. Bromo-p-xylene sulphonic acid

C₆H₂Me₂Br.SO₂II [1:4:2:5]. Formed by heating diazo-p-xyleno-sulphonic acid C₆H₂Me₂< N₂ [1:4:2:5] with strong IIBr. - BaA', 2aq: small white plates.

Chloride: [78°]; small white prisms.

Amide: [201°]; small white plates, v. sol. alcohol and ether, sl. sol. water, benzene, and chloroform (Nölting a. Kohn, B. 19, 141).

Bromo-p-xylene-sulphonic acid

 $C_0H_2(CH_3)_2(Br)(SO_3H)$ [1:4:2:x]. Pearly plates or flat needles. Formed by sulphonation of bromo-p-xyleno.

Salts.-NaA'aq: long thin prisms, trimetric plates, or six-sided plates.—BaA', : small prisms or thin six-sided plates.

Amide C₆H₂(CH₃)₂(B₄)(SO₂NH₂). [206°], flat prisms, v. sol. not alcohol (Jacobsen, B. 17, 2378). This acid is perhaps identical with the preceding.

Di-bromo-m-xylene sulphonic acid

C₆H(CH₃)₂Br₂(SO₃H) [1:3:4:6:2]. From di-bromom-xylene [72°] and funning II, SO, (Jacobsen a. Weinberg, B. 11, 1534). Scattlets, sl. sol. cold water. Reduced by sodium amalgam to (1,3,2)m-xylene sulphonic acid.

Salts.—BaA'2.—NaA' 2aq: leaflets. Chloride [107°]: rhombic leaflets. Amide [220°]: slender needles.

BROMO - m - XYLENOL C.H. (CH.) Br (OH) [1:3:x:4]. From Br and m-xylenol in HOAc. Liquid (Jacobsen, B. 11, 21).

Bromo-p-xylenol $C_6H_2(CH_3)_2Br(OH)$ [1:4:x:2]. [87°]. From p-xylenol and Br (Jacobsen, B. 11, 27).

Di - bromo - m - xylenol C.H(CH3),Br2(OH) [1:3:?:?:4] [73°] (J.).

 $\omega_1\omega_2$ Di bromo p-xylenol $C_0H_3(CH,Br)_2(OH)$ [1:4:2]. [74°]. S. (alcolol) 200. From p-xylenol and Br at 160° (Adam, Bl. [2] 41, 288). Needles; insol. water, but decomposed on boiling with it, HBr coming off.

Tri - bromo - o - xylenol C (CII3) Br3.OH [1:2:4:5:6:3]. [184°]. Fino necdles. Formed by bromination of o-xylenol C₆H₃(CH₃)₂.OH

[1:2:3] (Thöl, B. 18, 2562). Tri - bromo - o - xylenol Tri - bromo - o - xylenól $C_a(CH_3)_2Br_3(OH)$ [1:2:3:5:6:4]. [169°]. From (1, 2, 4)-o-xylonol. Felted needles (Jacobsen, B. 11, 28).

 $C_a(CII_a)_2Br_a(OII)$ Tri - bromo - m - xylonol [1:3:2:5:6:4]. [179°]. From (1, 3, 4)-m-xylenol (J.). Long needles.

Trl - bromo - m - xylenol Ca(CHa)zBrz.OH [1:3:2:4:6:5]. [166°]. Fine white noedles. From m-xylenol O₂H₃(OH₃)₂,OH [1:3:5] (Nölting a. Forel, B. 18, 2679; cf. Thöl, B. 18, 362).

Tri-bromo-p-xylenol C_a(CH₂)₂Br₄(CH). [175°].
Golden yell-aw needles (Jacobsen, B. 11, 26).
BROMQ¹— "XYLIDINE C₄H₁₆BrN i.e.
C₄H₄(CH₃)₂Br(NH₄) [13:5:4]. [37°]. From acetyl-m-xylidino by bromination and saponification. Minute needles (from diluto alcohol). Converted by the diazo-reaction into s-bromoxylene (Gev.z., B. 3, 225; Wroblewsky, A. 192, 215).

Di - bromo - o - xylidino C₆11(CH₂)₂Br₂.NH₂ [1:2:4:5:3]. [103°]. Obtained by redugition of the corresponding nitro- compound with iron and acetic acid. Colourless needles. V. sol. alcohol ether, and acctic acid. Does not form salts. By sodium-amalgam it is debrominated (Thöl, B. 18, 2562).

Di-bromo-m-xylidine C,11(C113)2Br2(NII2). From acetyl-m-xylidino by brominating and saponifying (Genz, B. 3, 225). Needles (from alcohol).

Di-bromo-p-xylidine C.HMc.Br.(NH.) [1:4:5:x:2]. [65°]. Formed by acidifying an alkaline solution of (1 mol. of) p-xylidine-sulphonio acid $C_0H_2Me_2(NH_2)(SO_3H)'[1:1:2:5]$ and (2 mols. of) bromine. Also formed by bromination of p. xylidine (Nölting a. Kohn, B. 19, 142)

BROMO-m-XYLIDINE-SULPHONIC $C_6HMe_2(Br)(NH_2)(SO_3H)$ [1:3:57:4:6]. Small white needles. Sol. hot, v. sl. sol. cold water, insol. alcohol. Formed by bromination of mxylidine-sulphonic acid C_sH_sMe_s(NH_s)(SO_sH) [1:3:4:6] (Nölting a. Kohn, B. 19, 140).

Bromo-juxylidino-sulphonic acid

C. HMc. Br(NH2)(SO3H) [1:4:c:6:2]. Small white plates. Nearly insoluble in cold water. Formed by bromination of p-xylidine-sulphonic acid C_aH₂Mc₂(NH₂)(SO₃H) [1:4:6:2]. —A'K (Nölting a. Kokn, B. 19, 143).

DI-BROMO-m-XYLOQUINONE C.Br.Me.O. [1:3:4:6:2:5]. [174°]. From mesitol, C. H. Me. (OH) and Br in water (Jacobsen, A. 195, 271). Golden laminæ (from alcohol), insol. water and Na CO3Aq, decomposed by KOHAq.

Di-bromo p-xyloquinone $C_aBr_aMe_aO_a$ [1:4:5:2:3:6]. [181°]. Formed by the action of bromine upon p-xyloquinono under water. Thin golden plates. Insol. water and cold alcohol; sol, ether and benzene (Carstanjen, J. pr. [2] 23, 434).

BRONZE v. COPPER, ALLOYS OF.

BRUCINE C₂₃IL₂₀N₂O₄4aq. [105°, hydrated]; [178°, anhydrous] (Cians, B. 14, 773); [151°] (Blyth). S. 12 at 15°; 2 at 100°. [a]₀ = -85° (in alcohol); -110° to -127° (in CHCl₃) (Oudemans, A. 166, 69).

Occurrence:-Together with strychnine in nux vomica (the seeds of Strychnos nux vomica). in the bean of St. Ignatius (the seed of Strychnos Ighatii), in the wood of Strychnes colubrina, in upas tiente, extracted from the bark of S. tiente, and in the bark of Strychnos nux vomica (False Angustura bark) (Pelletier a. Caventou, A. Ch. [2] 12, 118; 26, 53; Pelletier a. Dumas, A. Ch. [2] 24, 176; Corriol, J. Ph. 11, 495; Liebig, A. Ch. [2] 47, 172; A. 26, 50; Regnault, A. Ch. [2] 68, 113). Brucine, free from strychnine, BRUCINE.

occurs in the bark of Strychnos Ligustrina (Bidara Laut) (Greenish, Ph. [3] 9, 1013).

Preparation.—Nux vemica seeds (56 lbs.) are powdered and exhausted with alcohol, to which one-sixth part of water has been added. The alcohol is boiled off and the residue treated with water (40 lbs.) containing H.SO, (42 lbs.). The filtrato is neutralised with Na.CO₂. The pp. is collected after a few hours, dissolved in chloroform, and the solution shaken with very dilute H₂SO₄ which dissolves brucine. The solution is placed under a bell jar together with a beaker containing ammonia so that neutralisation The crystalline pp. is proceeds very slowly. oxtract with disute alcohol, which dissolves brucine, and the solution allowed to evaporate. The brucine, containing stryelmine, is dissolved in dilute H.SO, and the faintly alkaline solution mixed with KI. The brucine hydricalide that is ppd. is crystallised several times from alcohol. 1t is then shaken with aqueous Na₂CO₃ and chloroform, the chloroform is then shaken with dilute acid and the base pp. with ammonia (W. A. Shenstone, C. J. 39, 143).

Properties. Monoclinic efflorescent prisms (from dilute alcohol); v. sol. alcohol, chloroform, and essential oils, insol, ether and fatty oils. Feebly alkaline to litmus or phenol-phthalein (Plugge, Ar. Ph. [3] 25, 45). It is havorotatory, the extent of rotation depending upon the nature of the solvents and the concentration of the solution. In presence of free acids [a]_D varies from -29° to -36.5° (Tykociner, R. T. C. 1, 144). Commercial benefits a negative angular contains Commercial brucine usually contains strychnine, but the colour which strychnine gives with oxidising agents is masked by the presence of brueine. The stryel are may, however, be detected by adding dilute HNO, and then extracting the stryclmine with chloroform in presence of excess of KOH (Shenstone, Ph. [3] 8, 445; Hanriot, C. R. 97, 267). Brucine is affected by heating with acids, alkalis, or even water, and hence much is lost by the usnal method of preparation, but the products being usually amorphous, the unaltered bracine is easily separated in a pure state.

Colour tests. - V. p. 124. The most characteristic is the red colour with nitric acid, which, after warming, is turned violet by excess of ammonium sulphide (v. also Cotton, Z. [2] 5, 728; J. Ph. [4] 10, 18; Luck, Z. [2] 6, 275; Le Linde, C. M. 37, 98; Elinckiger, Fr. 15, 312; Hager, Fr. 11, 201; Dragendorff, Fr. 18, 108; Pandis, C. C. 1872, 410; Struve, Fr. 12, 16;

Buckingham, Ph. [3] 3, 881).

Reactions.—1. Bracine (60 g.) heated with alcohol (600 e.e.) and NaOH (30 g.) at 100° for 12 hours forms a solution which, after neutralisation with CO₂, filtration and evaporation, loaves a varnish containing some crystals. The crystals (7 g.) separated from the varnish by washing with water and purified by solution in very little dilute HCl and ppn. by NH, are thrown down as microscopic crystals of C₂,H₂,N₂,O₃ (2 g.). This gives a yellow colour with HNO₃; a pp. with bromine, and on boiling a magenta colour. An ammoniacal solution turns purple in air and finally green (Shenstone).—2. HNO₃ (S.G. 1·2 to 1·4) forms a red selution, producing 'caectheline' C₂,H₂N₂O₉, oxalic acid, and methyl nitrito (Streeker, A. 91,

76; Hanssen, B. 20, 4(1). K₂Cr₂O, and H₂SO₄ oxidise eacotheline to (b₁cH₁₁N₂O₄, which is also formed by the exidation of brucine.—8. Brucine (1 mel.) heated with H H at 140° gives off McCl (between 1 and 2 mel.). Brucine is therefore possibly C₂₁H₂₀(OMe⁷,N₂O₄, stryclinino being C₂₁H₃₀(OH)₂N₂O₄ (Shr instone, C. J. 43, 101; cf. Hanssen, B. 17, 226° .—4. Yields on distillation with potash several pyridine bases. Amongst offices (β)-di-methyl pyridine and (β)-tri-methyl pyridine, quinolin tetrahydride, together with probably a diag thyl-pyridine (Occlusner de Coninck, A. Ch/§ | 27,507; C. R. 99, 1077; Bl. [2] 42, 100). 5. Distilled in vacuo with zinedust it yields carbaxol (Lacbisch a. Schoop, M. 7, 613).—6. Alkalino KMnO₁ gives off about half the nitrogen in the free state (Wanklyn a. Chapman, G. J. 21, 161).—7. 1Cl forms a light flocculent pp. in solutions of salts of brueine (Dittmar).

Salts .- B'HCl: crystalline tufts, m. sol. water .- B'HClllgCl.: long needles (from alcohol). -B',H,PtCl, : yellow pp. -B'HI aq : rectangular lamine or very short prisms; m. sol. hot water.—B'III₃: brown violet needles (Jörgensen, A. Ch. [4] 11, 114; J. pr. [2] 3, 160).— B'HILL: unstable orange needles. B'HNO, 2aq: four-sided prisms; less soluble in water than strychnine nitrate. B'2H2SO17aq: long needles, v. sol. water, sl. sol. alcohol. -- B'_H_S_O_55aq; S. 1 at 15° (How, N. Ed. P. J. [new] 98).--B'aH.S. Gaq: [125]; insoluble yellow needles, formed by atmospheric oxidation of an alcoholic solution containing H.S (Schmidt, A. 180, 296; B. 8, 1267; 10, 838, 1288). — $B'_{3}H_{4}S_{12}$. B' H 3PO ma (Anderson, P. M. [3] 33, 163).-B' H FeCy, 2nq: hygroscopic needles, m. sel. hot water (Brandis, A. 66, 266). B',H. FeCy, 4aq. -13 H₁ FeCy₈: white powder (Holst a. Beckurts, Ar. Ph. [3] 25, 313). — B'₃H₄FeCy₈ 6aq. — B'₃H₄CoCy₄ 10ac (Lee, Am. S. [2] 2, 44). — B'₃H₄NiCy₄ (h.).—P'HCyS: scales, sol. water, — B'₄H₄NiCy₄ (h.).—P'HCyS: scales, sol Periogate: needles (Bödeker, A. 71, 64; Langlos, A. Ch. [3] 31, 278). Nitroprusside: S. 15 at 15° (Davy, Ph. (3 | 11, 756). Acctate: crystalline, but gummy if impure (Shenstone). -Dextro-tartrates: $B'_2C_1H_aO_a$ $5_2^1aq_1$ — $B''_3C_1H_aO_a$ $6_2^1aq_1$ — $B''_3C_1H_aO_a$ $11aq_1$ — $B''_4C_1H_aO_a$ $11aq_1$ — B''_4C_4 $11aq_1$ — B''_4C_4 $11aq_2$ — B''_4 $11aq_3$ — B''_4 $11aq_4$ — B''_4 $11aq_5$ — A_5 A_5

Melly to-iodide B'Mel 8aq: lamino (from water); resinitied by boiling KOH (Stahlselmidt, P. 108, 513). — B'Mel, — B'Mel, — B'Mel, — Methy to-browide B'MeBr 23aq.—Methy to-chloride B'MeCl 5aq: v sol. water and alcohol.—B', Me, PtCl, — 3HAuCl, — Methy to-sulphate B', Me SO, 8aq: radiate crystalline mass, v. sol. water. B'Me, SO, 2aq.—Methy to-nitrate B'MeNO, 2aq. The physiological action of these salts has been studied by Crum Brown (T. E. 25).

Ethylo-iodide B'Etl laq: not decomposed by KOII but converted by moist Ag.O into a very seluble alkalino hydroxide:—B'Etl.—B'Etl. aq. — Ethylo-platino-chloride B'_Etl_PtCl₆. Silkyneelles (Gunning, J. pr. 67, 46).

Bromo-ethylo-bromide B'C,H,Br, 3aq: from brucine and ethylene bromide at 100°; forms fureate groups of nacreous lamine (from water), AgNO, pps. half the Br. Moist Agn

torms the vinylo.h.droxide B'O.H.OH which is strongly alkaline; two of its salts are B'2(C2H2Cl).PtCl, al.d B'C2H2SO.H8aq.—
Bromo-ethylo-platinpchloride
B'2(C2H3P).PtCl, (Schadt A. 118, 207).
Allylo-iodide B'O.I.Iaq.—B'C2H.I.—
B'C.H.I.aq.—Allylo-iolatinp-chloride

B'C₃H₃C₄B₄C₄D₄C₄D₅C₅D₄C₄D₅C₅D₄C₄D₅C₅D₆C₅

adding an alcoholic solution of Br to an aqueous solution of brucine sulphate. Small needles (Laurent, A. Ch. [3] 24, 314; cf. Beckurts, B.

18, 1238).

Nitro-brucine C23II25(NO2)O4N2. Formed by nitration of brucine methylo iodide suspended in absolute alcohol. Large red trigactric crystals. Carbonises at about 210° without melting; sol. water. Salts .- B'HNO2: glistening yellow needlos, v. sol. water, sl. sol. alcohol and ether. B'2H2Cl2PtCl4: fine yellow needles (Haussen, B. 19, 520).

Amido - brucine C23H24(NH2)O4N2. Formed by reduction of nitro-brucine with tin and HCl. V. sol. water. Not isolated. Fo₂Cl₆ produces a green colouration changing to brown. Very dilute K2Cr2O, produces a fugitive blue-violet colour. Strong IINO, gives a yellow solution which is turned red by SnCl.—B"H₃Cl₂: colourless prisms.—B"'₂H₄Cl₅Pt₅Cl₁₂*: yellow amorphous pp. (Hanssen, B. 19, 523).

Dinitro-brucine C., H., (NO2), O, N2. Prepared by the action of IINO3 on an alcoholic solution of brucino. Red amorphous powder. sol. water, sl. sol. alcohol, insol. ether. -(B'HCl)2PtCl4: yellow pp. (Claus a. Röhre, B.

14, 765).

BRYONIN C48H80O19. An amorphous bitter substance in the root of the red-berried bryony (Bryonia dioica) from which it finay be extracted by boiling water. It appears to be resolved by dilute H.SO, into glucose and two amarphons bodies, bryoretin C21H35O7 sol. ether, and hydro - bryoretin C₂,H₃,O₈ insol. ether (Brandes a. Firnhaber, Brandes Ar. Ph. 3, 356; Walz, C. C. 1859, 5).

BUCHU. The leaves of several species of Barosma growing at the Cape contain a volatile oil and a crystalline substance [85°] (Flückiger, Ph. [3] 4, 689; Jones, Ph. [3] 9, 673).

BURETTE v. Analysis, p. 248.

BUTALANINE v. AMIDO-BUTYRIC ACID.

n-BUTANE Call to i.e. CH3.CH2.CH2.CH3. Diethyl. Methyl-propyl. Butyl-hydride. Mol. w. 58. (1°) (Butlerow, 2. 1867, 363). S.G. 2 60. V.D. 2-11 (cale. 2-01) (flougills, C. J. 18, 54; Pelouze a. Cahours. A. Ch. [4] 1, 5). S. (gas in alcohol) 18 at 14°. Occurs in potroleum (R.). Formed by the action of Zn on EtI at 150° (Frankland, A. 77, 221; Schöyen, A. 130, 233). Also, together with butylene, by heating butyl aloohol with ZnCl2 (Wurtz, A. 93, 112). Prepared by the action of sodium-amalgam on EtI (Löwig, J. 1860, 397).

Iso-butane (CH_s)_sCH. Trimethyl-methanc, Secondary butane. (-17°). H.F. p. 42,450. H.F. v. 40,180 (Th.). Prepared by slowly pouring tert-butyl iodide into water containing zino, the equation being 2(CH₃)₃CI + H₂O + Zn₂ =2(CH₂)₂CH + ZnO + ZnI₂. Also from iso-butyl

iodide and Al₂Cl₄ at 120° (Köhnlein, B. 16, 562). Colourless gas. • Chlorine converts it into (CH₃)₄C.Cl (Butlerow, A. 144, 10). Br at 100° forms di-brome-butane (Carius, A. 126, 195). Excess of Br at 300° forms C₂Br₄ (Morz a. Weith, B. 11, 2019). B. 11, 2244) BUTANE CARBOXYLIC ACID. v. BUTYRIG

ACID.

Butane di-carboxylio acid v. Propyl-Malonio ACID, ISO-PROPYL-MALONIO ACID, METHYL ETHYLmalonic acid, Ethyl-succinic acid, two Di-methyl-succinio acids, Adipic acid, a-Methyl-olutario

acid, and B-Methyl-Glutaric acid.

Butane αωω tricarboxylic acid C₂H₁₀Q ic. CH₂CH(CO₂H)CH(CO₂H)₂. Ethyl thenyl. MeCH_CH(CO_H)CH(CO_H)_2. Ethyl . Thenyl tri-carboxylic acid. [119°]. Got by saponifying the ether with alcoholic potash (Polko, A. 242, 115). White rhombic crystals. V. e. sol. water, alcohol, ether, acetone, v. sol. chloroform.—Salts.—A".Ba, insol. water and alcohol.— A"Ag, 11aq, sl. sol. water.—A" CaH, insol. aleohol, v. sol. water. - A"CaH 2 laq. - A"2Ca,

bygroscopic.—A". Zn, 6aq, v. e. sol. water, isol. alcohol.—A". Sn, 6aq.

Ethyl cther McCll.CH(CO_Et)CH(CO_Et). (276°). (189° at 60 mm.). S.G. 17.1°065. Formed by acting on sodium malonic ether with a-bromo-

butyric ether in alcoholic solution (P.). Butane aaß-tri-carboxylic acid

CH₃.CH(CO₂H).C(CO₂H)..CH₃. Butenyl-tri-carboxylic acid. [157°]. V. sol. water, alcohol and ether. On warming it yields s-di-mothylsuccinic anhydrido [87°] (Bischoff a. Rach, A. 234, 54). Salts.—Ba, A"2,—Ca, A"2,—Ethyl ether Et, A'". (c. 190°) at 50-60

mm.; (273°-275°) at ordinary pressure. Liquid. Formed by the action of a-brome propionic ether upon sodio-methyl-malonio ether. By boiling with HCl it is converted into s-di-mothylsuccinic acid [189°] (Leuckart, B. 18, 2346).

Iso-butane tri-carboxylio acid

Mc_C(CO_II)CH(CO_II)... [120°]. Prepared by saponifying the ether with alcoholio potash (Barnstein, A. 242, 128). Needle-shaped crystals (from water). V. o. sol. water, alcohol, ether, acetone, v. sl. sol. chloroform, benzene, petroleum etbor, carbon disulphide. Decomposed by boiling with water, giving off CO. Salts.—K₂A'''2m₁. Quadratic prisms: v. e. sol. water, insol. alcohol. -Ca₂A'''₂9aq: m. sol. water.

-CaÁ''', 2aq: v. e. sol. water. —Sr₃A'''₂7aq.

Ethylether Et₃A'''. (181°-185°) at 30-40 mm.; (279-281°) at ordinary pressure (L.); (273°) (B.). S.G. ½7 1-061. Liquid. Formed by the action of a-brome-isobutyric ether upon sodio-malonic ether. By boiling with HCl it is converted into u-di-methyl-succinio acid [1390] (Leuckart, B. 18, 2350; Barnstein, A. 242, 126).

Butane $\omega\omega\gamma\gamma$ -tetra-carboxylic ether $C_{16}H_{26}O_4$ i.c. $(CO_2H)_2$. CH_2 . CH_2 . CH_3 . CH_4 . CH_4 . CH_4 . CH_4 . CH_5 . $CH_$ at 225 mm. Formed, together with trimethylene dicarboxylic acid, by the action of ethylene bromide on sodium malchio ether (Perkin, jun., C. J. 51, 17; B. 19, 2038). The yield is very small. The corresponding acid splits up when heated into CO, and adipio acid. By treatment with sodium ethylate it gives a di-sodio-derivative which by the action of bromine yields tetra-

CH₂.C(CO₂Et)₂ methylene-tetra-carboxylic other CH2.C(CO2Et)2

Iso-butane tetra-carboxylic ether (CO,Et),CH.CHMe.CH(CO,Et), (c. 211°) at 20 mm. A by-product in this preparation of ethylidene-malonic ether from aldehyde and malonio ether; it is formed by the addition of malonic ether to ethylidene malonic ether CH₂.CH:C(CQ_Et)₂ (Komnenos, A. 218, 158; Claisen, J. pr. [2] 35, 414). The corresponding acid is split up by distillation into CQ₂ and β methyl-glutarioaoidCO2H.CH2.CHMe.CH2.CO2H.

Butane aaß\$-tetra-carboxylie ether (CO2Et)2CMe.CMe(CO2Et)2. Di-methal-acetylenetetra-carboxylic ether. S.G. 19 1-114. From sodio-methyl-malonic ether by the action of ioane of chloro-methyl-malonic ether. From sodio-ethano tetra-carboxylic ether and McI (Bischoff a. Raoh, B. 18, 1202; A. 234, 70).

Butane tetra-carboxylic ether CH(CO₂H)₂.C(C₂H₃)(CO₂Et)₂. Ethyl-acetylenctetra-carboxylic acid. Thick colourless oil; formed by the action of chloro-malonic ether on sodio-othyl-malonic other (Bischoff a. Rach, B. 17, 2785).

Bntane hoxa-carboxylio ether

CO₂Et.CH₂.C(CO₂Et)₂.C(CO₂Et)₂.CH₂.CO₂Et. [57°]. Six-sided tables. From sodium ethane tricarboxylio ether (CO2Et)2CNa.CH .CO2Et by the action of iodino or of (CO,Et), CCI, CH, CO,Et (Bisohoff, B. 16, 1046; 17, 2786).

ISO-BUTANE SULPHINIC AUD

(CH3) CH.CH SO.H. From iso-butane sulphoohloride by zine-dust (Pauly, B. 10, 912). Liquid. Iteduced by nascent hydrogen to isobutyl mercaptan .- ZuA'2. Plates (from alcohol).

22-BUTANE SULPHONIC ACID C, 11, SO, II. Formed by the action of 11NO, on n-butyl meroaptan (Grabowsky, A. 175, 341). Chick syrnp, v. sol. water and alcohol, m. sol. ether. -AgA': plates. — PbA'₂: plates, sl. sol. alcohol. — PbA'₂,2Pb(OH)₂: crystallino powder, sl. sol. water.—BaA'₂aq: plates.—CaA'₂, 2aq.—NaA': plates.

Iso-butano snlphonic acid

(CH₃)₂CH.CH₂SO₃H. From iso-butyl mercaptan and 1INO₃ (Mylius, B. 5, 978). Syrup. Its salts are v. sol. water. - AgA': scales. - BaA'2: needles.

Chloride Call SO2Cl (190°) (Pauly, B. 10,

ISO-BUTENYL ACETATE CH,: CMe.Cl1,OAc (120°) (Scheschukoff, J. R. 16, 502).

BUTENYL ALCOHOL C411,0 i.c.

CH3.CH:CH.C112OII. (117°). A product of the action of iron aud acctic acid upon erotonic and tri-chloro-butyrio aldehydes (Lieben a. Zeisel, M. 1, 825). 111 gives CH₃.CH₃.CH₁.CH₁.CH₂.

Isobutenyl alcohol CH2:CMe.CH2OH. (113°). S.G. 2 8695. Formed by boiling isobutenyl chloride with vory dilute K₂CO₃ (Scheschukoff, J. R. 16, 499). Dilute H.SO, changes it to isobutyric aldohydo. HI forms tert-butyl iodide.

Ethyl ether CH2: CMe.CH4.OEt (78°-85°). From isobutenyl chloride and KOEt.

Sec-isobutenyl alcohol CMe.: CH.OH. Methyl ether CMe :: CH.O.Me. isocrotyl oxide. (70°-74°). From brome-iso-butylene and NaOMe at 140°. Dilute H,SO₄ at

140° givss HOMe and isobutyric aldohyde (Elte-keff, B. 10, 705; J. R. 9, 163). Ethyl ether CMo₂:CHOEt. (93°). Similarly prepared (E.; S.; Butlerow, Z. 1870, 524). BUTENYL-TRI-AM NE CH(CH2NH2)2.

BUTENYL-TRI-AM' NE CH(OH₂NH₂)₂.

(above 150°). From the nitrile of methane tricarboxylio acid CH(ON₂) tin, and HCl.—
B'₂3H₂PtCl₄ (Fairlie, Cl. J. 16, 362).

BUTENYL-BENZEL E v. PHENYL-BUTYLENE.
BUTENYL CHLOR² v. Chloro-Butylene.

(a) BUTENYL-CUTYENE. O.H-PrC, H., Isopropyl-butenyl-benzégé. (213°) S.G. 8875.

Obtained from brance annyl-valeric acid Chlyr CH. Chlory Co. H. and Parkin

Obtained from br no -annyl valeric acid C_aLi,Pr.CH_a. GBrEt.Cv_aHand N₁ CO_aAq (Perkin, C. J. 32, 662). Fo, as a dibror ide [77°].

(8)-Butenyl-outleno (14, H. Pr.C. | H distilled with water and then over solid KOII and Na (Perkin, C. J. 35, 141). It forms a liquid dibromide.

BUTENYL-GLYCERIN v. TRI-OXY-BUTANE. DI-BUTENYL-KETONE v. DI-ALLYL-ACETONE

(p. 134).

o-BÚTENYL-PHENOL.

Methul ether [2:1] MeO.C, II, CH:CH.CH, CII, (233°). S.G. 13 9817; 30 9740. From the methyl derivative of oxy-phenyl-angelic acid by successive treatment with III and Na₂CO₂Aq (Perkin, C. J. 33, 213). Oil. Combines with bromine. Forms a rod

solid with 11,80.

p - Butenyl - phenol. Methyl ether

[4:1] McO.C.,11,C.,14,... [17]. (213°). S.G. 50 1973. Prepared like the preceding (Porkin, C. J. 32, 671).

Salicylio aldehydo (30 g.) sodie isobutyrata (22 g.) and isobutyric anhydride (45 g.) are heated together for 4 hours. Water is added to the product, and the oily butenyl-phenyl isobutyrato saponified by alcoholic KOII. The alcohol is boiled off, dilute 11Cl is added and the oil which separates is distilled (Perkin, C. J. 35, 142). I roperties .- Oil, with smoky and cedarlike odour. Sel. potash but insol. NH3. With salicylic aldehyde and conc. H2SO, it forms a

purple solution. p-Iso-butenyl-phonol C₁₀H₁₂O i.e. C₁H₁,C₀H₁,OH. (230°-235°). From p-oxy-benzoic aldehyde, sodic isobntyrate and butyric anhydride. Yield small (Perkin, C. J. 35, 145) An oil which solidifies in a freezing mixture.

Meth / ether C. H. C. H. OMe. [9°]. (237°). From misic aldehydo (20 g.), sodic isobutyrate (15 g.) and isobutyric anhydride (30 g.) by boiling at 180°. The product is distilled with steam and the oil defedover K.CO., Cooled by ico and HCl it solidities. On oxidation it yields anisic and accide acid.

. BUTENYL - STYRENE C.II, CH:CH.C.H. (245°-248°). Pheny! hexinenc. Cinnamic aldc. hyde (10 g.), sodium isobutyrate (15 g.) and isobutyrio anhydrido (10 g.) are heated together for two hours. CO, comes off, and the residue is distilled with steam, washed with NH₂, dried and rectified over Na (Perkin, C. J. 35, 141).

Properties.—Lighter than water. Rapidly oxidises in air, becoming a rosin. It combines with bromino.

Iso-BUTENYL-TOLYLENE-o-DIAMINE

C,H,
NII
C.C,H,. [158°]. Formed, together with small quantity of tolu-

butyraldehydine C.H. N:OH.C.H., by shaking a cold acetic acid rolution of tolylene-odiamine (1 mol.) with in aqueous or alcoholic solution of butyric aldehyde (2 mols.). Small colourless needles. V. 551. alcohol and other, al. sol-water. Very bit is taste (Hinsberg, B. :**20, 1**589).

BUTINENE C.H. P. CH.: CILCH-CH.
Erythrene. Vinyl-cth the Pyrrolylene.
Ocenre in the liquid got by ampressing coal-gas
(Caventou, B. 6, 70; Grima va a. Clocz, C. R.
104, 118). Formed also (?) by passing fusel oil through a red-hot tube (Caventou, A. 127, 93). Obtained by boiling crythrite with conc. formic acid (5 pts.) (Henninger, B. 6, 70); and by the action of KOII on di-methyl-pyrrolidine methylo-iodide (Ciamician a. Maganghi, B. 19, 569). A gas. It forms a tetrabrounde [119'].
Pyrrolyleuc and the butinene from crythrite form also a second tetrabromide [40°] so that they are probably mixtures of two butinenes.

Butinene CH. CIL C:CH. Ethyl-acctylene Crotonylene. (184). From methyl ethyl ketone by encessive treatment with alcoholic KOII and PCl_s (Bruylants, B. 8, 412). Gives a white pp. in PO₃ (Bruylants, B. 8, 412). Gives a write pp. in ammoniacal AgNO₂ and a yellow pp. in aumoniacal cuprous chloride. Mercuric chlorido solution gives a pp. of (C₁11_{...})₄(figOHgCl₂)₄ (Kutscheroff, B. 17, 21), where IfO produces methyl ethyl ketone. The same buttuene appears to be formed when a mixture of acctylene and ethyleno is passed through a rod hot tube. It forms a tetrabromide [113] (Berthelot, A. Ch. [4] 9, 406; Prunier, Bl. 20, 72; A. Ch. [5] 17, 17; C. R. 76, 1410).

Butinene CII, C:C.CII, (?). Crotonylene. (18°). From crude butylene bromide and alcoholió KOII (Caventou, Å. 127, 317). From aß-di-bromo-butane and alcoholic KOII (Almedingen, J. R. 13, 892). Also formed by distilling barium acctate with S (Pfankuch, J. pr. [2] 6, 110). 11.804 (3 mols.) diluted with water (1 mol.) converts it into hexa-methyl-benzene.

Butinene C, H_s. Caoutchin. [-10°]. (15°). S.G. = 2°65. Formed by the dry distillation of caoutchouc (q. v.) (Bouchardat, A. 27, 33).

BUTINENE GLYCOL v. DI-OXY-BUTYLENR. BUTONENE-AMIDO-PHENYL MERCAP-TAN C₁₆H₁₂N₂S₂ i.e.

 $C_0H_1 < \frac{N}{S} > C_0CH_2 \cdot CH_2 \cdot C < \frac{N}{C} > C_0H_1 \cdot C$ f137°1. From amido-phenyl-o-mercaptan and succinamide (Hofmann, B. 13, 1231). Needles (from

ulcohol).—B'HAuCl.
BUTYL DERIVATIVES of hydroxylic cempounds are described under the compounds of

which they are the others.
DI-BUTYL v. OCTANE.

BUTYL ACETATE C.H. O. i.e. C.H. OAc. (124-5°). S.G. § 9016. C.E. (0°-10°) 00113. S.V. 150 6 (Gartenmeister, A. 233, 259). Fran butyl iodide and AgOAc (Lieben a. Rossi, A. 158, 170; Linnemann, A. 161, 193; Pribram a. Handl, M. 2, 693).

Isobutyl acetate (CH₃)₂CH.CH₂.OAc. V.D. 4-073 (calc. 4-017). (110-3°) (Eleisser, A. 17-7°.

218, 326); (117° cor.) (Perkin, C. J. 45, 495; (112-8°) (R. Schiff, A. 220, 109). S.G. § *8921 (E.); \$\frac{15}{15}*8774; \$\frac{25}{25}*8688\$ (P.). C.E. (0°-10°) 129; A. 85, 197; 93, 107; \$\mathcal{Q}\$. R. 85, 810).

·001137 (E.). M.M. 6·623 at 10°. S.V. 150·10 (E.); 152.5 (S.). Formed from isobutyl iodide and AgOAc, or by distilling potassium isobutyl sulphato with KOAc (Wurtz, A. 90, 121).

Sec-Butyl acetate CH. CH. CH (OAc). CH. (112°). S.4.2.892. From sec-butyl iodide and AgOAc (De Luynes, J. 1864, 501; Lieben, A. 150, 112).

Tertiary Butyl acetate (CH3)3C.OAc. (93°-96°). From the iodide and AgOAc. Readily saponified by baryta-water (Butlorow, A. 144, 7).

1SO-BUTYL-ACETIC ACID v. HEXOIO ACID. ISOBUTYL-ACETO-ACETIC ETHER v. p. 24. BUTYL-ACRIDINE C1711,7N i.e.

 $C_{\epsilon}\Pi \underset{N}{\overset{C(C_{\epsilon}H_{0})}{\triangleright}} C_{\epsilon}\Pi_{\epsilon}. \quad \text{From valeric acid-130 g.), dis}$

phenylamine (30 g.) and ZnCl. (50g.) heated gradually for 20 hours up to 220' (Bernthsen a. J. Traube, A. 224, 41; B. 17, 1508).

Salts .- B'HCl [191°]: yellow columns, v. sel. water or alcohol, very dilute solutions show bluefluorescence. Insol. ether. -B'11NO, [139°]. -B'H2CrO4 [c. 100°].

Dihydride C, H, N(C, H,). [98°-100°]. Got by reducing the hydrochlorido with zino-dust. White plates (from alcohol).

BUTYLACTIC ACID v. OXY-BUTYRIO ACID.

BUTYLAL v. BUTYRIC ALDEHYDE.

BUTYL ALCOHOL C. H10 (). Mol. w. 74. The four butyl alcohols indicated by theory are known.

n-Butyl-alcohol CH, CH, CH, CH, OH. (117° cor.). S.G. % 8233 (Zander, A. 221, 79); 20 8096 Brühl, A. 203, 16). S. 8·3. C.E. $(0^{\circ}-10^{\circ})$ 00907 (Z.). μ_{β} 1·4040. S.V. 101·6 (Schiff, A. 220, 101). R $_{\odot}$ 35·45 (B.). Critical point 287° (Paulewsky, B. 16, 2634).

Occurrence. - In the heavy oils from brandy. It is completely absent from the products of the fermentation of sugar with elliptical yoast

(Claudon a. Morin, C. R. 101, 1187).

Formation .- 1. From butyrylculoride, butyric acid and sodium-amalgam (Saytzeff, J. pr. [2] 3, 76). 2. By the fermentation of glyceria by a Schizomycetes or by certain Bacteria in presence of CaCO, and ammonium tartrate; nbutyric acid and a little alcohol are formed at the same timo (Fitz, B. 9, 1348; Vigna, B. 16, 1438).-3. A product of the reduction of butonyl alcehol (Lieben a. Zeisel, M. 1, 825).

Preparation. - From butyrio aldehyde, water, and sodium-amalgam (Lieben a. Rossi, C. R. 68, 1561; 78, 1561; A. 151, 121; 158, 137; 165, 145; C. J. 21, 516; Linnemann, A. 161, 179).

Properties.—Liquid. Separated by CaCl₂ from aqueous solution. On oxidation it produces butyric acid. Fused ZuCl, forms the two n-butylenes (Le Bel a. Greene, C. R. 89, 413).

Combination. - (C4H10O)3CaCl2 (Heindl, M.

lso-butylalcohol (CH₃)₂CH.CH₂OH. (106·6°-106·8°) at 763·2 mm. (R. Schiff, A. 220, 102) (100°c) at 763°2 mm. (R. Schiff, A. 220, 102) (1698°) (Linnemann, A. 160, 238). S.G. Ψ 8062 (Brühl); $\frac{15}{15}$ 8069; $\frac{25}{15}$ 8009 (P.). S. 9.5 at 18°. S.V. 101°63. H.F. p. 71,150. H.F. v. 68,580 (Th.). $\mu_{\rm B}$ 1 4007. R $_{\odot}$ 35°41. M.M. 4°936 at 17.7°.

yl angelate and isobutyrate occur in Roman

oil of chamonile (Köbig, A. 195, 96).

Formation.—1. Isobutylene combines with ClOH forming (OH₃)₂:CCl.CH₂OH, which is reduced by sodium amalgam and water (Butlerow, A. 144, 24).—2. Isobutyl alcohol is produced by the action of Bacillus butylicus Ipon sugar, glyoerin, &c., even in presence of 8 p.c. alcohol (Ordonneau, C. R. 102, 219; Claudon a. Morin, C. R. 104, 1187).

Properties .- Liquid; smelling like fusel oil; separated from its aqueous solution by CaCl. The rate of etherification has been studied by

Menschutkin (A. Ch. [5] 23, 14).

Reactions.—1. GrO, produces isobutyrio, costic ad carbonic acids, and acctone (Krämer, B. 7, 252; Schmidt, B. 7, 1361).—2. Distilled over zinc dust splits up into isobutylene and H₂O (Jahn, B. 13, 989).—3. Dropped upon fused ZaCt, it forms isobutylene and CH₃OH:CILCH₃ (Le Bel a. Greene, Am. 2, 23). 4. Heated with ammoniacal ZnCl, at 260°-280° it yields a mixture of mono-, di- and tri- isobutylamino (the latter in smaller quantity), the yield of mixed bases amounting to 50 to 70 p.c. of the alcohol used (Merz a. Gasiorowski, B. 17,

Combinations. - (C, H100) CaCl2 (Heindl, Jl.

2, 208).

Metallic derivatives. — KOC_1H_0 . — $NaOC_1H_0(C_1H_0O)_3$ (De Forerand, C. R. 104, 169).— $\Delta L_0(OC_1H_0)$. [140°]. S.G. ± 9825 (Gladstone a. Tribe, C. J. 39, 6). From $\Delta L_1(DC_1H_0)$. iodine (2 g.) and isobutyl alcohol 40 c.c. at 100%. The yield is good (16 g.). Once fused it remains long fluid at 70 .

Secondary butyl alcohol

CH3.CH2.C(CH3)II.OH. Methyl- hyl-carbinol. **Butylene hydrate. (99°) at 740 mm. S.G. °
**827 (Licben, A. 150, 114).

**Formation. — I. From n-butylamine by

nitrous acid; at the same time some n-butyl alcohol is also formed (Linnemann a. Zotta A. 162, 3; Meyer, B. 10, 130).-2. By treating the compound of HClO with isobutylene, OH₃.CHCl.CH(OH).CH₃, with sodium amalgam (Lieben, A. 151, 121). -3. By the action of zine ethido on glycolic iodhydrin (Butlerow a. Ossokin, A. 145, 263).—4. Symmetrical dichloro-di-ethyl oxide, (CH₃, CHCl)₂O is converted by ZuEt₂ into (CH, CHEt) O, a butyl ether which, on treatment with 111 at 130 gives secondary butyi iodide (Lieben, A. 141, 236; Kessel, A. 175, 11). 5. Zine ethide forms a crystalline compound with aldehyde, CH_CHEt(OZnEt), which is decomposed by water into secondary butyl alcohol, ethane, and Zu(Oil) ... (Wagner, J. R. 8, 37; A. 181, 261) .- 6. From formic other by treating with a mixture of ZnEt, and ZnMc, and decomposing the product with water (Kanonnikoff a. Saytzeff, A. 175, 374).

Preparation.—Secondary butyl iodide (q. v.)

is treated with silver acctate rend the product saponified by potash (Do Luynes, A. 128, 330;

132, 274).

Properties. - Liquid with strong odour, ppd.

from aqueous solution by K2CO3.

Reactions .- 1. Oxidises to methyl-ethylketone and acetio acid (Saytzeff, Z. 6, 327).-2. The pure alcohol is unaltered when heated at but the presence of histrace of HCl, HBr, or especially HI, is suffice at to split it np, forming TI reaction commences at pseudobutyleno.

220° and is complete 5. 5 or 6 hours (Bongaieff a. Wolkoff, Bl. [2] 45 f. 9).

Tertiary butyl ald hol (Cll.) 0.OH. Trimethyl-carbinol. [25° (83° cor.). S.G. 27.779 (Linnemann); 22 .77 (Butlerow); 20 .786; 24 · 780 (Brühl); 15 · 886; 25 · 7761 (Perkin), M.M. 5·122 at 24·3° , 1·8:21. R & 35·53. Cri-tical point 235° (Pav. ewski. B. 16, 2634).

Formation. -1. inc methide (2 mols.) and acelyl chloride (! mol.) mixed at 0', form, after some hear, a crystalline contound, CH, CMe(OZnMe) sie, which is decomposed by water into tertiary butyl alcohol, Zn(OH), and CH, (Butlerow, A. 141, 1; Wagner a. Saytzeff, A. 175, 361; Pawloff, A. 188, 118). - 2. Together with isobutyl alcohol by treating isobutyl iodide with acetic acid and Ag₂O (Linnemann, A. 162, 12; Butlerow, A. 168, 143). - 3. From isobatylamine and HNO, -4. From isobutyl evanate and KOH (Linnemann, A. 162, 12) .- 5. From tertiary butyl iodide (q.v.) and water, even in the cold (Dobbin, C. J. 37, 238).-6. A mixture of iso- and tert- butyl chlorides is got by beating isobutyl alcohol with HCl; when heated with water (6 vols.) at 100° the chloride of tert-butyl alcohol is the only one converted into its alcohol (Freund, J. pr. [2] 12, 25).

Preparation.-Liquid isobutylene is scaled up with twice its volume of a mixture of equal parts of water and sulphuric acid, and the contents are left till homogeneous and then distilled

(Butlerow, Z. [2] (237; A. 180, 246). Properties. — Trunctric prisms. Forms & hydrate (C,II,0O)2I12O which boils at 80° (But-

lerow).

Reactions. -1. CrO_a mixture gives acctone together with acetic, carbonic and a little isobutyric acid (Butherow, Z. 1871, 485) .- 2. Heating with arhydras H₂C₂O₄ produces butylene (Cahours a. Demarçay, C. R. 86, 991). -3. When taken internally it is excreted in the urino as butyl-glycuronic acid C₁₀H₁₈O, (Thierfelder a. Mering, H. 9, 514) which is decomposed by boiling dilute IICl into tert-butyl alcohol and glycuronic acid.

DI-ISOBUTYL ALDEHYDATE v. p. 105.

ISO-BUTYL ALDEHYDE v. 1sobutyric alde-

ISO-BUTYL-ALDOXIM v. ISOBUTYRIC ALDE-HYDE.

ISO-BUTYL-AMIDO-ISO-BUTYL-BENZENE $C_4\Pi_g$ NH. $C_4\Pi_4$. $C_4\Pi_g$. (260° 270°). From anilino hydrochloride (10 g.) and iso butyl alcohol (13 g.) at 230° (Studer, A. 211, 210; B. 14. 1473). Oil. Does not give the carbamine reac-

Nitroso-derica tive S.G. 24 . 991. Solidifies on keeping.

Acetyl-derivative. • [74°]. (above 300°).

Needles (from benzoline).

BUTYL-AMIDO.TOLUENE v. METHYL-BUTYL-PHENYL-AMINE.

n BUTYLAMINE C.H.NH. i.c. Ir.CH.NH. Mol. w. 73. (76°). S.G. 2.755, 22.733.

Formation.—1. From butyl cyanato and

KOHAq (Lieben a. Rossi, A. 158, 172; Meyer, 240°-250° during 8 to 16 hours in a scaled tube, | B. 10, 131).-2. From butyronitrile by reduction

(Linnemann a. Zotta, 4, 162, 8).—3. From nitrobutane, Sn and Hd. (Züblin, B. 10, 2093).

Properties.—Miscibl, with water; dissolves freshly ppd. Cu(OH) and Ag.O. Reduces alkaline solutions of coppe, silvor, and mercury.

Nitrous acid converts it tuto see-butyl alcohol.—
Platino-ohlorido (N. HCl) _PtCl.; yellow crystalline plates, n. sol. Ald water.

Zrimary isobutyl-amil e Pr. CH_NNIL. (682)

R. Schilf, B. 19, 565. S. 12, 736. S. V. 106: 16

(R. Schiff, B. 19, 565). S., 12 736. S.V. 106:16 (S.). H.F. p. 38,460. 147. v. 35,560 (Th.). Formation.—1. By dist ing potassium isobutyl sulphate with potassium evanato and treating the product with KOH (Viirtz, A. 93, 124; Linnemann, A. 162, 23).—2. By heating isobutyl sulphate with kohl (Viirtz, A. 93, 124; butyl bromide with alcoholic NII, and separating the mono-, di-, and tri- butylamine; by exalic ether (Reimer, B. 3, 756; Hughes a. Römer, B. 7, 511; Malbot, C. R. 104, 63, 228; On heating isobutyl chloride with ammonia (molecular proportious) in isobutyl alcoholio solution or in aqueous solution very nearly similar results are obtained; namely one part of mono-butylamine, four of di-, and five parts of tri-butylamines (M.) .- 3. Formed, together with di- and tri- isobutylamine, by heating isobutyl-alcohol with ammoniacal ZuCl. at 260 -280; the yield of mixed bases amounts to 50-70 p.c. of the alcohol used (Merz a. Gasiorowski, B. 17, 623). 4. By reducing nitro-isobutane (Demole, A. 175, 142).-5. A mixture of equal mols. of valeramide (ordinary) and bromine is run into an excess of a 10 p.c. solution of KOH at 60°; the

yield is 90 p.o. (Hofmann, B. 15, 769). Salts.—B'HCl. [160°]. S. 133 at 15°.— (B'HCl) PtCl4: microscopio rhombic tablets. B'HAuCl.—Sulphate: cauliflower-like groups

of needles, not deliquescent.

Secondary butyl-amine (63°). CH₃.CH₂.CH(NH₂).CH₃. From dilute H.SO, and sec-butyl thio-carbinide from volatile oil of sourvy-grass) (Koyman, B. 7, 1289). Also from the iodide or cyanate (Hofmann, B.

Also from the form.
7, 513).—B'_4II_PtCl_a.
Tertiary butyl-amine Mo_CNII_ (46° cor.).
C.E. (0°-7-8°) 0014. Formed together with isobutylamino by the successive action of silver eyauate and aqueous KOH on isobutyl iodide. Colourless ammoniacal liquid, attacks indiarubber and cork. Miscible with water, but separated by K2CO3 or KOH from its solution (Brauner, A. 192, 72; cf. Linnemann, A. 162, 19; Hofmann, B. 7, 513).

Salts .- B'IICl melts at [-70°-280°] and boils soon after. On solidifying it increases greatly in bulk .- (B'HCl), PtCl, Large monooliuio prisms (from alcohol) -B'HI. B'HNO. Sulphate: six-sided prisms, not deliquescent.

 $\hat{\mathbf{Di}} \cdot n$ -butyl-amine (Pr.CH₂)₂NH. (160°). Formod, together with n-butylamine, by treating butyl cyanate with KOH (Lieben a. Rossi, A. 158, 175). Converted by nitrous acid into primary and secondary n-butyl alcohols and n-butylene (Meyer, B. 10, 130).—B'uH₂PtCl₂. Nitroso derivative (C₄H_e)₂N.NO. (236° cor.). Di-iso-butyl-amine (Pr.CH₂)₂NH. (137°).

Formed, togother with mono- and tri- iso-butylamine, by heating iso-butyl alcohol with ammoniacal ZnCl2 at 260°-280°. The secondary amine is isolated from the mixture of bases (which amounts to 50-70 p.o. of the alcohol soluble in water (Kahn, B. 18, 3365).

used) by means of its nitrosamine (Merz 2. Gasiorowski, B. 17, 623). Prepared by heating iso butyl iodide or bromide with alcoholic NH, to 150° (Ladenburg, B. 12, 948). Butyl iodido (1 mol.) in the cold acts upon di-isobutylamine (1 mol.) forp ing di-isobutylamine hydrochloride and free triviso-butylamine (M.).

Salts. -B'HCl: plates or scales. S. 62.5 at 15°; S. (ether) 07 at 15°; S. (alcohol) 06 at 14° (Malbot, C. R. 104, 366). — B'₂H₂PtCl₆. — B'HClAuCla. Yellow tables, sparingly solublo

in cold water.

Nitrost- derivative N(NO)(C,H,)2. [0°]. (213'-216°). Oil. Prepared by the action of KNO, on a solution of di-isobutylamino hydrochloride.

Tertiary di-butyl-amine (C,H,)2NH i.e. (CMe,)NH. From tertiary butyl iodide and tertiary butylamine at 50°. But above 70° isobutylene is given off: C,H,NH, + C,H,I =

C₄H₄ + C₄H₅NH₅,HI (Rudnew, Bl. [2] 33, 299). Salt.—B'Hl. Very soluble in water and alcohol. Converted by potash, or even boiling

water, into tertiary butÿlamine.

Tri-n-butyl-amine $(3,1\mathbb{I}_9)_4$ N. Mol. w. 185. (c. 213° cor.). S.G. 2–791; 22–778. From n-butyl iodide and NH $_3$ (Lieben a. Rossi, A. 165, 115).—B'.H.PtCl.,

n.Butylo-iodide (C,II,),NI: plates (Lie-

beu a. Rossi, A. 165, 113).

Tri - isobutyl - amino (PrCH2), N. S.G. 21 785 (Sachtleben, B. 11, 733). Formed, together with mono- and di- iso-butylamine, by heating isobutyl alcohol with ammoniacal ZnCl at 260°-280°. The tertiary amine is isolated from the mixture of bases (yield 50-70 p.c. of the alcohol) by means of its sparingly soluble ferrocyanido (Merz a. Gasiorowski, B. 17, 623). Also from di iso-butylamine and isobutyl bromide. Also from isobutyl iodide (1 mol.) and NH₃Aq (1 mol.) at 160° (Malbot, C. R. 105, 575). Does not mix with water. With isobutyl bromide it gives off butylene and forms tri-isobutylamine hydrobromide (Reimer, B. 3, 757). Isobutyl iodide (I mol.) at 180° forms hydriodides of di- and tri-isobutylamine and butylene (M.). Isobutyl chloride (I mol.) at 170° gives puro di-isobutylamine hydrochlorido and butyl-

Salts.-B',HCl, B',HNO, and B',,H2SO, are oxtremely soluble.—(B'IICl) PtCl, orange plates, sol. hot water. - B'HClAuCla: amorphous, insol. water.

Tert - BUTYL - tert - AMYL . AMINE

(C,H,)(C,H,)NII. The iodide of this base is formed by the slow action of tertiary amyl iodide on tertiary butylamine in the cold. It is vory unstable, being decomposed by solution in water (?) (Rudnell, Bl. [2] 33, 297).

n - BUTYL - ANILINE C₁₀H₁₀N C₆H₅.NHC₄H₉. (235° at 720°). Colourless oily fluid. Easily volatile with steam.

Salts .- B'HCl: very soluble white needles. -B'HNO,*: easily soluble.—B',H,SO,*: easily soluble fine white ucedles.

Acetyl derivative $C_6H_5.N(C_4H_9)Ao$: (274°) at 718 mm., colourless fluid.

Nitrosamine CoH3.N(C,H0)NO: yellow fluid; easily soluble in alcohol and ethor, in-

Isobutyl-aniline Pr.CH, NHPh. (242°) (G.); (226°) (N.). S.G. 11 926 (G.). From isobutyl bromido and aniline (Gianotti, G. 12, 268). B'HCl.-B'HBr.-B'HI.

Acetyl derivative (267°) (Nölting, J. 1883, 703).

p-Nitroso-derivative

[44] C₆H₄(NO).NHCH₂Pr or C₆H₄ NH.CH₁Pr [94°] France

[94°]. From isobutyl aniline, HCl, and NaNO₂ (Wacker, A. 243, 297). Steel-blue crystals, v. sol. alcolol, sl. sol. water. Reactions.—1. Reduction gives isobutyl-phenylene diamine. - 2. The chlorthe boiled with aqueous NaOH gives iso-butylamine and p-nitroso-phenol.—3. HCl and NaNO, givo a nitrosamine Calla(NO).N(NO).CHaPr crystallising in bright green plates, v. sol. alcohol and ether, insol. water.

Di-isobutyl-anilino (PrCH2)2NPh. (c. 218'). From auiline and isobutyl bromide (Studer, A. 211, 235).

DI-BUTYL-ANILINE-AZYLINE v. Di-butylamido-benzene-Azo-di-butyl-aniline.

BUTYL-ANISOL v. Methyl ether of Butyl-

ISO - BUTYL - ANTHRACENE Cisting i.e. Fluorescent

needles. Prepared by the action of zine-dust, isobutyl bromide and NaOH on authraquinone (Liebermann a. Tobias, B. 14, 802; A. 212, 107). The picric acid compound forms long brownish-red needles.

Di-hydride C6H4 CH CH CHI, From isobutyl-oxanthranol, III, and P (i.). Oil; oxidised by CiO3 in HOAc to isobutyl-exauthranel and finally to anthraquinoue.

BUTYL-ANTHRANOL-DIHYDRIDE

 $C_6H_4 < \frac{CH(OH)}{CH(C_4H_9)} > C_6H_4$, [72°]. From an hraquinone, aqueous NaOII and zinc-dust, boiled for some time and then iso-butyl bromide added

(Liebermann, A. 212, 103).
ISO-BUTYL-ANTHRANYL CHLORIDE

 $C_6H_4 < CO(C_4H_9) > C_6H_4$. [78°]. Tables. Prepared by the action of PCl₅ on isobutyl-exauthranol (Liebermann a. Walder, B. 14, 463).

n-BUTYL-BENZENE C₁₀H₁₄ i.e. C_aH₅.CH_aPr. Phenyl-butane. (180'). S.G. ¹² 862. From n-propyl bromido, benzyl chloride, and Na (Radziszewski, B. 9, 260). Also from n-butyl bromide, bromo-benzenc, and Na (Balbiano, B. 10, 296).

Bromination.-By the action of bromine in the dark, or in presence of iodine, the product is probably a mixture of o- and p- brome-butylbenzene. By the stom of bromme in direct sunshine, the substitution takes place in the γ -position of the side-chain giving C_aH_a .CIIBr. C_aH , or CaH3.CBr2.C3H7. If the γ-mono-bromo-derivative is further brominated in the dark at 100° the second Br atom probably enters the & position, the product being identical with the butylene-benzene-dibromide [70°] of Radziszewski (Schramm, B. 18, 1275).

ISO-BUTYL-BENZENE CaH CH2Pr. (167°). S.G. 2 ·880 (G.); ·858 (R.). V.D. 4·72 (C.).

Formation.—1. Fr m isobutyl bromide, bromo-benzene and a (Rioss, B. 3, 779; Wreden a. Znatowicz B_A 9, 1606).—2. From benzyl chloride, isopro-yl fodide, and Na (Köhler a. Arophein, B. 8, 54).—3. By the action of 50 g. iso-butyl ohlor to on 150 g. benzene in presence of about 50 x. Al Cl. (Hossin B. 193). presence of about 5% g. Al.Cl. (Gossin, Bl. [2] II, 446).—4. By he, and benzene with isobutyl alcohol and ZnCl₂ ? Joldschmidt, B. 15, 1966). 5. My distilling more prescoutyl-benzoic acid with lime (Kelbe a Reiffer, B. 19, 1728).

Properties, -C. ourless liquid; CrO₃ oxidises it to benzoic acic Passed over red-hot PbO it forms naphthale e.

Sec-butyl-benzene CH₃,CH₂,CH(C,H₃),CH₃, (171°). S.G. ¹⁰ S73. From C.H.,CHBr.CH₃ and ZnEt, (Radziszewsky, B. 9, 261)

(a)-n-BUTYL-BENZENE SULPHONIC ACID CaH, (CH, PASO, H. Formed by sulphonating n-hutyl-benzene (Balbiano, G. 7, 343). -BaA'2: small lamine, st. sol. cold water. - ZnA', 7aq. -PbA aq. -- MnA', 6aq.

(β)-n-Butyl-benzene sulphonic acid. Formed at the same time as the preceding (B.) .-BaA'_22aq: nodules, more soluble than the Ba salt of the (a) acid. -PbA'. 2aq.

Iso-Butyl-benzene-sniphonie acid

C₆H₄(C₄H₉).SO₄H. Formed by subphonation of isobutyl-benzene. A' Ba 2aq : glistening plates. A'K aq : plates.

Amide C₆H₁(C₁H₂).SO₂NH₂: [137°]; glistening needles (Kelbe a. Pfeiffer, B. 19, 1728).
BUTYL BENZOATE v. p. 470.

BUTYL BENZIMIDO-ETHER v. p. 479. m-ISO-BUTYL-PENZOIC ACID

 $C_4H_1(C_4H_9)CO_2H$ [1:3]. [127]. Long stout needles. Formed by oxidation of m-isobutyl-toluene with dilute HNO_3 . By further oxidation with dilute HNO_a at 170° 200° isognthalic acid is formed Cives a nitro-derivative [140°] .--

AgA': white 1:p.

Amide C_aH₁(C_e C_aONH₂: [130°]; hair-fine needles from water (Kelbo a. Pfeiffer, B. 19, 1725)

p-Isobutyl-benzoic acid Calla (Calla).COall [1:4]. $[164^{\circ}]$.

Formation.-1. By oxidation of p-isobutyltoluene with dilute HNO₃₀. By further oxidation with diluto IINO₃ terephthalic acid is formed (Kelbe a. Pfeiffer, B. 19, 1725). 2. By saponification of its nitrile (Pahl, B. 17, 1237).

Properties. - Monoclinic crystals; gives a

nitro-derNutive [16.10].
Sults. — AgA': white flocculent pp. — BaA' ang: plates, sol. hot water .-- CaA' a xaq: sl. sol. cold water.

Amide C, II, (C, II,). CONH2: [1710]; long

hair-tine needles (from water).

Methyl ether MeA': (247°); oil.

Nitrite C.I., (C.H.), CN. (249°) (G. a. M.);
(244°) (K.); (238°) (P.). V.D. 547 (obs.) (K.);
535 (obs.) (P.). Colouress oil. Formation.— L By distilling the formyl derivative of isobutylphenyl-amine with zinc-dust; the yield is about 12 p.c. (Gasiorowski a. Merz, B. 18, 1009).-2. By heating p-isobutyl-phenyl-thio-carbimide with copper powder at 200° (Pahl, B. 17, 1236). 3. Formed by heating tri-isobutylphenyl-phos-phate with dry KCN (Kreysler, B. 18, 1707).

ISO-BUTYL-BENZOYL-ACETIC ETHER v. p. 482.

ISO-BUTYL BORAT. B(OC.H.). (212°). Formed by heating B. (7, with isobutyl alcohol for 8 hours at 170° Hirns with green flame. Insol. water and slowly decomposed by it (Councier, J. pr. [2] 18, 382). Not acted upon by ammonia.

n-BUTYL BROMIDE 7,H Br i.e.
CH,CH,CH,CH,CH,Br. (101 cor.). S.G. 21:305;
22 1 299. From n-butyl alcohol, Br, and P
(Lieben a. Rossi, A. 153, 161). With Br at
150° it gives C,H Br (160°) (Linnemann, A. 161, 199). With bromine containing iodine at 250° it reacts thus C.H.Br + 8Br, = 2C.Br, + 9HBr (Merz a. Weith B. 11, 2244).

Isobutyl bromide Pr.CH.,Br. (92°) (L.); (91°) at 758 mm. (Schiff, B. 19, 563). S.G. \(\frac{1}{2}\)? 12722; \(\frac{2}{3}\). 12598 (Perkin, C. J. 45, 457). S.V. 11039. M.M. 8003 at 16° (P.). From the alcohold of the control of the hol and III or P and I (Pierre a. Pachot, J. Ph. [4] 13, 9; Wurtz, A. 93, 114; Chapman a. Smith, C. J. 22, 153). At 220° it partially changes to tertiary butyl bromide (Eltekoff, B. 8, 1241).

Tertiary butyl bromido CMc₃f. (72°). S.G. 2º 1·215; ½ 1·2020; ½ 1·1892. V.D. 4·7 (obs.). M.M. 8·238 at 18°. From isobutylene and IIBr (Rozeboom, B. 14, 2396). From tri methyl envising and Phys. (Phys.) 4.7 (2°). tri-methyl carbinol and PBr, (Reboul, C. R. 93,

Reactions. -1. Readily decomposes into HBr and isobutylene. This occurs when it is treated with Ago, with KHO, with NEt, or with Za and water (Butlerow, Z. 1867, 362) .-- 2. With ZnO it forms tri-isobutylene, C, H, -3. With water, even in the cold, it forms tertiary butyl alcohol.

BUTYL BUTYRATE v. BUTYRIC ACID.

ISO-BUTYL CARBAMINE C. II, NC. (c. 116°). S.G. 2 787. Preparation and properties like those of ethyl carbanine. Less readily attacked by HCl than cthyl carbamines (Gautier, A. 152, 221; Bl. [2] 11, 211; Z. [2] 5, 445).
n - BUTYL CARBOPATE (PrCП_s), CO₃.

(207° cor.) at 740 mm. S.G. 2 941 (Lieben a.

Rossi, A. 165, 112).

Isobutyl carbonato (l'r.CH.), CO3. eor.). S.G. 15 919. From isobutyl iodide and Ag₂CO₃ (De Clermont, A. Ch. [3] 44, 336). From isobutyl alcohol and CyCl (Humann, A. Ch. [3] 44, 340). From sodium isobntylate and chloropierin (Rese, A. 205, 253; cf. Wurtz, A. 93, 119).

Isobutyl-orthocarbonate (PrCII,O),C. (245° cor.). S.G. 900. The chief product of the action of chloropicrin on sodium isobutylate (Rose, A. 205, 253).

BUTYL-CHLORAL v. Tri-chloro-butyrio ALDEHYDE.

n-BUTYL CHLORILE CAHACI i.c.

CH, CH, CH, CH, Cl. Mol. w. 92.5. (78° cor.). S.G. 2 .907; 14 .897 (Linnemann, A. 161, 197). S.V. 114.3 (Ramsay). From n-butyl alcohol and IICI (L.; cf. Lieben a. Rossi, A. 158, 161). From n-butane and chlorino (Schöyen, A. 138, 235).

Isobutyl chloride (CH₃)₂CH.CH₂Cl. (69°). S.G. ¹⁵ 980 (Limemann); \$\frac{1}{2}\cdot 880 (Limemann); \$\frac{1}{2}\cdot 880; \$\frac{1}{2}\cdot 880; \$\frac{1}{2}\cdot 840; \$\frac{1}{2}\cdot 880; \$\frac{1}{2}\cdot 840; \$\frac{1

Tertiary butyl chloride (CH.), C.Cl. (51°).
S.G. 15 '8471; 35 '8368. M.M. 6'257 at 15°.

Formation.—F. By action of chlorine on (CH.), CH (Butlerow, J. 1864, 497).—2. By heating isopartyleon (CH.). C.Cl. (1997).—2. By heating isopartyleon (CH.). C.Cl. (1997).—2. By heating isopartyleon (CH.). ing isobutylene, (CH₁), C:CH₂, with cono. HCl at 100° (Salesky, A. 165, 92; B. 5, 480; Le Bel, Bl. [2] 28, 462; Puchot, A. Ch. [5] 28, 549).— 3. From tri-methyl-carbinol and AcCl or PCl. -4. From isobutyl iodidc and ICl (Linnemann, A. 162, 18).

Properties .- With water (5 vols.) at 100° it is readily converted into tertiary butyl alcohol (Butlerow, A. 144, 33). It partakes, therefore, somewhat of the character of an acid chloride.

BUTYL-CHLORO- v. CHLORO-BUTYL BUTYL-CINCHONIC ACID v. BUTYL-QUINO-LINE-CAUBOXYLIC ACID.

BUTYL-CRESOL v. METHYL-BUTYL-PHENOL. ISO-BUTYL CYANATE Pr.CII...N.CO. (110°). From isobutyl iodide and silver cyanate (Brauner, B. 12, 1877).

Tertiary butyl oyaneto (CII,), C.N.CO. cor.). S.G. 9 8676. The chief product of the action of isobutyl iodide on silver cyanate (B.)

BUTYL CYANIDE v. VALERONITRILE.

n-BUTYLENE C,H, i.c. CH2.CH2.CH:CH2. Vinyl-ethane. Ethyl-ethylene. Mol. w. 56. (c.-1°). Occurs in the oils deposited from compressed coal-gas (Colsen, Bl. [2] 48, 52; C. R. 104, 1286).

Formation. - 1. By boiling n-butyliodide with alcoholic KOII (Grabowsky a. Saytzeff, A. 179, 325; Lieben a. Rossi, A. 158, 164).—2. From bromo-ethylene and ZnEt₂ (Chapman, C. J. 20, 28; Wurtz, Z. [2] 5, 407).—3. Together with secondary brityl alcohol by the action of HNO. on n-butyl-amine (Meyer, B. 10, 136).

Properties .- Gas. Forms with Br a dibromide (167°). HI forms CII, CH, CIII.CH,

ψ-Butylene Clf. CH:CH.CH., (β)-Butylene. s-Dimethyl-ethylene. (1°).

Occurrence.—In the oils from compressed

coal-gas (Colson, C. R. 104, 1286).

Formation .- 1. From secondary butyl iodide and alcoholic KOH, Ag.O and water, or AgOAc (De Luynes, A. 129, 200; 132, 275; Lieben, A. 150, 108; 151, 121).—2. Together with isobutylene by distilling n- or iso- butyl alcohol with ZuCl. (Nevolé, Bl. 24, 122; Le Bel a. Greone, Am. 2, 23; Bl. [2] 29, 306).—3. By heating secondary butyl alcohol .- 4. From McI, allyl iodide, and Na, small quantities of the two other butylenes being also formed (Wurtz, Bl. [2] 8, 265; Grosheintz, Bl. [2] 29, 201).—5. ILSO, acting on isobutyl alcohol forms (1 vol. of) CH3.CH:CH.CH, and (2 vols. of) (CH3)2C:CH2 (Konovaloff, Bl. [2] 34, 333; cf. Puchot, A. Ch. [5] 28, 508).-6. By boiling isobutyl iodide with PbO (Eltekoff, Bl. [2] 34, 347).

Properties. - Gas. Br gives C.H.Br. (156-

159°). HI gives CH₂.CH₂.CH₁.CH₃.

, Isobutyleno (CII3)2C:CH2. u-Di-methyl-ethylcne. (-6°). S.G. = 14 ·637 (Puchot). H.F. p. 10,660. H.F.v. 8920 (Th.). V.D. 1.93 (calo. 1.94). Occurs in the oils obtained by compressing oil-gas (Faraday, T. 1825, 440) or coal-gas (Colson, Bl. [2] 48, 52; cf. Prunier, Bl. [2] 19,

Formation.—1. By electrolysis of potassium

valerate (vilbe, A. 69, 269).—2. From sec. or tert butyl alcohol and H.SO, or ZnCl2 (Wurtz, 93, 107; Konovaloff, Bl. [2] 34, 333; Lermon' v. A. 196, 117; Puchof, A. Ch. [5] 28, 508; C. R. 85, 757; Scheschukoff, Bl. [2] 45, 181; cf. Nevolé, Bl. [2] 24, 122). 3. Together with ethylone and propyleno by passing fusel oil through a red-hot tube (Wurtz, A. 104, 249; Butlerow, A. 145, 277).—4. From iso or tertbutyl iodide and alcoholic KOH (De Luynes, C. R. 56, 1175; A. Ch. [4] 2, 385; Butlerow, A. 144, 19; Z. [2] 6, 236).—5. By heating dimethyl-acrylic acid to 210°-220 during 25-30 hours (Gorboff a. Kessler, Bl. [2] 41, 392).

Preparation. - Isobutyl iodide is run into boiling 2 cholic K'III and the gas collected.

Properties.— (tas, sl. sol. water, v. sol. alco-

hol, v. e. sol. ether and HOAo. Reactions.—1. Conc. HCl at 100° forms tertbutyl chloride.—2. Conc. HI aq absorbs it, forming tert-butyl iodide and tri-methyl carbinol (Soheschukoff, Bl. [2] 46, 823). - 3. Conc. H.SO. 3 pts.) mixed with water (1 part) absorbs it, and on diluting with water and distilling tertbutyl alcohol is got. H SO, (5 pts.) mixed with less water (1 pt.) forms much dodceylone.—
4. Br in CS₂ forms C₄H₂Br₂ (148°) and C.H.Br₃ (173°-183°) at 235 mm. (Nevolé, C. R. 83, 65; Norton a. Williams, Am. 9, 88).-5. HClO forms a ohloro-butyl alcohol (137°) (Butlerow, A. 144,
1; Z. [2] 6, 236; Henry, Bl. [2] 26, 23).-6. CrO₃ gives acctone, oxalic acid, acctic acid, &c. (Zeidler, A. 197, 251).—7. Cl forms isobutenyl chlorides C, II, Cl (Scheschukoff, J. R. 16, 488).

Combinations.—(C,II,)_Al,Cl, From cthylenc, Al,Cl, and IICl (Gustavson, J. R. 16, 97).—(O,H,),Al,Br,: oil. S.G. 22·1. V. p. 147.
BUTYLENE ALCOHOL v. Dr. 5 [-BUTANE.

BUTYLENE DIAMINE C,H,2N

i.e. $C_4H_a(NH_2)_a(?)$. (above 140°). From ethylene cyanide, tin, and HCl (Fairley, C. J. 17, 362; could not be obtained by Ladenburg, B. 16, 1150, or Lollmann a. Würthner, A. 228, 221).— B'2H2PtCl8

BUTYLENE BROMIDE v. DI-BROMO-HUTANE. BUTYLENE TRI-CARBOXYLIC ACID

CH₁,CH:C(CO₂H).CH(CO₂H)₂. Ethylidene-ethenyl-tri-carboxylic acid. [185°]. Formed by the action of a-chloro-crotonio ether upon sodiomalonic ether, and saponification of the product (Hjolt, B. 17, 2833). V. sol. water, m. sol. ether.— $A'''Ag_3$: flocculent pp.— A'''_2Ca_3 × and A'''_2Ba_3 ×: easily soluble powders.

Mono-ethyl ether A"H, Et 3aq: [70°]: large triclinic crystals, a;b;c=0111:1: 7553.

A"H,Et aq [145°].

Tri-ethyl ether Et, A". (286°). BUTYLENE GLYCOL v. DI-OXY-BUTANE.

BUTYLENE GLYCOL CHLORHYDRIN v. CHLORO-BUTYL ALCOHOL.

BUTYLENE "YANAMINE C.H., N. [173°] Formed by heating guanidine isovalcrate at 225° Trimetrio needles, m. sol. cold water. Converted by oone. H.SO₄ into butylene guananaido C.H.₁N₂O₂, whence HNO₂ forms oyanurio acid (Bandrowski, B. 9, 240).—B'HC!—B'AgNO₂.—

B .H.SO. BUTYLENE HYDRATE v. Sec-BUTYL ALCOHOL. BUTYLENE NITRITE (?) C₄H₂N₂O₄. From isobutylene and cono. HNO₂ (Haitinger, M. 2, 287).

Entylene nitrite (?) [96°]. Formed by boiling petroleum of Tiffis ith HNO. (Beilstein a. Kurbatoff, B. 14, 1621)

ISV-BUTYLENE ON THE CIH, O i.e. O CMe, (52°). S.G. 2 ·8311. From isobutylene with successive 1. ment with IClO and KOH (Eltekoff, 31.27 L., 23; J. R. 1, 368). Water, at ordinary temperatures, unites with it, forming di-axy-

s-Butylene oxide $O<_{\rm CHMe}^{\rm CHMe}>$. (57°). S.G. 2 *8314. Formed sin ilarly from CII₃.CII:CH.CII₄ (E.). Water a. 100° forms di-oxy-lettano CΠ_z.CH(OH).CH(zH).CH_z.

BUTYL ENNYL KETONE CullesO (?) or C₁H₉,CO.C₁H₈(C₅H₁₁). Amyl valerone. (209°). S.G. ¹² ·845. From CO on sodium amylate, NaOC₂II₁₁, at 160° (Geuther a. Fröhlich, d. 202, 301). Liquid, does not combine with NaIISO₃. BUTYL ETHER v. BUTYL OXIDE.

BUTYL-FORMIC ACID v. VALERIC ACID. BUTYL-GLYCERIC ACID v. Di-oxy-nurving

BUTYL-GLYCIDIC ACID v. Dr-oxy-Butyrio ACID

BUTYL GLYCOL v. DI-OXY-BUTANE

BUTYL-GLYOXALINE C,113(C,H,)N2. Gly-[121°]. (274°). Flat needles. oxal-amyline. Sl. sol. water. Prepared by the action of valerio aldehyde-ammonia on glyoxal (Radziszewski, B. and cnyde-anmonia on glyoxal (flatiziszewski, B. 16, 747; 17, 1291). Br forms C, II, Br, N₂ [158°] and C, H, Br, N₂ [217°].

Salts.—B'HCl [136°].—B'HBr [100°].—
B'H, C, O, [196°].—B', H, PtCl,

ISO-BUTYL DIGJANIDE C, II, N₃ i.e.

C, H, N₃ (C, H₃). The sulphate of the copper

derivative Cu(C,H,N,),H,SO, is formed by tho action of aqueous CuSO, and isobutylamine on dicyandiamido at 100° (Smolka, M. 4, 815). Alkaline syrup, acsorbing CO2 from the air.

Salts. — B' H₂SO₄ I laq. S. 26·3 at 16°. Colourless transparent leaflets.—B'H₂SO₄ I laq. —B'HGi. [216°]. S. 40 at 16°. —B'2HGl. [191*]. Deliquescent.—B',H,P'lCl, aq: golden four-sided tables.—B',H,CrO, aq.—l',H,C,O, Metallic derivatives Cu(C,H,N,);:

silky red needles, sl. sol. water, sol. dilute acids. -Cu(C,H,N,), 2HNO, - Cu(C,H,N,),H,Cl, aq. -Cu(C₆H₁₄N₃)₂H₂SO₄: commine red grains. S. \cdot 26 at 18°. -Cu(C₆H₁₄N₃)₂H₂SO₄aq: palo red grains. -Cu(C₆H₁₄N₃)₂H₂SO₄3aq: light rose red crystals. n.BUTLL HEPTYL-OXIDE C.H. O.C.H.

(205·7°). S.G. § ·8023. S.V. 271·3. C.E. (0°-10°) 00097 (Dobriner, A. 243, 8)

BUTYL-HYDRO-ANTPRANOL v. BUTYL-ANTHRANOL DINYDRIDE,

ISO-BUTYL HY"OPHOSPHATE (C,H,),P,O,. S.G. 15 1.125. From isobutyl iodide and Ag, P,O, at 140° (Sänger, A. 232, 12). Oil. Decomposed by hot water.

Iso-butyl-hypophosphate of barium

(C,H,)BaHP,O, 5aq. Needles. BUTYLIDENE-ACETO-ACETIC ETHER .

ISO-BUTYLIDENE-AMIDO-BENZOIC ACID C₁₁H₁₅NO₂ i.e. Pr.CH:N.C₆H₄CO₂H. [145°-150°]. From amido-benzoio acid and iso-butyrio aldehydo (Sohiff, A. 210, 114). Slender needles, decomposed by cold dilute alkalis.

TRI-ISO-BUTYLIDENE DI-AMINE v. Hydro-

butyramide under Isosutkric Aldenyde. 180-BUTYLIDENE CELORIDE v. Di-chloro-ISOBUTANE.

ISO-BUTYLIDENE-ETHYLENE-ANILINE v. DI - PHENYL - ISOPROPYL . METAPYRAZOL *TETRA-HYDRIDE.

**BUTYL IODIDES CH₃.CH₂.CH₂.CH₃.L Mol. w. 184. (131° ctr.). S.G. $\frac{2}{3}$ 1·647 (Brühl); $\frac{2}{5}$ 1·6476. S.V. 188-2. C.E. (0°-10') 00098 (Dobrinor, A. 243, 20). μ_{β} 1·510. 1254.47. From n-butyl alcohol and III (Linnemann, A. 161, 196). ICl, t 250° acts on it thus re C₄H₉I + 11ICl₂ = 2C₄H₉I + 9HCl + 12ICl (Krafft, B. 10, 805).

Socondary w-butyl iodide CH₃CH₂CH₂CH₃CH₄CH₄(118°). S.G. ⁵₀ 1·626; ²⁰₀ 1·592; ³⁰₀ 1·579 (Lieben); ⁹ 1·632; ²⁰ 1·600 (Luynes). V.D. 6·6 at 20° (obs.). Formation.—1. By distilling erythrite with III (De Luynes, A. 125, 252; BL. 3) .- 2. From ethyl-chloro-ethylic ether CH,Cl.CEtH.O.Et and HI at 140° (Lieben, A. 150, 87). — 3. From n-butylene and HI

(Wurtz, A. 152, 23).

(Witt2, A. 103, 20).

Isobutyl iodide (CH₃), CH.CH.I. (120°).

S.G. Ψ 1-606 (Brühl); ½3 1-6130, ½5 1-6007 (Perkin, C. J. 45, 462). μ_B 1-506. R_∞ 51-41. S.V.

128-28 (Schiff, B. 19, 564). C.E. (0°-20°) 0110 (Brauner, A. 192, 69). M.M. 12 199 at 19 1°. From isobutyl alcohol and HI. Distils constantly with 21 pts. water at 96° (Pierre a. Puchot, C. R. 74, 224). Treated with AgNCS gives a product, (CII,) CH.CH2NCS, which on saponification yields isobutylamine and, in greater quantity, tertiary butylamine (B.). Heated with lead oxide it yields pseudo-butylehe besides iso-butyleno (Eltekoff, Bl. [2] 34, 347).
Tertiary bntyl iodide Me₃CI. (100°). S.G. 2

1.571. Formation.—From tertiary butyl alcohol

and HI.

Preparation.-By passing isobutylene (q. v.) into funning III, cooled with iccound well shaken

(Markownikoff, Z. [2] 6, 29). Reactions.—1. If the iodide (20 g.) be shaken with water (30 g.) for two days it dissolves, being eonverted into the alcohol: Me₃CI+1I₂O= Me_sC.OH + HI (Dobbin, C. J. 37, 237).--2. With dry ZnO it forms tri-isobutylene -- 3. With sodium it forms isobutylene, tri-isobutylene, and hydrogen, with small quantities of a hydrocarbon not absorbable by H₂SO₄ (Dobbin).—4. Moist Ag₂O, Zn and water, and AgOAc, form isobutylene (Butlerow, Z. [2], 6, 237). 5. With McOII at 100°-110° it gives McI and trimethyl carbinol (Baucr, A. 220, 163).-6. With MeOAo at 110° it forms isobutylene, HOAe and MeI.

DI-BUTYL-KETINE v. DI-METHYL-DI-BUTYL-

PYRAZINE.

DI-ISOBUTYL KETONE C,II,80 i.e. C.H., CO.C.H., Valeronc. Mol. w. 142. (182°). S.G. 20 833. Formed, in small quantity, by distilling calcium valerate (6 pts.) with CaO (1 pt.) (Löwig, P. 42, \12; Ebersbach, A. 106, 268; Schmidt, B. 5, 600). Does not combine with NaHSO,

DI-ISOBUTYL-KETONE-SULPHONIC ACID C₄H₈(HSO₃)—CO—C₄H₈(HSO₃). The sodium salt is formed by leaving phorone in contact with a saturated solution of NaHSO, for 2 or 3 weeks. It forms colourless prisms (A'Na, 2 aq): soluble in water and alcohol (Pinner, B. 15, 593). BUTYLLACTIC ACID v. OXY-BUTYRIC ACID. BUTYL-LUTIDINE v. DI-METHYL-BUTYL-PYBL

n-BUTYL-MALDNIC ACID C,H,2O, i.e. C₁H_n.CII(CO₂H)₂ [101°]. Prepared by saponification of the nitrilo C₁II_n.CH(CN).CO₂Et obtained by the action of KCN on chlorohexoic-ether. Thick prismatic crystals. V. sol. water, alcohol, and ether. It gives a reddishviolet colouration on warming with H.SO. At about 150° CO. is evolved and it is converted into caproic acid.

Salts. -4."Ba: white plates. S. 2.98 at 24°. A"Pb: very sparingly soluble white glistening Plates. S. 011 at 20°. A"Guaq: blne glistening plates. S. 086 at 22°. A"Ag., hite pp. S. 119 at 23° (Hell a. Lumpp, B. 17, 2217).

Isobutyl-malonic acid PrCH, CH(CO, H).

[107°]. From its ether (Hjelt, J. 1882, 875). Ethyl ether Et.A". (225°). S.G. 17.983. From sodium malonic other and isobutyl iodido.

n-BUTYL MERCAPTAN C.H.SH. Mol. w. 90. (98°). S.G. $\frac{9}{6}$ ·858; $\frac{16}{9}$ ·843 (Saytzeff a. Grabowsky, A. 171, 25T; 175, 35I). HNO₃ gives butane sulphonie acid.

Isobuty1 mercaptan (88°). S.G. 12 ·848 (Humann, A. 95, 256); 🙄 ·8357 (Nasini, G. 13, 301). V.D. 3·10 (obs.). B_∞ 27·47 (N.). Formed by distilling K(C,H_o)SO, with aqueous KHS.—KSC₁H_o. — Hg(C,H_oS); pearly scales.—Pb(C,H_oS)_o; yellow crystalline pp.

Sceondary butyl mercaptan (85°). S.G. 17 830. From sec-butyl iodide and KHS.-C.H.SAq. - (C.H.S).Hg [189°] (Reymann, B. 7, 1287).

TETRA - ISOBUTYL - METHYLENE - DI-AMINE C₁₇H₃₈N₂ i.e. N(C₁H₃)₃.CH₂N(C₄H₉)₃. (215°-255°), 4 From 'trioxymethylene' (formio paraldehyde) and di-isobutylamine (Ehrenberg, J. pr. [2] 36, 117).—B"H_PtCl_s [1982]. Forms with CS₂ a compound C₁₇H₃₈N₂CS₂ [54°].

BUTYL - MUSTARD OIL v. BUTYL TRIO-CARBEAUDE.

ISO-BUTYL-NAPHTHALENE CHILL, i.e. $C_{16}\Pi_1(C_4\Pi_4)$. (280°). Prepared together with $(\alpha-\alpha)$ and $(\alpha-\beta)$ -dinaphthyl by heating naphthalene and isobutyl chloride in presence of aluminium chloride. Colourless oil, very slightly volatile in steam. Pieric acid compound; [96°] aggregates of golden needles (Wegscheider, M. 5, 236).

ISO-BUTYL NITRATE C.II., NO. S.G. 2 1038. From AgNO, nrea, and isobutyl igdide (Wurtz, A. 93 120; Chapman a. Smith, Z. 1869, 433).

ISO-BUTYL NITRITE (CII,) CH.CII O.NO. (67°). S.G. 9 891. H.F.p. 47,800. H.F.v. 44,900 (Th.) (Chapman a. Smith, %. 1869, 433; Pribram a. Handl, M. 2, 658; Bertoni a. Truff, G. 14, 23).

Tertiary butyl nitrite (Crt.), C.ONO. (63°) (B.); (76°-78°) (R). S.G. 2 ·8914 (B.). Formed together with a little of the isomeric nitrobutane (110°-130') by distilling tertiary butyl iodide with silver nitrite (Tscherniak, A. 180, 155; B. 7, 962). Prepared by distilling tertiary butyl alcohol (1 mol.) with glyceryl tri-nitrite (1 mol.) (Bertoni, G. 15, 357)

BUTYL-PSEUDO-NITROLE v. NITROSO-NITRO-

BUJANE.

BUTYL-NITROLIC ACID v. NITROSO-NITRO-

n-BUTYL-OCTYL-OXIDE C,H, O.C,11, (225.7°). S.G. 6 8069. S.V. 295.7. C.E. (0°-10°) ·00097 (Dobrinor, A. 243, 9).

ISO-BUTYL OXALATE v. OXALLO ACID. ISO-BUTYL-OXAMIC ACID

PrCH2NH.CO.CO2H. From oxalie ether (1 mol.) and dry isobutylamine (1 mol.) at 160° (Malbot, C. R. 104, 229). CaA'2.

Di-isebutyl-examic acid

(PrCH2)2N.CO.CO2H. Similarly prepared from di-iso-butylamine (M.).

DI-ISO-BUTYL-OXAMIDE (PrCH_NII)_C_O_. [1678]. From oxalic ether (1 mol.) and isobutylamin (2 mols.). Acute plates, insol. boiling water; may be sublimed (Malbot, C. R. 104, 228).

DI-BUTYL OXIDE $(C_1H_0)_2O$. Mol. w. 130. (141°). S.G. 2 784 (Lieben a. Rossi, A. 165, 110); n · 7865. S.V. 197-3. C.E. (0°-10°) · 00109 (Dobriner, A. 243, 8).

Di-isobutyl oxide (100°-104°). From isobutyl iodide and KOC, H, or Ag,O (Wurtz, A.

Di-see-butyl oxide (121°). S.G. 21 .756. From aldehyde hydrochloride and ZnEt₂ (Kessel,

A. 175, 56; B. 7, 291).
p.ISO - BUTYL - PHENOL C.H., C. II, OH.
[99°]. (231°) (S.); (237°) (L.).
1. From amidoiso-butyl-benzene by the diazo- reaction (Studer, A. 211, 242; B. 14, 1474, 2187).—2. From phenol (100 g.), isobutyl alcohol (80 g.) and ZuCl₂ (240 g.) (Liebmann, B. 14, 1842; 15, 150, 1991; Dobrzycki, J. pr. [2] 36, 390). Needles (from alcohol). Volatile with steam. V. sol. Gives a pp. with bronnine-water, but alkalis. no colour with Fe₂Cl₆. Is antiseptic. PCl₃ gives chloro-iso-batyl-benzene, waich on oxidation gives p-chloro-benzoic acid. Gives a dinitro-derivative [93°]. Fused with P₂O₃ it gives } phenol and is butylene. Anmonia and ZnCl, give C₄H₅,C₆H₄NH₂ (Lloyd, B. 20, 1254). C₄H₉.C₆H₄OMe. (215.5°). Methyl ether

S.G. 27 .937. Ethyt ether C,H,C,H,OEt. (235°) (S.);

(242°) (L.).

C, H, C, H, OBz. Benzoyl derivative[83°] (S.); [80°] (Kreysler, B. 18, 1717); (335°); (344°) (K.). White plates (from alcohol).

**Acetyl derivative C₁H₂,C₆H₄OAc. (245°). S.G. 24 .999.

ISO-BUTYL-PHENOL SULPHONIC ACID C₄H₉·C₆H₃(OH).SO₃H. From isobutyl phenol and H₂SO₄ (Liebmann, B. 15, 1990).—BaA', 2aq. 1SO-BUTYL-PHENYL-AMINE v. Ambo-

PHENYL-BUTANE.

Di-isobutyl-di-phenyl-amine Thick oil. (C,H,C,H,)2NH. (290°-315°). Formed together with $C_6 H_1(C_4 H_9)$.NH, by heating oxy-phenyl-isobutane $C_6 H_4(C_4 H_9)$.OH [1:4] with ammoniacel Zzzce (or ZnCl) and NII Br (or NH,Cl) at 320°-330°; the yield is 20 25 p.c.—

B',II,Cl,PtCl,: yellow needles.
Acetyl derivative (C₆H₄,C₄H₉),NAc: [75°]; glistening white plates; sl. sol. water, v. sol. aloohol and benzene (Licyd, B. 20, 1255). DI-ISO-BUTYL-DI-PHENYL CYANAMIDE

C21H26N2 i.e. C(NC6H4.CH2Pr)2. Carbo-di-phenisobutyl-imide. [189°]. Formed by boiling a solution of di-isobutyl-di-phenyl-thio-urea in benzene with PhO (Pahl, B. 17, 1242). Colour- insol. water.

less crystals. Sol. ho benzone, sl. sol, ether. By warm dilute alcohol it is converted into diisobutyl-di-phenyl-ures. With amines it combines to form guanidines. Heated with CS2 at 170' it rields isobutyl-phenyl-thio-carbimide.

ISO-BUTYL-PHENYLENE-DIAMINE C₆H₄(NH₂)(NHCH₂Pr'. [39°]. Formed by the reduction of p-nitros e-iso butylanilino (Wacker, A. 243, 299). Plates (from ether); can be distilled. Its chloride forms white plates; v. sol. water, sl. sol. alcohol, insol. ether.

DI-p ISO-BUTYL-DI-PHENYL-GUANIDINE 1IN:C(NH.C_aH_aC_cH_a)_x. Di-phenisobutyl-guarid-ine. [173°]. If med by heating an alcoholic solution of di-p-isobutyl di-phenyl-thiourca with NII, and lead oxide (Pahi, B. 17, 1240). Colourless plates. V. sol. alcohol and benzene.— B'_H_Cl_PtCl_4: yellow crystalline pp.

Tri-p-is. butyl-tri-phenyl-guanidine

C₁H_p,C₁H₁N:C(NH,C₁H₁,C₁H_p)₂. Tri-pheniso-butyl-quantitine. [164°]. Obtained by digesting an atcoholic solution of di-p-isobutyl-di-phenylthiourea with p-isobutylphenyl-mine and lead oxide (P.). Small white needles. V. sol. benzene and hot alcohol. B'allaClaPtCla: yellow needlos.

ISO-BUTYL-PHENYL-(B)-NAPHTHYL-THIO-UREA $C_{10}\Pi_1$, NH.CS. NH. $C_{6}\Pi_1$, $C_{1}\Pi_9$, Phenisobutyl- (β) -naphthyl-thiourea. [152], Prepared by warming an alcoholic solution of (β) -naphthylthio-carbinide and isobutylphenyl-amine (Mainzer, B. 16, 2026). White plates. Sol. boiling alcohol. By phosphoric acid it is split up into isobutylphenyl-thio carbinide, (B)-naphthylthio-carbinide, isobutylphenyl-amine, and (B)naphthylamine

1SO-BUTYLPHENYL-PHENYL-THIOUREA v. Phenyl-isobutylphenyl thiourea.

TR1-ISO-BUTYLPHENYL-PHOSPHATE

PO(OC, HrC, II,),. (above 400'). Obtained by heating isobe'y! phenol with POCl₃; the yield is 90 p.c. of the theoretical (Kreysler, B. 18, 1700). Thick liquid. V. s !. ether, benzene, and warm alcolai, sl. sol. petroleum ether.

TE' RA-ISO-BUTYLPHENYL SILICATE Si(OC, II, C, II,),. (e. 380°). Obtained by heating isobutyl-phenol with SiCl₄; the yield is 70 p.c.

of the theoretical (Hertkorn, B. 18, 1692). Thick colourless oil. V. sol. benzene, chloroform, CS., etc. p-ISO-BUTYL-PHENYL-THIOCARBIMIDE

SC:N.C.II.(C.H.). Phenisobutyl-mustard-oil. [42°]. (277°). Formed by leading di-p-isobutyldi-phenyl-th ourea with phosphorio acid (Pahl, B. 17, 1235). Long white needles. V. sol. alcohol. and ether.

DI-j)-ISO-BUTYL-7/I-PHENYL-THIOUREA SC(NH.C, II, C, II,)2. Di-phenisobutyl-thiourea. [193°]. Formed li cohobiting an alcoholic solution of p-isobutylphenyl-amine with CS₂ (Pahl, B. 17, 1235). Fine white needles. Sol. ether and benzene, sl. sol. alcohol, insol. water.

DI-p-ISO-BUTYL-DI-PHENYL-UREA $OC(N11.C_611..C_1H_p)_2$ Di - phenisobutyl - urea. [281°].

Formation.-1. By the action of carbonyl chloride on p-isobutylphenyl-amine dissolved in benzene .- 2. By cohobating an alcobolio solution of the thiourea with mercuric oxide (Pahl, B. 17, 1240). Colourless needles. Sol. hot alcohol.

ISO BUTYL PHOSPÄINE O.H.PH. (62°). Prepared, together with the following, from isobutyl iodide, ZnO, and PH,I at 100° (Hofmann, B. 6, 296). Resembles ethyl-phosphine in preparation and properties.

Di-isobutyl phosphine (C.H.).PH. (1.3°).

Tri-Isobutyl phosphine (C,H,),P. (215°). From the preceding and isobutyl iodide.
Tetra-isobutyl-phosphonium iodide

(C.H.) PI. Crystalline. ISO-BUTYL-PHOSPHINIC ACID C.H.PO.H. [100°]. Parafiin-like solid (Hofmann, B. 6, 301). –A'Ag. Amorphous pp.

Dr isobutyl phosphinic spid (C,H,),PO.H.

Oil .- A'Ag. Amorphous.

ISOBUTYL PHOSPHITE. Dichloride. (155°). S.G. º 1·191. A by-C.H.O.PCl2. product in the conversion of isobutyl alcohol into isobutyl chlorido by PCl, (Mencehutkin, A. 139, 347)

DI-ISO-BUTYL-PINACONE C₁₁ll₁₀O₂ i.e. Pr.CH₂.CIL₂.CMc(OH).CMc(OH).CIL₂.CH₂.Pr. Dioxy-tetra-decane. Tetra-decylene glycol. [30°]. (268°). A product of the action of Na upon methyl iso-amyl ketono (Rohn, A. 190, 305: Purdie, C. J. 39, 468). Needles, insol. water.

BUTYL-METAPYRAZOLONE v. DI-OXY-

BUTYL-METAPYRAZOL.

(Py. 3)-ISO-BUTYL - QUINOLINE $C_{13}H_{13}N$

CH:CH i.e. C.H. (271°). Colourless oily $N = \dot{C}(C_1H_9)$ liquid. Formed by distilling its (Py. 1)-carboxylio acid with limo (Doebner, B. 20, 280; A.

242, 282). Piorio acid compound B'.C.H.N.O. [161°]. Lomon-yellow plates (from alcohol). -

(B'HCl) PtCl. Orange-red needles (from water). (Py. 3)-ISO-BUTYL-QUINOLINE—(Py. 1)— CARBOXYLIC ACID

C(CO₂II):CH C(C,II,) O, H, NO, i.e. C, H, . Isobutyl-

cinchonic acid. [186°]. Formed by the action of isovalerio aldehydo upon pyruvic acid and aniline. White satiny plates (containing 1 haq) (Doebner, B. 20, 279; A. 212, 280).—B'HClaq: plates. - B'2H2PtCl6. - AgA'.

ISO-BUTYL SILICATE (PrCII,), SiO,. (c. 258°). S.G. 15 953. From SiCl, and isobutyl alcohol (Cahours, C. R. 77, 1408). Slowly decomposed

by water.

BUTYL SULPHATES

n-Butyl-sulphurio aoid Pr.CH., SO, H.

Salt.—BaA'2aq: erystalline plates. S.G. 21 1.778 (Lieben a. Rossi, A. 165, 116; Clarke, B. **11, 1**506).

Isobutyl sulphurio acid PnCII, SO.H. From the alcohol and H2SO, (Wurtz, C. R. 35, 310).-

KA': laminæ (from alcohol).

Chloride C.H. O.SO, Cl. From isobutyl

n-BUTYL SULPHIDE (C.H.),S. Mol. w. 146. (182°). S.G. § 852; § 839 (Saytzoff, A. 171, 253). From butyl iodide and K.S.

Isobutyl sulphide (Pr.CH₂),S. (171° i. V.). B.G. 12 836. Formed by distilling isobutyl sulphate with conc. aqueous KHS (Beckmann, J. pr. [2] 17, 445). Secondary butyl sulphide (CMeEtH) 3 (165°). S.G. 23 832. Combines with McI (Rey-(OMeEtH).3 mann, B. 7, 1288).

Isobutyl disulphide (CH2Pr)2S2. (220°)(Spring a. Legros, B. 15, 1938). ISO-BUTYL SULPHOCYANIDE CH₂Pr.S.CN.

(175°) (Reimer, B 3, 757).

n-BUTYL SULPHONE (C,H,)2502. [449]. From fuming HNO, and (C,H,),S (Grabowsky, A. 175, 348).

Di-isobutyl sulphone (C₄II₃)₂SO₂, [17°]. (265° i. V.). S.G. ½ 1·006. Di-isobutyl sulphoxide (100 pts.) is dissolved in water (1000 pts.) and oxidised by KMnO₄ (63 pts.) dissolved in water (1900 pts.). Excess of permanganate destroyed by sodic formiate and the sulphone oxtracted by other. The yield is quantitative (Beekmann, J. pr. (2 | 17, 448).

Properties. White plates. Not affected by

reducing agents

n-BUTYL SULPHOXIDE (C,II,),SO. [32°]. From IINO, (S.G. 1.3) and (C, H,)2S (Grabowsky, A. 175, 348).

Di-isobutyl sulphoxide (C,H,),SO. [69°]. From di-isobutyl sulphide (1 pt.) and (2 pts. of) HNO₃ (S.G. 1.4) in the cold. Extracted with ether after neutralisation. The yield is quantitative (Beckmann, J. pr. [2] 17, 116).

Properties.—Yellow needles. Soluble in 2 parts of cold water, but separates as an oil on warming. Readily reduced to di-iso-butyl sulphide.

BUTYL SULPHURIC ACID v. BUTYL SUL-PHATE.

DI-ISO-BUTYL-THETINE

CO2H.CH2.S(C,H9)2(OH). The hydrobromide is a syrup formed by the action of isobutyl sulphide on bromo-aectic acid at 100°. Lead converts it into crystalline C₁₀H₂₀SO₂5PbBr₂ and C₁₀H₂₀SO₂3PbBr₂ (Letts, Pr. E. 28, 588).

n BUTYL THIO CARBIMIDE CILPIN.CS. Mol. w. 115. (167°). Formed by boiling n-butylamine with CS₂ and alcohol. Evaporating the alcohol and heating the residue with aqueous HgCl₂ (Hofmann, B. 7, 508). NH₂ gives butylthio urea [79°].

Isobutyl thio-carbimide CH, Pr. N.CS. (162°). S.G. 14 937. Similarly prepared. (II.). NH, gives isobutyl-thio-urea [94°].

Secondary isobutyl thio-carbimide

ClimeEt.N.CS. (160°). S.G. 12 944. Similarly prepared. It is the essential constituent of the oil of scurvy-grass or spoon-wort (from Cochlearia officinalis) (Hofmann, Z. [2] 5, 400; B. 2, 102). NH₃ gives butyl-thio-uroa [135°

Tertiary butyl thio carbimide CMe. N.CS. [11°]. (140°). S.G. 34 '900 (Rudneff, J. R. 11, 179; B. 12, 1023).

BUTYL THIO-CARBONIC ACID v. Thio-

CARBONIC ACID.

n-BUTYL-THIOPHENE G.C. (C.II.) (181°). S.G. 12 957. Colourless oil. Formed by the sloohol and SO₂Cl₂ (Belliend, J. pr. [2] 15, 34). section of sodium on a mixture of iodothiophene and butyl bromide (Meyer a. Kreis, B. 17, 1561).

BUTYL THIO-UREAS. The preparation and properties of these bodies are analogous to those of the corresponding ethyl thio-ureas (q. v.).

n-Butyl thio-urea CH_Pr.NH.CS.NH... [79°]. From butyl thio-oarbimide and NH. (Hofmann, B. 7, 512).

lsobutyl thio-urea [94°] (H.).

Sec-butyl thio-urea [138°] (H.).
Tert - butyl thio-urea CMs, NH.CS.NH..
[185°] (Budnsff, Bl. [2] 33, 300). Prisms, sol.

Di-tert-butyl thio-ursa SC(NHCMe₃). [162°]. From tert-butyl-amine, CS₂, and alcohol. Formed also by the action of tert-butyl thiocarbimide on tert-butyl-amine (Rudneff, Bl. [2] 33, 300).

m-ISO-BUTYL-TOLUENE CaH (CIIa)(C,Ha) [1:3]. (187°) (K. a. B.); (194°) (R.); (185°) (E.). Methyl - isobutyl - benzene. Colourless liquid. Occurs in oil of resin (Kelbe ne Baur, B. 16, 619, 2559; Renard, A. Ch. [6] 1, 250). Prepared by the action of isobutyl bromide on toldene in preserve of Al.Br. (K. a. B.). Formed by diazotising methyl-isobutyl-phenyl-amine and treating the diazo- salt with an excess of SnCl2 (Effront, B. 17, 2329). On oxidation with HNO, it first gives m-iso-butyl-benzoic acid and finally isophthalic acid (Kelbo a. Pfeiffer, B. 19, 1723).

p-(Iso?) - Butyl - toluene $C_6H_1(CH_3)(C_1\Pi_9)$ [1:4]. (178°). Occurs in oil of resin. oxidises it to p-toluic acid (Kelbo a. Baur, B.

16, 2562).

p-Isobntyl-toluene (c. 195°)? Formed by heating toluene with isobutyl alcohol and ZnCl₂ (Geldschmidt, B. 15, 1067). Formed, together with the m-isomeride, by the action of isobutyl bromide on toluene in presence of Al₂Br_e (Kelbe a. Pfeiffer, B. 19, 1725). IINO₃ oxidises it to p-isobutyl-benzoic acid.
m-ISO-BUTYL-TOLUENE-SULPHONIC ACID

C₆H₃(CH₃)(C₄H₅)(SO₃H) [1:3:x]. [76°]. Small hygroscopic plates. Formed by sulphonation of

m-iso-butyl-toluene.

Salts.-KA'aq: large soluble pearly plates. -NaA'aq: needles. - CuA', 4aq: large blue soluble plates. -BaA', aq: small plates, sl. sol. cold water and alcohol. -PbA', 3aq: pearly plates.

Amide: ['.5°], small plates, soluble in water (Kelbe a. Baur, B. 16, 2560).

p-(Iso?)-Butyl-tolusne-sulphonic acid $C_8H_3(CH_3)(C_4H_9)(SO_3H)$ [1:4:x]. Obtained by

Salts.— Kh'1½q; small easily soluble plates.—Nah'2aq: Bah'2aq: small sparingly soluble plates.—Cuh'2xq: easily soluble blue orystals. - PbA'23aq: small plates, sol. hot wator.

Amide: [113°], largo pearly plates, sl. sol. hot water (Kelbe a. Baur, B. 16, 2563).

ISO-BUTYL-o-TOLUIC ACID

 $C_4H_5(CH_5)(C_4H_9)CO_2H$ [1:3:6]. [140°]. Formed by saponification of its nitrilo (Effront, B. 17, White needles. V. sol. alcohol and ether, sl. sol. hot water, insol. cold water .-A'Ag: colourless plates, v. sol. hot water.

Nitrile C.H. (CH2)(C,H2)CN [1:3:6]. [60°]. (248°). Formation .- 1. By distilling the formyl derivative of metnyl-isobutyl phenyl amine with zino-dust .- 2. By heating isobutyl-tolyl-thiocarbimide with copper powder (E.). Long white needles. V. sol. alcohol and ether, sl. sol. hot petroleum-ether, insol. water.

Iso-butyl-toluic acid CaH(CH3)(C,H9)CO.H [1:5:6]. [132°]. Formed by saponification of its nitrile (Effront, B. 17, 2343). White silvery platss. V. sol. alcohol and ether, sl. sol. hot platss. water. -A'Ag: colouriess plates, sol. hot water.

Nitrile C.H. (CH.) (C.H.) (C.H.) (C.H.) (C.H.) (248°).

Formation.—1. By distilling the formyl derivative of methyl isobutyl-phenyl-amins with zinc-dust .- 2. By heating isobutyl-tolyl-thiocarbimide with copper-powder (Effront, B. 17, 2343). Colourless oil, solidifies on freezing to a white crystalline mass. V. col. alcohol and ether.

ISOBUTYL-TOLUIDINE v. METHYLFISORUTYL-

PHENYL-AMINE

BUTYL-TOLYL-AMINE r. METHYL-BUTYL-PHENYL-AMINE

ISO-BUTYL-TOLYL-THIOCARBIMIDE

C_aH_a(CH_a)(C₁H_a).NCS [1:3:6], [46°], (275¾ 260°). Formed by the ection of pho-phoric acid on diisobutyl-di-tolyl-thiourca (Effront, B. 17, 2336). Long white needles. V. sol. alcohol and ethor. Iso-butyl-tolyl-thio-carbimide

Call₃(CH₃)(C₄H₃)NCS [1:5:6]. [44°]. (267°). Formed by heating di-isobutyl-di-tolyl-thioures with phosphonio acid (Effrout, B. 17, 2345).
White crystalline solid. V. sol. alcohol and White crystalline solid.

DI-ISOBUTYL-DI-TOLYL-THIOUREA

amine with CS. in alcoholic solution (Effront, B. 17, 2335). Long thin silky needles. V. sol. ether, sl. sol. alcohol.

Di-isobutyl-di-tolyl-thiourea

SC(NH.C_aH₃(CH₃)(C₄H₃)[6:1:5])₂. [175°]. White needles. Sol. hot alcohol. Formed by digesting methyl-isobutyl-phonyl-amine with an

alcoholic solution of CS₂ (Effront, B. 17, 2344). ISO-BUTYL-UREA. Valeryl derivative ISO-BUTYL-UREA. Valeryl derivative NH(C₁N_u).CO.NH(CO.C₄H_u). [102]. Flat needlos. Sol. alcohol and ether, sl. sol. water. Formed by the action of KOII on a mixture of (2 mols. of) valeramide (isopropyl-acetamide) and (1 mol. of) bromine (Hofmann, B. 15, 758).

Iso-lert-di-butyl-urea CMe, NII.CO.NH.CIL,Pr. [163°]. From tcrtbutyl cyanate and sobutylamine (Brauner, B. 12, 13, 5).

Di-Mert-butyl-urea (CMe, NII), CO. From tert-butyl cyanate and tert-butylamino (B.).

ISO-BUTYL-XANTHAMIDE v. THIO-CARBONIC ETHERS.

BUTYRAL v. BUTYRIC ALDEHYDE.

BUTYRAMIDE v. Amide of Butyrio Acid. [1740] Di-isobutyramide (C,11,.CO),NII. Formed by the action of ammonia on isobutyryl chloride LiIofmann, B. 15, 981). Long needles. Sublimable. Sol. alcohol. On distillation it splits up into isobutyric acid and isobutyro-

n-BUTYRIC ACID C.H.O. i.c.

R_∞ 35-50. S.V. 107-85 (R. Schiff, A. 220, 105); 108-3 (L). S.H. 440 26 0° (Schiff, A. 234, 300).

Occurrence.—1. Butter contains 2 p.c. of glyceryl butyrate (Chevreul, A. Ch. [2] 23, 23). Rancid butter contains free n-butyric acid (Grunzweig, A. 162, 193). -2. In crude wood vinegar.-3. In perspiration (Schotten, J. 1852, 701).—4. In muscular juice (Scherer, A. 69, 196).—5. In the contents of the large intestine.—6. As hexyl butyrate in oil got from fruit of Heracleum giganteum.-7. The fruit of Pastinaca sativa contains octyl butyrate. -8. In oheese (e.g. of

Limburg) (Iljenko, A. 63, 268).

A great many vegetable and animal juices and oils contain butyric acid, but in most cases it has not been determined whether the acid is n- or iso- butyric acid (Gorup-Besanez, A. 69, 369; Klinger, A. 106, 18; Krämen Ar. Ph. [2] 54, 9; Wagner, J. pr. 46, 155; Scherer, A. 69, 196; Rebling, Ar. Ph. [2] 92, 83; 93, 300).

Formation .- 1. A product of the fermentation of sugar, starch, milk, fibrin, and most kinds of vegetable and animal matter (Pelonze a. Gelis, A. Ch. [3] 10, 436; Lerch, A. 49, 216; Bonaparte, C. R. 21, 1070; Erdmann a. Marthand, J. pr. 29, 468; Wurtz, A. 52, 291; Scharling, A. 49, 313; Schubert, J. pr. 33, 256; Sullivan, J. 1858, 280; Ritthausen, \hat{Z} . [2] 4, 314). -- 2. By fermentation of calcium lactate: $2C_1H_2O_2 = C_4H_2O_2 + 2CO_2 + 2H_2$ 3. Along with n-butyl alcohol by the fermentation of glycerin through a Schizomycetes in presence of CaCO₃ (Fitz, B. 9, 1348).—4. By the action of CrO₃ on albuminoids (Guckelberger, A. 64, 68).— 5. By the action of HNO₃ on fats (Redtenbacher, A. 59, 49) and on Chinese wax (Buckton, C. J. 10, 166).—6. By oxidation of conine (Blyth, A. 70, 89).—7. By the aceto-acetic ether synthesis (Franklanda, Duppa, A. 138, 218) v. Aceto-Acetic ETHER. -8. By passing CO over a mixture of NaOEt and NaOAcat 200°: $C_1H_1NaO + C_2H_2NaO_2 + CO = CHNaO_2 + C_1H_1NaO_2$ (Fröhlich, A. 202, 306).

Preparation. - Sugar (6 kilos.), water (26 litres), and tartaric acid (30 g.) are left for some days, after which there is added putrid cheese (250 g.), sour skimmed milk (8 kilos.), and chalk (3 kilos.). The mixture is kept at 30° to 35° with occasional stirring. Calcium lactate is first formed, and this is afterwards decomposed with evolution of hydrogen (v. Formation 2); at the end of six weeks the evolution of gas ceases and the whole is now converted into calcie Latyrate (Bensch, A. 61, 177, cf. Grillone, A. 165, 127; Lieben a. Rossi, A. 158, 146; and Fitz, B. 11, 52). Crude butyric acid may be purified by etherification, followed by saponification of the butyric ether (121) (Bannoff, B. 19, 2552).

Properties .- Liquid, miscible with water, of powerful unpleasant smell. Separeted from aqueous solution by CaCl... Its barium salt is more soluble in alcohol than those of formic, acetic, or propionic acids. Its calcium salt is ppd. by boiling a solution saturated in the cold.

Reactions.—1. Boiled with HNO sit gives succinic acid (Dessaignes; Erlenn yer, A. 180, 207).—2. With CrO, it gives CO, and acctic acid (Grünzweig a. Hecht, B. 11, 1053).—3. With MnO₂ and dilute H₂SO₄ it gives propyl butyrate (Veicl, A. 148, 164).—4. The silver salt decomposes on dry distillation according to tho equation:

 $4C_3H_7.CO_2Ag = 3C_3H_7.CO_2H + CO_2 + C + 4Ag$ (Iwig a. Hecht, B. 19, 240). - 5. Distillation over zino-dust at 350° gives di-propyl ketone, propylene, CO, H, and other products (Jahn, B. 13,

2115).

Salts.-NaA'.-KA'. S. 125 at 15°. Very deliquescent.—MgA' ōaq. Very soluble plates.-CaA'₂. S. 20 at 0°; 18 at 22°; 15 at 60°; 16·2 at 100° (Chancel a. Parmentier, C. R. 104, 474; Hecht, A. 213, 69). Trimetric needles.— CaA', aq. S. 19.6 at 22°.—(CaA',),2(C,H,O,) 5aq. CaA' CaCl (C, H, O2), - CaCl2(C, H, O2), 2aq (Lie-

ben, M. 1, 926). — SrA'₂. S. 40 at 22°.— BaA'₂ 4aq. S. 40 at 14°. S. (alcohol) 11.7. Trimetrio. — BaA', 22aq. — BaA', 2C, H₈O₂ (Mixter, Am. 8, 343).—BaA', 2CaA', Pogular octahedra (Fitz, B. 13, 1314). ZnA', 2aq. S. 11 at 16°. Monoclinic prissis. Gives pp. of basic salt on boiling. —PbA'₂: oil. —PbA'₂: 2PbO. —PbA'₂:2CaA'₂: eules. — CuA'₂ 2aq. Monoclinic. — CuA'₂ aq. Triclinic (Alth. 4. 91, 176). —CuA'₂ 2Cu(AsO₃) Wöhler, A. 94, 44).—AgA'. S. 343 at 4.65 Needles or monoclinic prisms.

Methyl ether (192°). S.G. ⁶ · 9194 (Gartenneister, A. c233, 249); ⁶ · 9194 (Elsässer, A. 218, 314); ⁴ · 948 (Kahlbaum, B. 12, 344); ² · 8962 (Brühl); ¹⁵ · 9937; ²⁶ · 8945 (Perkin, C. J. 45, 483). M.M. 5·387 at 16·4° (k.). S.V. 120°35 (R. Schiff, A. 220, 332). μ_β 1·3936 (Brühl). R _∞ 43·11 (B.). C.E. (0° · 10°) · 001156 (E.).

43 11 (B.). C.E. (0° -10°) Vol116 (E.). Ethylether.—Mol. w. 116. (121° cor.). V.D. 3-99 (for 4-00) (S.). S.G. $\frac{9}{6}$ -9004 (Gartenmeister); $\frac{9}{6}$ -8996 (E.); $\frac{12}{8}$ -898 (Linnemann a. Zotta, A. 161, 178); $\frac{39}{4}$ -8892 (B.); $\frac{11}{12}$ -8849; $\frac{32}{82}$ -8762 (P.). M.M. 6-477 at 16-1° (P.). S.V. 150-23 (S.). μ_{β} 1-4007. 1t $_{\infty}$ 50-33 (B.). C.E. (0°-10°) ·001162 (E.). Smells like pine apples, in which it occurs as well as in other fruits.

Allylether (140°) (Caliours a. Hofmann, Tr. 1857, 555; A. 102, 296); (145°) (Berthelot

a. De Luca, A. 100, 360).

Propyl ether (113° cor.). S.G. 2 8930 (E.); 12 1.879 (Linnemann, A. 161, 53). S.V. 173.85 (S.). C.E. (0°-10°) 001077 (E.).

Iso-propyl ether. (128°). S.G. 2 ·879; 14 ·865 (Silva, A. 153, 135). n-Butyl ether (165° cor.). S.G. § ·8878

(G.); 12 ·876 (Linnemann, A. 161, 195); 2 ·889; 20 ·872 (Lieben a. Rossi, A. 158, 170). S.V. 197·8 (G.). C.E. (0° 10°) ·00105 (G.).

Iso-butyl ether (158°, (S.); 157° (E.). S.G. ? 8818 (E.); 2 880; 26 866 (Grünzweig, A. 162, 207). S.V. 197 66 (S.); 200 53 (E.). C.E. (0 -10°) 001093 (E.). Velocity of bromination: Urech B. 13, 1693.

n-Amyl other (184.8°). S.G. 9 8832. C.E. (0°-10°) .00099. S.V. 222.3 (Ga.).

Iso-amyl ether. (176°) (Delffs, A. 92, 278); (1786°) (E.). S.G. 4 8823 (E.). S.V. 221 52 (E.). C.E. (0°-10°) 001014 (E.).

Hexyl ether. (205°). S.G. 8 8825. C.E. (0°-10°) 00096. S.V. 246'4 (Ga.). From Hera-eleum (Franchimont a. Zincke, A. 163, 198). Heptyl ether (225°). S.G. § 8827. C.E. (0°-10°) 00093. S.V. 270-2 (Gartenmeister).

Octylether. (245°) (R.); (242°) (G.). S.G. 3 8794. C.E. (0°-10°) 00091. S.V. 295·6 (G.). From Pastinaca sativa (Renesso, A. 166, 80).

Cetyl ether. [20°]. (c. 265°) at 200 m.m. S.G. 29 .856 (Dollfus, A. 131, 285). Ethylene ether v. Glycol.

Ethylidence other v. Di-butyryl ortho-Алденчое р. 106.

Glyceryl ether v. GLYCERIN.

Amide C₃II₇.CONH₂. [115°]. (216°). Prepared by heating ammonium butyrate under pressure at 230°; the yield is 75 p.c. (Chancel, A. 52, 294; Buckton a. Hofmann, C. J. 9, 241; B. 15, 982). - Hg(C,H,NO).

Anilide C, H, CO.NH(C, H,). [92°]. Pearly plates (from dilutealcohol). Sol. ether. Formed by heating butyramide, butyric anhydride, or chloride with aniline (Gerhardt, A. Ch. [3] 37,

829 : Kelbe, B. 16, 1200)

Ohlorids C₃H..COCl. Mol. w. 106·5. (101°). S.G. 2 1 2 2 2 3 2 1 2 2 2 3 2 3 2 1 2 2 2 3 2 Linnemann, A. 161, 179). Converted by sodinmamalgam into dibutyryl (C,II,O)... Al.Cl., forms crystalline $C_{12}H_{18}O_3$ [107°] 'butyro - butyryl - butyric anhydride.' NaOH forms $C_{12}H_{19}NaO_4$ (Combes, C. R. 104, 853).

Bromide C.H.O.Br. (128°) (Berthelot, J.

1857, 344).

Iodide C4H,O.I (147°) (Cahours, A. 101,

111).

Aniv; dride (C,H,O),O. (192°). S.G. 12 978.

V.D. 538 (obs.). From sodium butyrate (4 pts.) A. 87, 155). Also from butyryl chloride and butyric acid (Linnemann, A. 161, 179). Heated with sodium butyrate at 180° it forms di-propylketone (Perkin, C. J. 49, 325).

Peroxide (C.H.O), O. From butyric an-hydrido and Bz.O. Oll (Bredie, Pr. 12, 655). Nitrile C.H. C.N. Propyl cyanide. Mol. w.

69. (119°). S.G. 13 '795. Formed by distilling the amido or ammonium butyrate with P2O3 (Dumas, A. 64, 334; Henke, A. 106, 272).

(Bulmas, A. 04, 554; itelike, A. 100, 272).

Isobutyric acid (CH₃),CH CO₂H. Mol. w.
88. (153° cor.). S.G. $\frac{1}{4}$: 9539; $\frac{2}{5}$: 9157 (Perkin, C. J. 45, 487); $\frac{2}{5}$ ° 9190 (Brühl); $\frac{2}{5}$: 9651 (Zander).
C.E. (0°-10°) · 00110 (Z.). S. 20 at 20°. M.M. 4·179 at 17·8° (P.). S.V. 108·57 (R. Schiff, A. 220, 105). μ_{β} : 1·3979. R_{20} : 35·48 (B.). S.H.
435 at 0° (Schiff, A. 234, 300). Heat of solution of the content of the 973. Heat of neutralisation in dilute solution 13989 (Gal a. Werner, Bl. [2] 46, 801). Vapour-pressure: Richardson (C. J. 49, 174).

Occurrence .- 1. In St. John's bread, the fruit of Ceratonia siliqua (Grünzweig, A. 158, 117; 162, 193) .- 2. In the root of Arnica montana (Sigel, A. 170, 345). - 3. As an ether (isobutylic?) in Roman oil of chamomile (Kopp, A. 195, 85; Köbig, A. 195, 96).-4. In human excrement

(Brieger, E. 10, 1029).

Formation. -1. From iso propyl cyanide and potash (Markownikoff, A. 138, 361).-2. By saponifying di-methyl-aceto-acetic ether (Frankland a. Dnppa, A. 138, 337). -3. Aqueous calcium butyrate which had been heated and cooled in a sealed tube 30 or 40 times in 10 years was found to have changed to the extent of 10 p.c. into calcium isobntyrate (Erlenmeyer, A. 181, 126) .- 4. By the oxidation of pyroterebic acid (Williams, B. 6, 1094).

Preparation.-By adding K.Cr.O. (4 pts.) to a cold mixture of isobutyl alcohol (3 pts.), H.SO. (5½ pts.) and water (15 pts.). Isobutyl isobutyrate separates. It is distilled with moist potash, and the potassium salt is distilled with strong H2SO4

(Pierre a. Pucbot, A. Ch. [4] 28, 366).

Properties.—Unpleasant smelling liquid.
Reactions.—1. Oxidised by CrO, mixture at 140° to CO2, acetone (Poposs, Z. 1871, 4) and acctic acid (Erlenmeyer, Z. [2] 7, 57).--2. Oxidised by KMnO, in alkaline solution, to B-oxyisobutyric acid, (CII,)2C(OH).CU2H, according to Richard Meyer's rule that when the group CH is united to three carbon atoms it may be exidised to C.OH.-3. Calcic isobutyrate on distillation gives di-isopropyl-ketone, with smaller quantities of methyl tert-butyl ketone, isobutyric aldehyde, and isobutyric acid (Barbaglia

a. Gucci, G. 11, 84).

Salts.—More soluble than those of n-butyric acid.—CaA'₂ aq: small plates. S. (of CaA'₂) 20 at 0'; 8 at 80°; 25 at 100°.—CaA' 5aq: long monoclinic prisms (Chancel a. Parmentier, C. R. 104, 477). —SrA'₂5a₁. S. 44 at 17° (hydrated) (Grünzweig). —BaA'₂1aq. Monoclinic. — BaA',11A' [74°] (Mixter, 4m. 8, 346). — BaA',2Ba(C.H.₂O.), aq. —ZnA',2aq. S. (hydrated). 17 at 20° —PbA'₂. S. 9 at 16°. Trimetric plates. Melts under hot water.—AgA'. S. 93 at 16°. Plates.

Methyl et er. (92°). S.G. 2 9112 (Elsisser, A. 218, 332). C.E. (0°–10°) 901223 (E.). S.V. 126.5. H.F. p. 109,660. H.F. v.

116,760 (Th.).

Ethyl ather. (110°). S.G. 1 8904 (E.); 15 8758; 25 8670 (Perkin, C. J. 45, 487). M.M. 6·479 at 21·8° (P.). C.E. (0° 10°) ·001156 (E.). S.V. 148·86 (E.); 150·68 (Schiff, A. 220, 333).

Propylether. (135°). S.G. 9 8813 (E.). C.E. (0³-10°) ·001039 (E.). S.V. 173·7 (E.);

171·2 (S.).

Iso-propyl ether (120°). S.G. 2 879

(Pribram a. Handl, M. 2, 691).

Iso-butyl ether. (116·6°) (E.); (149°) (S.). S.G. 5 8750 (E.). C.E. (02-102) 000094 (E.). S.V. 198·2 (S.); 196·0 (E.). Iso-amyt ether. (169°) (E.). S.G. § 8760. C.E. (0°-10°) 001031. S.V. 223 04.

Benzyl ether v. p. 493.

Amide [129°]. (c. 218'). Prepared by heating annuonium isobutyrate at 230° under pressure; the yield is 90 p.c. (Hofmann, B. 15. 982; cf. Letts, B. 5, 672; Münch, A. 180, 340; and Disiso-Butyramor).

Bromo-amide C.H.CO.NIIBr. Prepared by the action of bromine and KOH on isobutyramide (Hofgann, B. 15, 755). Large colourless needic, soi, other, sl. sol, water. Decomposed by caustic alkalis into propylamine, HBr, and CO₂, but by carbonated alkalis the reac-tion stees half way with production of propyl cyanate.

Iso-propyl-amide PrCO.NHPr. [102°]. (210°). Formed by the action of acetyl chloride on di-isopropyl acetoxim (Meyer and Warrington, C. J. 51, 685). Also by the action of isobutyryl chloride on isopropylamine. Colourless transparent needles v. sol. alcohol and ether, ne sol. water Sublimes at ordinary temperatures as I distils without decomposition. Is decomposed by prolonged boiling with alcoholic potash into isobut ric acid and isopropylamine.

Anilide Call., CO. HC. H., [103°]. From isobutyric acid and amiline (Norton, Am. 7, I16).

p-Bromo-anilide CaH, CO.NH.C.H, Br [1:4]. [128]. From the preceding and Br (N.). Chloride C.H., Colcl. (92) (Markowniskoff, Z. 1866, 501). S.G. 20 1-0174 (Brühl, A. 203, 20). $\mu_{\rm S}$ I-4135. R. 41-41 (B.). ZnMe₂ (I mol.) followed by water converts isobutyryl chloride into a ketone C₁₀H₁₈O (190°). S.G. 2 870 (Pawlow, A. 188, 139). ZnMe₂ (2 mols.) followed by water forms tertiary butyl alcohol and sometimes a ketone C₁₂H₂₂O (218°). S.G. 4 .864.

Bromide (117°). Anhydride (C.H.O),0. (182*). S.G. 13 9574 (Markownikoff, Z. 1866, 501; Tönnies a.

Staub, B. 17, 850).

Nitrile (CH3)2CH.CN. (108°). From icopropyl iodide, KCN, and alcohol (Marko inikoff, Bl. 1866, 53). From isobutyric acid and potaesium culphocyanide (Letts, B. 5, 669).

neBULYRIC ALDEHYDE C.II.O i.e. CH₃·CH₂·CH₂·CHO. (74°). Mol. w. 72. S.G. $\frac{10}{2}$ ·8170 (Brühl). μ_{θ} 1·3893. $\frac{1}{1}$ R₂₀ 32·93. S. 3·7. Got by distilling calcium butyrate with calcium formate (Linnemann, A. 161, 186; Lipp, A. 211, 355). From casein by oxidation with H.SO, and MnO, (Guckelberger, A. 64, 39).

Reaction. -- Aqueous NaOII and NaOAc form oily C_sH₁₁O (173°). It is probably Pr.CH₂CH:CEt.CHO as it reacts with phenylhydrazine and combines with bromine (Raupen-

strauch, M. 8, 108).

Ammonia compound

(CH₃)₂CH.CH(OH)(NH₂)3 aq. [31°]. Trimetrio pyramide. V. sl. sol. water, v. sol. alconol, ...
sol. ether. Deliquesce above 4°, giving off water.

Bisulphite compound C.H. ONAHSO,

17 0505 Kahn. B. 18, 3304).

Butyraldines. Dibutyraldino C₈H_{1,1}NO and tetra-butyraldino C₁₈H₂NO₂ are formed by the protracted action of alcoholic ammonia on butyric aldeliydo (Schiff, A. 157, 352). Butyral-

dine, on distillation, gives para-contine C.H.N. Butyral C.H.O (?) (95°). S.G. 22 821. A productof distillation of calcie butyrate (Chancel, A. Ch. [8] 12, 146; Limpricht, A. 90, 111; 93, 241). Reduces Ag.O. Does not combine with NH₂. Combines with NaHSO₃.

Reactions.-1. Air or Ag₂O forms butyric acid.—2. Chlorine forms C₄H,ClO (141°) and C₄H₂Cl₂O (200°).—3. PCl₅ forms C₄H,Cl (c. 100°). Isobutyrio aldehyde (CH3) CH.CHO. (64°). S.G. ½ 7972; ½ 7879 (Perkin, C. J. 45, 476); 2 7938 (Brild), A. 203. 18) S. 11 at 20°. μ₂ 13777. R_∞ 32×89. H.F.p. 61,340. H.F.v.

59,310. M.M. 4.321 at 19.3°

Formation.-1. From isobutyl alconol by chromic mixture (Pfeiffer, B. 5, 699; Michaelson, A. 133, 182; Pierre a. Puchot, C. R. 70, 434).—2. By heating ico-butyler.e bromide, (CH₂)₂CHBr.CHBr, with water (20 vols.) at 160° (Linnemann a. Zotta, A. 162, 36).-3. By distilling calcium isobutyrate (Popoff, B. 6, 1255; Barbaglia a. Gucci, B. 13, 1572).—4. By distilling calcium isobutyrate with calcium formate Linnemann a. Zotta, A. 162, 7). 5. By dietilling colophony (Tilden, B. 13, 1604).

Preparation. - A mixture of conc. aqueous K2Cr2O7 with an equal volume of H2SO4 is slowly run into a flast containing iso-butyl alcohol (100 g.) and water (200 g.) until the layer of alcohol has disappeared. The product is distilled. The yield is 55 p.c. of the theoretical (W. H. Porkin, jun., C. J. 43, 91; cf.

Fossek, M. 4, 660).

Properties.-Pungent liquid. Forms a compound with NaHSO, from which it is separated

by potash without change.

Reactions.-1. By the action of potash (4 g.) in alcohol (140 g.) upon the aldehyde (50 g.) the following bodies may be obtained; isobutyrio acid, an acid $C_{12}H_{22}O_{1}$ (245°-255°) and an aidebyle $C_{11}H_{22}O_{2}$ —2. If more potach (8 g.) and a Ag_2O . V.D. 167 (Theory 842). With Ac_2O it

higher temperature be used, the neutral products are: $C_{12}H_{22}O_{2}$, $C_{12}H_{24}O_{2}$?, $C_{22}H_{34}O_{4}$, $C_{23}H_{44}O_{4}$, $C_{23}H_{44}O_{4}$ and $C_{23}H_{44}O_{4}$ (W. H. Perkin, jun., C. J. 43, 101).—8. Aqueous potash forme an acid $C_{23}H_{14}O_{4}$ [75°–80°], a crystalline body C₅H₁₆O₂ [90°], and di-oxy-octane (Fossels M. 3, 622).—4. PCl₈ gives chloroisobutylene Me C:CHCl (68°) and di-chloro-isobutane (104°) Me CH.CHCl2 (Oeconomides, C. R. 92, 884).—5. H.S and aqueous ammonia form isobutyraldine C₁₂H₂₃NS₂ (Pfeiffer, B. 5, 700). 6. CS₂ and cone. NH₃Aq give NH₂CS.SN(C₄H₉). [91°]. Prisms, insol. water, v. sol. alcohol.-7. Alcohol and HCl followed by NaOEt form di-cthyl-ortho-isobutyrio aldehyde CMc_CH CH(OEt)₂ (135°). S.G. 12 996, V.D. 143.5, and, when some water is also present, a compound $C_{10}H_{20}O$ (223°) (Occonomide, Bl. [2] 36, 210; C.R. 92, 886).—8. Gives with ammonia 30, 217, 314; B. 13, 906; 14, 1746).

205, 1; 211, 344; B. 13, 906; 14, 1746).

7C₂H₁CHO + 6NH₂ = 6H₂O + (C₂H₁CH)₂N₂H₂O. When the product, 'oxy-hepta-iso-butylideneamino [32°], is heated, it first splits up into 2NII, C₄II₈O and 2(C₄H₈)₃N₂. The latter is hydro-butyramide, an oil, nearly insoluble in water, v. sol. alcohol or ether. If quickly heated it distils at 154°, but if heated slowly it splite up into NH, and C.H., N. Hydro-butyramide or tri-isobutylidene-diamino ie not affected by boiling KOH, but dilute IICl splits it up into butyric aldehyde and NH₃. It is, therefore, C₄H₈: N.C₄H₂. N:C₄H₈. Dry HCN added to its othereal solution forms the dihydrochloride of (Cy.C, II, NH)2C, H, a body that ie decomposed by water into isobutyric aldehydo and amidovalero-nitrile. When hydro-butyramide ieslowly heated it does not, like hydro-benzamide, change into an isomeride, but splits up, giving CaH15N. This compound, 'iso-butenyl-butylidene-amine,' ie a liquid (145°-147°) at 715 mm., nearly insoluble in water, miscible with alcohol or ether. It is not affected by aqueous KOH, but steids eplit it up into isobutyric aldehyde and NH₃. It would thue appear to be (CH₃)₂CH.CH:N.CH:C(CH₃)₂. It combines with Br, forming C₃H₁₃NBr₂, a body that, when kept for a long time, and then treated with water, gives NH, Br, ieobutyric aldehydo and bromo-isobutyric aldehyde, or rather a polymerido of the lattor [129°]. If C₈H₁₅NBr₂ be at onco treated with water, the unstable liquid bromo-butyric

aldehydo is probably formed (Lipp.).

Oxim C₂H₃.CH:N(OII). [139°]. Colourless liquid. Sol. water. Formed by the action of an asueouecolntion of hydroxylamine on isobutyrio

aldehyde (Petraczek, B. 15, 2784).

Description of contensation products, obtained as above (W. H. Perkin, jun., C. J. 43, 90).

Acid C₁₂H₂₁O₃ (245°-255°). Brownish oil.

Reduces ammoniacal Ag.O.

Compound C₁₂H₂₂O₂ (154°-157°). Oil. Ethereal odour. Reduces ammoniacal Ag₂O. Combines slowly with NaHSO,. Decomposee on prolonged heating. Is probably Urech's C₂H₁O (B. 12, 191). With Na and wet ether, it is reduced to C₁₂H₂₈O₂ (170°-175°), an alcohol (?) which does not combine with NaHSO,

Compound C₂₀H₃₅O₄ (223°-225°). Oil Smells of camphor. Very elowly combines with

forms C, Ha, AcO, (240°-242°). Oil, which with Ac.O at 200° gives $O_{20}H_{14}Ao_{1}O_{4}$ (248°-252°). When saponified by KOH a body $O_{20}H_{42}O_{4}$ (c. 220°) is formed.

Compound C20H44O4 (250°-253°). Oil. V.D.

(air = 1) 12.9 (Theory 13.7).

Compound C₂₈H₄₈O₈ (227°-229°) at 100 mm. Thick oil. Decomposed when heated under

atmospherio pressure.

Di-isobutyrio di-aldehyde C_sH_{1a}O₂. (138°) at 18 mm. V.D. 5·2 (calc. 5·0). This polymeride of isobutyrio aldehyde is obtained, together with octenoio aldchyde (?) C_sH₁₄O (15•°) by heating isobatyric aldehyde with cone. aqueous NaOAe at 50° (Fossek, M. 2, 622). It is an oil, sol. alcohol and ther, forming a crystalline compound with NaHSO.

Iso-butyrie paraldehyde (C,H₀O)₃. [60°]. (195°). V.D. (H=1) 104.8. From iso-butyraldehyde by H₂SO₄, HCl, PCl₃, Cl, Br, or I (Barbaglia, B. 5, 1052; 6, 1064; G. 16, 430; Deintschenko, B. 6, 1176). Needles (from water or by sublimation). Difficultly attacked by oxidising agents (Urech, B. 12, 1749). Does not combine with NaIISO, or react with NH3. At 150° it partially changes to ordinary isobutyric aldehyde.

Iso-butyric poly-aldehyde (C, fl,O), S.G. 24 *969. Prepared by leaving isobutyric aldchyde in contact with dry Na CO3. Thick liquid. Sl. sol. water. Decomposed on distillation, with separation of water and formation of isobutyrie aldehyde and condensation products (Urech, B.

12, 191, 1744; 13, 483, 590). BUTYRIN v. GLYCERIN.

BUTYRO-CHLORAL v. TRI-CHLORO-BUTYRIO

BUTYRO-COUMARIC ACID v. OXY-PHENYL-

BUTYRO-CREATININE v. METHYL-AMIDO.

BUTYRIC ACID.

BUTYRO-FURONIC ACID C, II, O5 i.e. CO.H.CH:CH.CO.CH..CH..CH..CH..CO.H [142°]. Prepared by treating furfurvalers acid with bromine-water and subsequent action of silver exide. White crystals. Sol. water and alcohol, sl. sol. ether. By III and P it is reduced to azelaio acid (Toennies, B. 12, 1200).

BUTYRO-LACTONE v. γ -Oxy-BUTYRIC ACID. BUTYROLIC ACID v. TETROLIC ACID.

BUTYRONE v. DI-PROPYL-KETONE.

BUTYRONITRILE v. Nitrile of BUTYRIC ACID.
BUTYRO-PINACONE C14H30O2 v.e.

CPr₂(OH).CPr₂(OH). Di-oxy-tetradecanc. [68°]. (260°). From di-propyl-ketone, water, and Na (Kurtz, A. 161, 215). Crystals, smelling of camphor, sl. sol. water.

BUTYRO-THIÊNONE v. THIÊNYL PROPYL

KETONE.

DI-BUTYRYL C₂II₁₄O₂ i.e. Pr.CO.CO.Pr. Di-propyl di-ketone. (245°-260°). From butyryl chloride and sodium-amalgam or zine (Freund, A. 118, 35). Yellow oil. Boiled with potash it forms butyrate of potassium and a liquid C. H., O which does not unite with NH2 or NaHSO,

Mono-oxim C,II, CO.C(NOH).C,H,: thick oil; can be distilled in small quantity without decomposition. A di-oxim has not been obtained (Münchmeyer, B. 19, 1846).

BUTYRYL-ACETOPHENONE

C₄H₃.CO.CH₂.CO.C₃H₇. Benzoyl-methyl-propylketone. (174° at 24 mm.). S.G. 1.061 at 15°. tondi, G. 4, 192; Alessandri).

Colourless oil. Formed from acetophenone and butyrio ether by EtONa (Beyer a. Claisen, B. 20, 2181).

2181).
Isobutyryl-acetophenore
Benzoyl-methyl-isoColourless. C₆H₄.CO.CH₂.CO.C₃H
7. propplectone. (170° at 26 mm.). Colourless liquid. Formed from acetophenone and isobutyrio cther by EtONa (Beyer a. Claisen, B. 20, 2181).

BUTYRYL-AMIDO-BENZOIC ACID PrCO.NH.C, H, CO.H. [209°]. From n-butyric ether (20 c.c.) and m-anddo-benzoic acid (10 g.) at 180° in scaled tubes (Pellizzari, A. 232, 148). Sol. water and alcohol.

BUTYRYL LROMIDE v. BUTTRIC ACID. BUTYRYL CHLORIDE v. BUTYRIC ACID. BUTYRYL CYANAMIDE v. CYANAMIDE. n-BUTYRYL CYANIDE C.II NO i.e.

PrCO.CN. (133°-137"). From AgCN and PrCOCI

(E. Moritz, C. J. 39, 16).
n-Di-butyryl di-cyanide (PrCO)₂(CN)₂ (c. 234°). By-product in preparation of above.

Iso-batyryl cyanido Pr.CO.CN. (117'-120°). From PrCO.Cl (40 g.) and AgCN (50 g.). Bad yield (E. Moritz, C. J. 39, 13). The greater part of the product is di-iso-butyryl di-cyanide (226°-228"). S.G. 96.
BUTYRYL IODIDE v. BUTYRIC ACID.

BUTYRYL-MALONIC ETHER

 C_3H_7 .CO.CH(CO₂Et)₂, (247° 252°). Formed by the action of butyryl chloride upon sodio-malonic ether. By nitrous acid it is converted into isonitroso-butyryl-acetic other (Lung, B. 20, 1826). BUTYRYL PEROXIDE v. Butyrae acid.

BUTYRYL-PROPYL-UREA v. Butyryl derivative of Propyl-ukea.

BUTYRYL SULPHOCYANIDE (180°). From hutyryl chloride and lead sulphocyanide. Decomposes when boiled (Miquel, A. Ch. [5] 11 295).

BUTYRYL-UREA v. Ukua.

BUXEINF. An alkaloid extracted by dilute oxalic acid from the back of the box-free Buxus sempercirens. Yehowish-white crystalline substance, sol. alcohol and other, st. sol. water. IINO, sives a greenish-yellow colouration turning brick-red. H2SO, gives a blood-red colour. Chromic acid mixture gives an orange pp. (Alessandri, G, 12, 96). It is perhaps identical with buxine. Barbaglia finds four alkaloids in the leaves and twigs of the box: buxine, parabuxine, buxidine, and buxinidine (G. 13, 249; B. 17, 2055)

BUYINE C19II. NO3(?). An alkaloid extracted by dilute ox lie acid from the leaves of the box tree. White crystalline substance, sol. alcohol and other, sl. sol. water. HNO3 gives a purple-red colouration. H₂S₁A₂ vives a brick-red colour. Chromic acid mixture gives a canary-yellow pp. (Alessandri, G. 13, 96; Barbaglia, B. 4, 757; Fauré, J. Ph. 16, 428; Concelle, J. Ph. 1854, 51). According to Wa'z (J. 1860, 548) buxine is identical with beneering (q, v).

Parabuxine C_{us}H_{us}: •QO'(?) An alkaloid occurring in both leaves and back of the box tree. It is a reddish-purple amorphous resin, sol. water and alcohol, insol. ether. HNO, gives a permanent greenish-yellow colouration. ILSO, gives a greenish-yellow colour becoming dark. Chromic acid mixture gives no pp. B"H2SO4.-B"2HCl.-B"H2PtCl (Pavesi a. KoCACAO v. THEOBREMINE.

CACODYL v. p. 318. CACOSTRYCHNINE C21H22N3O10. A hroduct of the action of HNO3 on strychnine (q. r.). Golden moedles (from dilnte HNO3) or hexagonal plates (from conc. HClAq). Sl. sol. most menstrna, sol. alkalis, forming red solutions.-B'2H_PtCl₆ (Claus a. Glassner, B. 14, 773).

CACOTHELINE C 2011 22 N On. A product of the action of HNO, on brucine (q. v.). Orange lamine (containing aq). Weak base; sol. alkalis, v. sl. sol. hot water, insol. alcohol and ether. -B'BaO 7aq. - B'.H.PtCl. (Strecker, .1. 91, 76; C. R. 39, 52; Rosengarten, A. 65, 111; Claus a. Röhre, B. 14, 765).

CADAVERIC ALKALOIDS v. PTOMATNES. CADET'S FUMING LIQUID r. p. 318.

CADMIUM. Cd. At. w. 1117. Mol. w. 111-7; gaseous molecule is monatomic. [320] (Person, A. Ch. [3] 27, 250; Rudberg. P. 71, 460; v. Riemsdyk, C. N. 20, 32). (763'-772°) (Carnelley a. Williams, C. J. 33, 284). S.C. (molten) 8-65, (hammered) 8-8 (Stromeyer, S. 22, 362; Schröder, P. 106, 226; 107, 113; Matthiessen, P. 110, 21, &c.). V.D. 558 (Deville a. Troost, C. R. 52, 920). S.H. (0° 100°) 0548 (Bunsen, P. 141, 1), 0567 (Regnault, 4. Ch. [3] 26, 268). C.E. (linear, 0°-100°) ·003223; (oubled for 1°) ·000094 (Kopp, A. 81, 32; Matthiessen, P. 130, 50; Fizeau, C. R. 68, 1125). T.C. (Ag = 100) 20 06 (Lorenz, W. 13, 422). E.C. (Hg at 0'=1) at 0', 13 46; at 100°, 9.5 (Lorenz, W. 13, 422 a. 582). Heat of fusion 13,660 (Person, P. 76, 426). S.V.S. abt. 12.8. Emission-spectrum characterised by lines 3609.6, 3465.4, 2717.7, 2572.2, 2313.6, 2288.9 (Hartley, T. 1884. 63).

Cd was discovered by Stromeyer in 1817 in a specimen of zine carbotate (S. 21, 297; 22, 362; v. also Hermann, G. A. 59, 95). The name cadmium was derived from cadmic, jossilis by which name zine ore was then known.

Occurrence.-With Zn in various native sulphides, earbonates, and silicates, especially in the Silesian zine ores (v. Damou., J. pr. 13, 354; Stadler, J. pr. 91, 359; Blum, J. 1858. 734; Bunsen, A. 133, 108). CdS occurs nearly pure as Greenockite at Bishopton in Renfrew.

Formation. - In the distillation of crude zine oxido with charcoal; the greater part of the Cd distils over before the Zn.

Preparation. — ZuO containing CdO, or metallic Zn containing Cd, is dissolved in dilute H_SO, Aq or HClAq; the warm solution is saturated with H.S; the CdS thoroughly washed and dissolved in conc. HClAq; most of the free HCl is removed by warming, the solution is diluted and filtered, and an exec is of (NH4)2CO3 is added; the pp. of CdCO₃ is well washed, dried, and strongly heated; the CdO thus produced is mixed with $\frac{1}{10}$ of its weight of pure powdered charocal and heated in a retort of hard-glass or porcelain when pure Cd distils ovor (Stromeyer, **S.** 22, 362).

Properties .- White with slight blue tinge: very lustrous; soft, but harder than zine; very

malleable, ductile, and flexible; more tenacious than tin; crystallises easily in monometrio forms, chiefly the octahedron (v. Kümmerer, B. 7, 1724; also G. Rose, P. 85, 293). Vapour is yellow. Cd does not decompose water even at 100°; but if Cd vapour and steam are passed through a hot tube the steam is decomposed (Regnault, A. Ch. 62, 351). Cd oxidises slowly on the surface by exposure to air; when heated in air it burns to CdO. The atomic weight of Cd has been determined (1) by finding the V.D. of, and by analysing, CdBr₂ (Mayer, B₂₀12, 1292; Dumas A. Ch. (3) 55, 158; Huntingt Ta, P. Am. A. 17, 28); analyses of CdC.O, (Lenssen, J. pr. 79, 281); reduction of CdSO, to CdS (v. Hauer, C. C. 1857, 897); analyses of CdO (Stromeyer, S. 22, 366); (2) by determining the S.H. of Cd (Bunsen, P. 141, 1; Regnault, A. Ch. [3] 26, 268); (3) by comparing, as regards crystalline form and general reactions, salts of Cd with salts of Zn, Be, Mg, and Hg. In the gaseous molecule CdBr. (this is the only compound of Cd whose V.D. has been determined) the atom of Cd is divalent. The gaseous molecule of Cd is monatomic. Cd is a distinctly metallic element; it acts on IIClAq, H.SO,Aq, &c., evolving H and forming salts of the form CdX, where X2 - Cl2, Br2, SO4, CO3, &c.; many of these salts combine with the similar salts of the more positive metals (K, Ca, Mg, &c.) to form double salts; but few basic salts of Cd are known, the most marked are derived from such weak acids as H.CrO, H.BO, &c. No compound of Cd exhibits any acidic functions. CdO, H. acts towards acids as a salt-forming hydroxide; its heat of neutralisation by H.SO, Aq is about the same as that of the corresponding hydroxide of Mn, Ni, Co, Fc, or Zn, [CdO II Aq, II SO Aq] = 23,8° (v. Th. 1, 339 a. 436). CdO₂H₂ is dehydrated by heat; the oxide CdO is not converted to CdO.H. by direct addition of H.O. Cadmium is closely related to Zu, it is less positive than that metal; it is also related to Mg on one hand and to Hg on the other (v. MAGNESIUM GROUP OF ELEMENTS).

Reactions.-1. Heated in air, or O, CdO is produced. -- 2. Heated nearly to redness in bromine, CdBr., is formed. -3. Aqueous solutions of hydrochloric, sulphuric, or nitric, acids are decomposed by Cd with formation of chloride, sulphate or nitrate of the metal. -4. Heated with SO.Aq to 200° CdS is formed (Geittner, A. 129, 351); possibly sulphite and thiosulphate are first formed (v. Schweitzer, C. N. 23, 293; Fordos a. Gélis, A. 50, 260).

Combinations .- Most compounds of Cd are formed from the exide or other salt. Cd combines directly with the elements O, Cl, Br, I, P, S, Se, Te, and with many metals (v. Cadmium, exide of, &c., and Cadmium, alloys of).

Detection and Estimation .- Formation of the yollow sulphide, CdS, insoluble in dilute HClAq and also in solution of ammenium sulphide, characterises Cd salts. Cd is usually estimated by ppn. as CdCO₃ (by K₂CO₃Aq), the pp. is strongly heated, and the CdO is weighed. Separation from other metals may be effected by repeated ppn. by H₂S, and solution of CdS in cone. HClAq. Cd may be ppd. as oxalate; on this fact is founded a volumetric method of estimation.

Cadmium, Alloys of. Usually prepared by melting the metals together. Several are characterised by low melting-points. An amalgam with Hg is formed at ordinary temperatures: by dissolving Cd in warm Hg, and pressing, a erystalline amalgam, having the composition Hg,Cd2, and S.G. 12.62, is formed; by completely saturating Hg with Cd, octahedral crystals of Hg2Cd, melting at 77%, are produced (Gaugain, C. R. 42, 430; Regnauld, C. R. 51, 779; Crookewitt, J. pr. 102, 65 a 129; Kopp, A. 46, 186). • Easily fusible alloys with Bi agreeing in composition with the formula BiCd, BiCd, and BiCd, are known (Matthiessen, P. 110, 21). Various alloys of Cd with (1) Bi and Pb. (2) Bi, Pb and Su, (3) Bi and Su, are also known ([1] Wood, D. P. J. 164, 108; v. Haner, J. pr. 94, 436: [2] Lipowitz, D. P. J. 158, 376: [3] Wood, Lc.). Alloys with Pb (Wood, C. N. 6, 135); Na (Sonnenschein, J. pr. 67, 169); Tl, and with Tl and Bi (Carstenjeu, J. pr. 102, 65 a. 129); and Sn (Rudberg, J. 1817, 71), have been described. An arsenide, AsCd., is said to be obtained as a faintly red-coloured alloy, S.G. 6.26, by reducing the arsenate by KCN (Descamps, C. R. 86, 1022 a. 1065).

Cadmium, Arsenates of, $-\mathrm{Cd}_{g}(\mathrm{AsO}_{i})_{g}.311_{i}\mathrm{O}_{i}$ and Cd, H2(AsO1)24112O; v. ARSENATES, under ARSENIC, ACIDS OF.

Cadmium, Arsenide of. Cd. As v. Cadmium,

Cadmium, Bromide of. CdBrg. Mol. w. 271.2. [570°] (Carnelley, C. J. 33, 278). (806°-812) (Carnelley a. Williams, C. J. 37, 126). S.G. 205 4.794 (Clarke, Am. 5. No. 4). H.F. [Cd, lir2] = 75,200; [Cd,Br²,Aq] = 75,610 (Thomsen).

Preparation .- By heating Cd to redness in Br vapour; or by dissolving CdCO, in HBrAq

and subliming.

Proporties and Reactions .- White, crystalline, non-hygroscopie, solid; soluble in water, ether, and alcohol; decomposed by HNO,Aq (Bodeker a. Giesecke, J. 1860, 17; Croft, I. M. [3] 21, 355; Berthemot, A. Ch. 44, 387; Rammelsberg, A. 44, 267).

Combinations. - Dissolved in H.O. and evaporated, yields long white needles of CdBr₃.4H₃O; these are dehydrated at 200° $[CdBr^2,4H^2O] \approx 7.730$ (Thomsen). ${\bf CdBr}_{2}{\bf Aq}$ and KBrAq mixed and evaporated yield 2CdBr...2KBr.H.O, and on further evaporation

CdBr. 4KBr. The double salts

2CdBr₂.2NaBr.5H₂O, and CdBr₂.BaBr₂.4H₂O, have also been obtained (v. Haner, J. pr. 64, 477; 69, 121. Croft, J. pr. 68, 399). Cillir,

Cadmium, Chloride of. CdCl. Mol. w. un-known; probably as represented by formula OdOl, [541°] (Carnelloy, U. J. 50, 2.5), 564). (Carnelley a. Williams, C. J. 35, 564). 140; (100°) 150 (Kremers, P. 103, 57; 104, 133; next article.

105, 860). H.F. $[Cd,Cl^2] = 93,240$; $[Cd,Cl^2,Aq]$ =96,250 (Thomsen).

Preparation.—By dissolving Cd in HClAq and heating to low redness the crystals of CdCl_2H_O which separate on evaporation.

Properties and Reactions .- Pearly, lustrous plates; white powder after exposure to air for some time. S. (208-100°) alit. 140 (Kremers, P. 103, 57; 101, 133). Insoluble in cone. HClAq.

Combinations .- 1. With water to form elilorescent prisms of CdCl, 2H,O; [CdCl, 2HO] =2,250 (Thomsen); best obtained by dissolving Cd, CdO, or CdCO, in HClAq, evaporating, and crystallising. (2. With hydrochtoric acid and water to form 3dCl 2HCl.7HO: obtained by passing HCl into CdCl_tq; casily decomposed in uir with evolution of HCl; [CdCl,211Cl,711-0] - 40,200 (Berthelot, C. R. 91, 1021). - 3. With ammonia, to form CdCl...6NII, and CdCl...2NII... CdCl...6NII, is obtained by passing NII, over CdCl...or by passing HCl into CdCl...Aq containing excess of NIt, until the NII, is partly neutralised. Loses 4NII, by exposure to air, leaving CdCl...2NII, this compound is also produced by exposing to air a solution of CdCL in excess of warm NII3Aq (Croft, P. M. [3, 21, 55; Schäler, A. 87, 34; v. Huner, J. pr. 61, 477; v. also André, C. R. 104, 908).—4. With various metallic chlorides to form double salts. These salts have been prepared chiefly by Croft (P. M. [3] 21, 55), and v. Hauer (J. pr. 63, 432; 61, 477; 66, 176; 69, 121); their crystalline forms, and the thermal conductivities of some of them, have been determined by Grailich (J. 1858, 182), and v. Lang (P. 135, 29). These double compounds are obtained by evaporating mixed solutions of the two chlorides; the chief are:-

2CdCl_2NH_Cl_H_O; CdCl_BaCl_c4H_0; 2CdCl_BaCl_coH_0; CdCl_4N1LCL, 2CdCl_2LCLTLO; 2CdCl_SrCl_7H_0; CdCl. 1KCl; CdCl. 2NaCl.3H,J; 2CdCl, CaCl, 7H.O; CdCl_2CaCl_12H_0; CdCl_2MgCl_12H_0; CdCl_.MCl_.12H_O when M = Co, Fe, Mg, Mn, or Ni; CdCl_wCuCl_w4H₂O.

5. With the hydrochlorides of many organic bases to form double salts; e.g. with toluidine 4(C,H,N.HC!).3CdCl2:: H2O (Williams, C. C. 1856, 47; Galletly, U. C. 1856, 606).

Cadmium, Cyanide of. Cd(CN). Prepared by evaforeting KUNAq with CdCl.Aq (v. CYANIDES).

Cadmium, Fluorido of. CdF. Mol. w. un-known; probably as copresented by formula. [520°] (Carnelley, C. J. 33, 280). S.G. $\frac{227}{40}$ 5.994 (Clarke, Am. 5, No. 4). Hard, white, crystalline absorbs NH, to form CdBr...4NH,; all NU, is removed by heat. Conc. OlBr.Aq saturated with NH, and evaporated, gives crystals of CdBr...2NH, (Croft, P. M. [3] 21, 355; Rammelsberg, A. 44, 267). CdF. 2ZrF, 6H₂O, were obtained by Marignao (Ann. M. [5] 15, 221; A. Ch. [3] 60, 257); and the double compound CdF. MoO.F. 6H₂O by Delafontaine (J. 1867. 236).

Cadmium, Hydrated exide of, CdO.H.O v.

Cadmium, Hydroxide of, CdO₂H₂. A white, amorphous solid, obtained by adding KOHAq to a dilute aqueous solution of a Cd salt, washing with warm water, and drying at 100²-200² (Schaffner, A. 51, 168). According to Kickles (J. Ph. [3] 12, 406) CdO₂H₂ is obtained it crystals by keeping Cd in contact with Fe in NH₂Aq. De Schulten (C. B. 102, 72) describes the formation of hexagonal crystals of CdO₂H₂, S.G. ½0 4.79, by dissolving 10 grams CdI₂ in 150 cc. H₂O mixed with 360 grams KOH containing 13 p.c. H₂O, heating till all is dissolved, and cooling. Thomsen gives the thermal value [CdJO, H²O] = 65,680; and the following values for the heat of neutralisation of solid CdO₂H₂ (Th. 3, 285):

Q	[M,QAq]	$M = CdO_2H_2$.
H,SO,	24,200	
H ₂ N ₂ O ₄	20,620	
$\mathbf{H}_{2}\mathbf{S}_{2}\mathbf{O}_{1}$	20,360	•
II,Cl,	20,290	
$\mathbf{H}_{2}\mathbf{Br}_{2}$	21,560	
$\mathbf{H}_{2}^{T}\mathbf{I}_{2}^{T}$	24,210	

The quantity of heat produced with the three haloid acids increases as the atomic weight of the halogen increases; in this respect $\mathrm{CdO}_{-}\mathrm{H}_{2}$ is analogous to the corresponding compounds of Hg and Pb , and differs from those of Ba , Ca , Sr , and Mg . $\mathrm{CdO}_{-}\mathrm{H}_{2}$ loses $\mathrm{H}_{2}\mathrm{O}$ at 300° (II. Rose, P , 20, 152); $\mathrm{CdO}_{-}\mathrm{H}_{2}$ loses $\mathrm{H}_{2}\mathrm{O}$ at 300° (II. Rose, P , 20, 152); $\mathrm{CdO}_{-}\mathrm{H}_{2}$ in the thermal values given by Thomsen (Th. 3, 285; and P. 143, 354 a. 497) the reaction $\mathrm{CdO}_{-}\mathrm{H}_{2}\mathrm{O}_{-}\mathrm{CdO}_{-}\mathrm{H}_{2}$ would require the expenditure of about 10,000 units of heat.

Expenditure of about 1,000 mins of heat.

Cadmium, Iodide of, GiL, Mol. w. nn-known, but probably as represented by the formula. [401°] (Carnelley, C. J. 33, 278). (708°-719°, with decomposition) (Carnelley a. Williams, C. J. 37, 126). S.G. $\frac{abc, 15}{4}$ 5·644, and 4·626 (v. Preparation a. Properties). H.F. [Cd,F] = 48,830; [Cd,F,Aq] = 47.870 (Thomson). $V_t = V_o$ (1 + 000087487), t^* tot greater than 40° (Fizeau, C. R. 64, 314). S. (20.) 92·6; (60°) 107·6; (100°) 133°3 (Kremers, P. 103, 54; 104, 133; 111, 60).

Preparation and Properties. 1. By heating together Cd and I, in the ratio Cd: I., in absence of air. -2. By digesting together Cd and I under water (Stromeyer, S. 22, 362).-3. By evaporating a solution of 20 parts KI and 15 parts CdSO, to dryness, dissolving in alcohol, and crystallising (Vogel, N. R. P. 12, 393).—4. By dissolving CdCO, in IIIAq, decolourising by addition of elippings of Cd, and crystallising (Clarke, Am. 5, No. 4) .- 5. By dissolving Cd in IllAq, evaporating, and crystallising (Clarke, Lc.). According to Clarke (Lc.) Cdl, exists in two forms; the normal salt is white, is unchanged by heating to 250°, and has S.G. 5 644; the other salt is brownish, loses weight even at 40°, and has S.G. 4.626. The conditions under which the less stable salt is formed have not yet been exactly determined; Clarke obtained it twice by the action of 111Aq on Cd and on CdCOs. The S.G. of the less stable salt increases by heating to 50° for some time. If the formulaweight CdI, is divided by the S.G., the results are, for the stabler salt 61.8, and for the less stable salt 79.2; new S.V.S. of Cd+S.V.S. of L = 64.3 (Clarke, l.c.).

Combinations.—1. With ammonia to form CdI₂.6NH₂, and CdI₂2NH₃; obtained as the corresponding CdCl₂ compounds (q. v.): both are decomposed by H₂© with ppn. of CdO₂H₂ (Rammelsberg, P. 48, 153).—2. With some metallic iodides to form double salts; Croft (J. pr. 68, 399) described CdI₂.2KI.2H₂O (aqueous solution of this salt pps. most of the organic bases from plants; Marmé, N. R. P. *16, 306);

from plants; Marmé, N. R. P. 16, 306); CdI, 2NH, I.2H,O; CdI, 2NaI.6H,O; CdI, BaI, 5H,O; and CdI, SrI, 8H,O. Clarke (Am. 5, No. 4) obtained CdI, 3HgI, as gold-

coloured plater.

Cadmium, Öxids of. CdO. Mol. w. unknown. S.G (amorphous) 6.95, (crystalline) 8.11 (Stromeyer, S. 22, 362; Worther, J. pr. 55, 148; Schüler. A. 87, 34; Sidot, C. R. 69, 201; Follenius, Fr. 13, 279). H.F. [Cd,O] = 75,500 (calculated from data given by Thomsen, Th. 3, 285; P. 143, 354 a. 497).

Preparation.—As a dark-brown, amorphous, infusible, powder, by burning Cd in air or O; or by strongly heating CdO₂II₂, or CdCO₃. As black-brown very small octahedra (or other forms of the monometric system), by strongly heating Cd.2NO₃, or CdSO₄ (Werther, l.c.; Schü-

ler, l.c.; Herapath, B. J. 3, 109).

Properties de.—Reduced to Cd by heating with C. At red heat Cl forms CdCl₂. Readily combines with CO₂ forming CdCO₃. Dissolves in aqueous acid with production of Cd salts. Thomsen (P. 143, 354 a. 497) gives the thermal values, [CdO,H'SO'Aq]=14,240 for crystalline CdO, and 14,510 for amorphous CdO.

Marchand (I'. 38, 145) supposed that a lower oxide than CdO was formed when CdC₂O₄ was heated; but Vogel's experiments (J. 1855. 390) seem to prove that the substance was a mixture, in varying proportions, of CdO and

By the action of H₂O₂Aq (about 3 p.c. H₂O₂) on resist CdO₂H₂, Hass (B. 17, 2219) obtained an oxide of Cd containing more of than CdO. Analyses gave results agreeing fairly with the formula Cd₂O₃, in one case with Cd₂O₄. These results were confirmed by Bailey (C. J. 49, 484) who obtained Cd₂O₅ by the action of H₂O₂Aq on CdSO₂Aq followed by addition of NH₂Aq. The pp. obtained by these methods may have been a mixture, or possibly a loose compound, of CdO and CdO₂ (c. Haas, Lc. 2255).

Cadmium, Phosphides of. Cd and P are said

Cadmium, Phosphides of. Cd and P are said to combine when heated together to form a grey, slightly metal-like, mass, which burns in all to phosphates, and dissolves in HClAq with evolution of PH₃ (Stromeyer, S. 22, 362). According to B. Renault (C. R. 76, 283) when P vapour is passed over hot Cd, two phosphides are formed, Cd₃P₂ and Cd₂P. Oppenheim (B. 5, 979) describes Cd₃P₂ as a grey, metal-like, crystalline substance, produced by heating CdO with KOHAq and P, and heating in H.

Cadmium, Salts of. Compounds obtained by replacing the H of acids by Cd. The Cd salts form one series CdX., when X.=Cl., (NO₃), (ClO₃)... SO, CO₃, HPO₃, &c. The V.D. of one salt, CdBr₂, has been determined; from this result, and from the similarities between the salts of Cd and Zn, it is probable that the gaseous molecules of the haloid Cd salts are

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correctly represented by the formula CdX2 where X=F, Cl, Br, or I. The greater number of the salts of Cd are soluble in water; the squeous solutions redden blue litmus paper; they are poisonous. The haloid salts are not decomposed by heat; salts of volatilisable aoids give CdO when strongly heated. Many Cd salts are isomorphous with corresponding salts of Zn; come, especially more complex double salts, are isomorphous with corresponding salts of Mg, Ni, Co, and Zn. A great many double salts containing CdX2(X=Cl,Br,I) are known; but few basic salts of Cd have been prepared. The rincipal Cd salts are borate; bromate; carbonates, a chlorate, perchlorate; chromate; cyanates, a cidate, periodate; molyblate; nitrates, nitrites; phosphates, phosphite; selenates, selenite; sulphutes, sulphite, &c.; tangstate; vanadate; v. Borates, Candonates, &c.

Cadmium, Selenide of. CdSe. Golden yellow, motal-like, crystalline, mass; by heating Cd in Se vapour; S.C. 8 79 (Stromeyer, S. 22, 362). The same body is said to be formed as a darkbrown pp. by passing H.Se into solution of a Cd salt (Vigier, Bl. 1861. 5; Ronault, C. R. 76,

283).

Cadmium. Silicofluoride of. CdSiF., Long, columnar, deliquescent crystals, obtained by action of H.SiF.Aq on CdO (Berzelius, P.

1, 26).

Cadmium, Sulphide of. CdS. Occurs native in hexagonal prisms (a:c=1:S1257) as Greenockite. Obtained as an amorphous yellow solid, by repeatedly heating to dull redness CdSO, in H₂S (v. Hauer, J. pr. 72, 338); by passing 11.8 into a slightly acid solution of a Cd salt; by heating Cd with SO.Aq (Geitm., A. 87, 31; Fordos a. Gélis, A. 50, 260; Schweitzer, C. N. 23, 293). Obtained also in crystalline form by fusing the amorphous CdS with K.COa and S (Schüler, A. 87, 54); by heating CdCl2 in 11 S; or by melting together CdSO, CaF₂, and CaS (Troost a. Deville, C. R. 52, 920). CdS is also produced in crystals, but in small quantity, by passing S vapour over strongly heated CdO, or Cd (Follenius, Fr. 13, 411; Sidot, C. R. 62, 999). Crystalline CdS is non-volatile at any temperaturo; it dissolves easily in boiling conc. HClAq, or dilute H.SO,Aq (Follenius, Lc.; Hofmann, A. 115, 286); S.G. 45, when melted 46. Schiff (A. 115, 74) described edS, as a vellow powder obtained by the action of K.S.Aq on a noutral Cd sall in solution; according to Follenius (Fr. 13, 411) this yellow solid is a mixture of CdS and S.

Cadmium, Sulphocyanide of, Cd(CNS), Obtained by action of HCNSAq on CdCO,; v.

SULPHOCYANINES, under CYANIDES.

Cadmium, Telluride of. Calle. Black crystals, S.G. 6:20, by heating Cd with Te, and subliming the product in I1 (Margottet, C. R. 85, 1142).

M. M. P. M.

CADMIUM ETHIDE CdEt,. Obtained in impure condition from Etl and Cd. Takes fire in air (Wanklyn, C. J. 9, 193; Sonneuschein, J. pr. 67, 169).

CESIUM. Cs. At. w. 132.7. [26°-27°] A. 211, 100).—2. The mixed chlorides of Cs and (Setterberg, A. 211, 100). S.G. 12° 1.88 (Sotterberg, A. 211, 100). S.V.S. 70.7. Discovered by Bunsen phates, and then into alums by adding and Kirohoff as ohloride in the water of a Al.3SO, Aq and evaporating. Rb alum is 4 mineral spring at Dürkheim (P. 113, 342). times more soluble than Cs alum; Cs alum Vol. 1.

Name given because element characterised by two sky blue (cæsius) lines in the spectrum.

Occurrence. Never free Salts very widely distributed, but in very subill quantities, along with Rb, ebicily as chloride and oxido; in many minerals and mineral waters, in sea water and sea weed, in residuce from saltpetro refining, in ash of tobacco, tea, coffee, and oak wood. Co. especially Laspeyres, A. 131, 349; 138, 826: also Smith, Am. S. [2] 49, 335; Erdmann, J. pr. 86, 377; Grandeau, C. R. 53, 1100; Loustadt, C. N. 22, 25 a. 44). The rare mineral Polluza, from Elba, according to analyses by Pisani, contains 34 p.c. % oxide combined with silica,

and is free from Rb (A. 132, 31).

Preparation.—1. The mother liquor, obtained

by repeatedly evaporating the water of the mineral spring at Nanheim, and separating from the erystals which form, contains nearly 2 p.o. CsCl. Fe, Al, and the alkaline earth metals, are removed in the usual way; the liquid is evaporated, and heated to volatiliso ammonium salts added in the preceding processes; the residue is dissolved in water and the Cs and Rb are ppd. as double chlorides of Cs, or Rb, and Pt, by addition of 1'tCl,Aq. The pp. is boiled in a very little water, and allowed to settle, the water is poured off while still hot; this process is repeated about 20 times, when the pp. will be quite free from K.PtCl, and will consist of Gs.PtCl, mixed with Rb.PtCl,. The pp. is now reduced in II, CsCl and RbCl are dissolved out in boiling H₂O (Böttger, J. pr. 91, 126). The mixed chlorides are converted into sulphates, these are dissolved in H2O, BaOAq is added, BaSO, is removed by filtration, and the filtrate is evaporated to dryness in a silver dish after addition of (NII), CO₃; the residue is dried, dissolved in water, BaCO₃ removed by filtration, and twice as such H₂C₄H₄O₆ is added as in required to neutralize the solution of Cs. CO. and Rb₂CO₃: the liquid is evaporated at 100° and crystal red; the pp. consists of CsH.C.H.O. mixed w h libH.C.H.O. As the latter salt requires 8 times as much H.O as the former for solution, the two salts may be completely separated by fractional precipitation; this process is continued until the crystals of Cs tartrate do not show a trace of Rb in the spectroscope (Bunsen, P. 119, 1; Allen, P. M. [4] 25, 189). By heating the tartrate, and dissolving the residue in H.SO, Aq, and crystallising, Cs.SO, m by be prepared; this is dissolved in H.O, decomposed by BaOAq, and the solution is filtered and evaporated to dryness in a silver dish, when CsOH is obtained. The CsOH is dissolved in absolute alcohol, and day HCN is passed into this solution : CsC : is thus obtained as a white solid (Setterberg, A. 211, 100). A mixture of 4 parts CsCN and I part Ea(CN), is heated just to melting in a porcelain crueible, and an electrio current from 2 or 3 Bunser cells is passed into the molten mass, in the manner described by Bunson (P. 155, 633). The contents of the crucible are thou warmed under petroleum when the metallic Cs melts into globules (Setterberg, A. 211, 100) .- 2. The mixed chlorides of Cs and Rb, obtained as in 1, are converted into sul658 CÆSIUM.

may be obtained quite free from Rb by a few orystallisations. The Cs alum is dispolved in hot H₂O, and ppd. by NH₂Aq, the liquid is filtered from Al₂O₃, evaporated to dryness in & Pt dish and strongly hoated to remove (NH₄)₂SO₄; the residue is dissolved in H2O, and BlCl,Aq is added so long as a pp. of BaSO, forms; the pp. is filtered off, NH, Aq and (NH,), CO, Aq are added to the filtrate, the liquid is kept warm for some time, and is then filtered from any BaCO, which has formed; the filtrate is eva-perated to dryness, and heated to fusion; solution in H2O, treatment with NH3Aq and (NH4)2CO3, evaporation, and fusion are repeated; finally CsCl is obtained by desolving the fused mass in 1120, and crystallising (Godeffroy, A. 181, 176; Redtenbacher, J. pr. 95, 148). This is converted into Cs.SO, and then into CsOH which is treated as described in 1 .-3. Lepidolite (a silicate of Al), from Hebrou, in Maine, U.S.A., contains about 4 p.c. Cs oxido and 2 p.c. Rb oxide. The powdered mineral is well mixed with 2 parts freshly slaked CaO, and very strongly heated for some time; the mass is powdered, half its weight of conc. H2SO4Aq is added, followed by water; the whole is boiled, filtered, and evaporated to dryness; the residue is dissolved in water, filtered from CaSO, and evaporated until the alums of K, Cs, and Rb crystallise out. About 4 kilos, of the crude mixed alums was prepared by Setterberg (A. 211, 100), and dissolved in hot water, so that the solution had S.G. = 20° Beaumé; this was cooled slowly to 45°, when the Cs and Rb alums ware deposited, as they are insoluble in cold cone, potash alum solution. The alums were dissolved in a little hot water and again cooled, and then solution and orystallisation was continued until the crystals were free from potash. Cs alum is 4 times less soluble in II,O than Rb alum, and is insoluble in a saturated solution . the latter; the mixed alums were dissolved in a little hot water, and allowed to cool, when Cs alum sepa ated with a little Rb alum; this process was repeated until pure Cs alum was obtained. The alum was dissolved in hot water, enough BaOAq added to ppt. Al₂O₃ and all the H₂SO₃, the solution was filtered off and evaporated to dryness: the CsOH thus obtained was dissolved in absolute alcohol, and CsCN was prepared; the CsCN was then electrolysed as described in 1. (For other processes for preparing pure salts of Cs v. Godeffroy, B. 7, 241; Cossa, B. 11, 812; Stolba, D. P. J. 197, 336; 198, 225; Sharples, Am. Ch. 3, 453. For an account of attempts to prepare the metal by various methods similar to those used for preparing Rb, v. Smith, Am. Ch. 6, 106.)"

Properties .- Silver white, soft, ductile, mctal; oxidises rapidly with production of heat and light in air; decomposes ILO at ordinary temperature with inflammation of H produced. Molts 26°-27°; S.G. at 15° 1.88 (Setterberg, A. 211, 100). Spectrum characterised by two lines In the blue, $Cs_a = 4560$, $Cs_b = 4597$; 000005 mgm. Cs may be detected by the spectroscope; 003 CsCl may be detected in presence of 300 400 parts of KCl or NaCl; 001 CsCl in presence of 1500 LiCl (Bunsen, l.c.). The atomic weight of

of V.D. of CsCl, and analyses of the same sals; by Bunsen (P. 113, 353; 119, 4), by Johnson a. Allen (Am. S. 35, 94), and by Godeffroy (A. 181, 185); (2) by comparing the reactions of Cs comporaids with compounds of Li, K, Na, and Rb. One gaseous compound of Cs has been obtained; the S.H. of the metal has not been determined. Cs is positive to all other elements

(v. Alkali Metals).

Combinations.—No compounds of Cs have as yet been prepared directly from the metal. When cone. CsClAq is electrolysed with Pt as the +, and lfg as the - electrode, an amalgam of Cs and Hg is formed, and solidifies to a white crystalline mass; the Cs in this amalgan very

quickly oxidises to CsOII.

Detection and Estimation. Most of the salts of Cs are easily soluble in water. Cs salts may be detected by the comparative insolubility in H₂O of Cs₂PtCl₆ (v. Irreparation, No. 1), and by the spectroscope. There is no satisfactory method for separating and estimating Cs salts; the pp. by PtCl, contains Rb_PtCl, and a little K.PtCla; by repeating the ppn. the pp. may be obtained almost free from K_PtCl_a; the pp. is then reduced in H, the CsCl and RbCl dissolved out, the liquid evaporated and the residue weighed; the Cl is then estimated and the quantity of CsCl is calculated.

Cæsium chloride CsCl. Mol. w. 168.07 (Scott, Pr. E. 1888). For preparation v. Casium, Preparation, No. 2. Small, white, cubes; not deliquescent when pure; partially decomposed by melting in air, residue is alkaline. Melts at low red heat, and volatilises at a higher temperature. Easily soluble in H.O and alcohol. CsCl forms several double compounds, insoluble in conc. IIClAq, with other metallic chlorides; they are obtained by adding CsCl in conc. HClAq to a solution of the other chloride also in cono. HClAq. The following are known: 2CsCl.CdCl_{2*} 2C₃Cl.HgCl., 2CsCl.MnCl., 2CsCl.ZnClg, 2CsCl.CuCl2, 2CsCl.NiCl., $2CsCl.PdCl_{2}$ 6CaCl.BiCl3, 6CsCl.SbCl3, 225; Godeffroy, B. 7, 375; 8, 9); 2CsCl.PtCl. S. (0°) ·024, (100°) ·377. When melten CsCl is electrolysed in an atmosphere free from O, a small blue mass is obtained which is dissolved by H2O with evolution of H; probably this is duo to fc mation of a subchloride.

Cæsium cyanide CsCN. Prepared by the action of dry HCN on CsOH dissolved in absolute alcohol; v. Cyanides.

Cæsium hydrexide CsOH. Mol. w. unknown. Prepared (as described under Casium, Preparation, No. 1) by decomposing Cs.SO.Aq by BaOAq, filtering off BaSO, and evaporating to dryness in a silver dish. Grey-white solid, melting below redness; undecomposed by heat; deliquesces in air, with production of much heat, to form strongly a kaline CsOHAq.

Cæsium exide. An oxido of Cs has not yet been prepared.

Casium, Salts of. Compounds obtained by replacing H of acids by Cs. CsOHAq acts as a very strong base. The salts belong to one 80, CO, &c.; scries CsX whore X = Cl, NO3, Os has been determined (1) by determination the formulæ are established from the vapour

density of CsCl, (Scott, Pr. E. 1888), and also by comparing the salts with those of the other alkali metals. The salts of Cs are very similar to those of Rb; they are well marked, stable, compounds; no basic sales are known; so far as investigation has gone the Cs salts show a marked tendency to form double salts. Most of the salts of Cs are soluble in water; the solutions are ppd. by PtCl,Aq (ycllow), by H₂.C₄H₄O₅Aq (white), by HClO₄Aq (white), and by silicotungstic acid (white). The chief salts are carbonates, nitrate, sclenates, siticotungstate, sulphates, tartrate (v. CARBONATES, Mc.)

M. M. P. M. CAFFEÏC ACID C.H.O. i.c. [4:3:1] C. (OH), CH:CH.CO.H. Di-ornamic acid. Di-oxy-phonyl-acrylic acid. Di-oxy-cin-

Formation. - 1. By boiling onfetamnic acid with aqueous KOH (Hlasiwetz, A. 142, 221).-2. From its acetyl derivative. -3. Powdered cuprea bark is extracted with ether followed by alcohol; the rosiduo is boiled with aqueous KOH, H2SO, is added, and the liquid filtered while hot. The filtrate, when cold, is exhausted with other, and the ethercal solution, after decolourising with animal charcoal, is set aside to crystallise (G. Körner, Ph. [3] 13, 216).--!. From hemlock (in which it is combined with conhydrine?) (Hofmann, B. 17, 1922).

Properties.—Yellow monoclinic tables (containing jaq), v. o. sol. alcohol. The aqueous solution is turned green by Fo.Cl., on adding Na₂CO₃ it then changes to blue and violet. It does not reduce Fehling's solution but reduces warm ammoniacal AgNOs. Its solution in

KOHAq turns brown in air.

Reactions.-1. Dry distillation gives pyrocatechin .- 2. Potash fusion form protocatechuic acid .- 3. Sodium amalgam reduces it to di-oxy-phonyl-propionic acid.

Salts.— G_1A_2' 3aq. S_1A_2' 4aq. $-BaA_2'$ 4aq. $Ba_3(C_9H_3O_4)_29a_4$.— $Pb_3(C_9H_3O_4)_2$ 2aq. Mono-methyl derivative v. Fractic

Di-methyl derivative

O_eH₃(OMe)₂.Cl1;Cl1.CO₂l1. [180]. Formed by saponifying the ether or by heating caffer or ferulic acid with Mcl and KOH. White needles. Sol. alcohol and other, nearly insol. water. On exidation with KMnO, it produces veratric acid. Methyl other A'Me. [64°]. Prisms. Prepared by methylation of isoferulic acid (Tiemann a. Will, B. 11, 651; 14, 959).

Methylenc ether CH. CO. C. Hs. CH. CII. CO. H. [232°]. Formal by beiling piperonal CH2O2:C0H3.CHO with NaOAc and Ac₂O (Lorenz, B. 13, 757). Minute crystals (from diluto alcohol). Conc. H.SO₄ forms a brick-red solution,—AgA'.

Acetul-matheria

Acetyl-methyl derivative v. Acetyl-FERULIO ACID.

Di-acetyl derivative C.H. (OAc) CH:CH.CO.H. [191°]. From caffeic acid and Ac.O or by heating protocatechuic aldehyde (2 pts.) with NaOAo (2 pts.) and Ao₂O (6 pts.). Slonder needles. V. sl. sol. water, v. sol. alcohol and other (Tiemann a. Nagai, B. 11, 659).

Hydro-caffeic acid v. DI-OXY-PHENYL-PRO-PIONIC ACID.

CAFFEIDINE C, H12N,O. Formed, together with methylamine, CO₂, and NH₃, by boiling caf-feine with conc. baryta-water (Strecker, A. 123, 360; 15, 1; C. R. 52, 1269; Schmidt, B. 14, 816; Schultzon, Z. 1867, 616). Alkaline liquid. Sol. wat, r, alcohol, and chloroform, sl. sol. ether. Long boiling with baryta-water gives methylamide-acetic acid, formic acid, CO, and NH, Chromic acid oxidises it to di-methyl manide, methylamine, CO2, and NII, (Maly a. Andreasoh, M. 4, 381) Etl forms C, H, EtN, O.

Salts.-B'HCl.-B',H.PtCl, 4aq. CAFFEIDINE CARBOXYLIC ACID

C_sH₁₂N₄O₂. Prepared by the gradual solution of caffeino in dilut NaOHAq; this solution is neutralised with HOAc and the collect salt ppd. with Cu(OAc), (Mnly a. Andreasch, M. 4, 369). Very soluble crystalline mass; its aqueous solution on boiling gives off CO, and leaves caffeidine. Salts. - KA': golden syrup. 11gA'22HgCl2:

bulky pp. — $\text{Cu}\Delta'_{2}$: minute crystalline granules. — $\text{Ca}\Delta'_{2}$ — $\text{Cu}\Delta'_{2}$. — $\text{Cd}A'_{2}$. — $\text{Mg}\Delta'_{3}$. — $\text{Ca}A'_{4}$. — $\text{Ca}A'_{5}$. — $\text{Cd}A'_{5}$. — $\text{Mg}\Delta'_{5}$. — $\text{Ca}A'_{5}$. maille, C. R. 81, 817). S. (alcohol) 61 at 16 ; 3.12 at 78°. S. (ether) 011 at 16'. S. (CS.) 06

at 16°. S. (chloroform) 13 at 16 '.

Occurrence.-1. In coffee berries and leaves (Runge, Materialien zur Phytologie, 1820; Stenhouse, P. M. [4] 7, 21; Pfaff a. Liebig, A. 1, 17). Coffee berries contain from 1 to 1 28 p.c. caffeine; roasted coffee about 1.3 p.c. (Paul a. Cownley, Ph. [3] 17, 565; cf. Stenhouso a. Campbell, C. J. 9, 33; A. 89, 246).—2. In tea leaves (Oudry, Mag. Pharm. 19 19; Jobst, A. 25, 63; Mulder, P. 43, 160). Tea contains 2 to 1 p.c.-3. In guarana, the dried pulp of Paulinia sorbilis (Martius, A. 36, 93). Guarana contains about 5 p.o. of caffeine.—4. In Maté or Paraguay tea the leaves and twigs of Hex Paragnayensis (Stenhonse, P. M. 3] 23, 426). 5. In the seeds of the Kola free (Col's acaminata) of West Central Africa, to the amount of 2.13 p.c. of the dried seed (Attield, Ph. [2] 6, 457).—6. Present to a small excut in cocoa (E. Schmidt, A. 217, 306).

Formation. By heating silver theobromine with MeI for 20 hours at 160': caffeine is thus shown to be methyf-theobromine (Strecker, A.

118, 151; E. Schmidt, A. 217, 282).

Preparation .- 1. Ten or coffee is exhausted with boiling water; tannin is ppd. by lead subacctate; the filtrate is freed from lead by H.S and eve porated to crystallisation (Péligot, A. Ch. [3] 11, 129). 2. Raw ground coffee (5 pts.) is mixed with moist lime (2 pts.) and extracted with alcohol, chloroform, or benzene, from which the caffeine crystallises or evaporation (Versmann, Ar. Ph. [2] 68 148; Vogel, C. C. 1858, 367; Payer A. Ch. [3] 26, 108; Paul a. Cownley, Ph. [3] 17, 565). -3. Tea or coffee is boiled with water and either the whole, or else the filtrate, is evaporated to a syrup, mixed with alaked lime and extraced with chloroform (Aubert, Pflüger's Archiv, 5, 589; Cazeneuve a. Caillel, Bl. [2] 27, 199).-4. By sublimation from tea (Heignsius, J. pr. 49, 317) .-- 5. A decoction of tea is evaporated with PbO to a syrup, K.CO, is added, and caffeine extracted by alcohol (Grosschoff, J. 1866, 470).

Properties.-Mass of slender silky needles (containing aq); begins to sublime at 79°

(Blyth). Sl. sol. cold water and alcohol, v. sl. sol. ether. The crystals from alcohol and ether are anhydrous. Weak base; the sale being decomposed by water; doss not affect red litmus. Tastes bitter. Produces tetatus and rigor in the voluntary muscles of frogs Anbert; Brunton a. Cash, Pr. 42, 238). In men it increases the heart's action, excites the nervous system, and diminishes metabolism (?) (Lohmann, A. 87, 205). Caffeine gives a yellow pp. with phosphomolybdic acid.

Estimation. - The various methods of preparation may also be used for estimation (Stenhouse, A. 102, 126; Lieventhal, C. C. 1872, 631; Weyrick. Fr. 12, 191; Péligot, Rep. Pharm. 82, 540; Claus, J. 1863, 708, Zöller, J. 1871, 818; Mulder, J. pr. 15, 280; Commaille, Bl. [2] 25, 261; Paul a. Cownley, Ph. [3]

Colour Test .- Evaperate with chlorine-water on platinum-foil. A yellowish residue is left, which on further heating becomes red, and is turned purple by ammonia (Schwarzenbach, J. 1861, 871; 1865, 730). Xanthine, theobromine and uric acid also give this test. evaporated with conc. HNO3 gives a yellow residue (amalic acid) which is also turned purple (murexide) by amnionia (Rochleder, A. 69, 120).

Reactions.-1. Gaseous chloring or HCl and KClO3 give in the first place di-methyl-alloxan and methyl-urea (E. Fischer, A. 215, 257): $C_8H_{16}N_4O_2 + O_2 + 2H_2O = C_6H_8N_2O_3 + C_2H_8N_2O$. Part of the di-methyl-alloxan becomes amalic acid. Chloro-caffeine, methylamine, and eyancgen obloride are also formed, and, if the reaction is prolonged, di-methyl-parabanic acid (cholesis prioringed, it meany regulation and consister optione). Bromine and water at 100° act similarly (Maly a. Hinterberger, M. 3, 85).—2. Cold HNO, attacks it slewly, giving off CO₂ (1 vol.) and N₂O (about 2 vols.) (Franchimont, D. 2002, 2003), 2 Het Alliet HNO gives di R. T. C. 6, 223).—3. Hot diluto IINO, gives dimethyl-parabanic acid (C'enliouse, A. 45, 366; 46, 227; Rochleder, A. 69, 120; 71, 1).—3. Chromic acid gives di methyl-parabaric acid, NH3, methylamine, and CO2 (Maly a. Hinterberger, M. 2, 87).—4. Boiling baryta water splits up caffcine into caffcidine and CO2; the caffcidine then breaks up into CO., NH., methylamine, formic acid, and methyl-amido-acetic acid (sareosine) (Rosengarten a. Strecker, A. 157, 1). 5. With conc. HCl at 250° it forms ammonia, methylamine, sarcosine, formic acid and CO, (E. Schmidt, A. 217, 270). The volume of NH, is to that of NMeH, as 1:2 Delow 200°, HCl has no action. Hence there are three NMo groups in oaffeine, $C_8H_{10}N_4O_2 + 6H_2O = 2CO_2 + 2NMeH_2 + NH_3 + CH_2O_2 + C_3H_3NO_2$.

Salts.—(E. Schmids, A. 217, 282; Herzog, A. 26, 344; 29, 171; Bicder, nann, Ar. Ph. [3] 21, 175; Tilden, O. J. 18, 99; 19, 145.) B'HCl.—B'HCl 2aq: monoclinic; decomposed by moist air into HCl and caffeine.—B'2HCl.—B'4HCl.—B'₄H₂, tCl₄ (at 100°).—B'HAuCl, 2aq; glittering plates.—B'HBr 2aq.—B'HI.—B'2HI.—B'2HI.—B'HI.4aq (Tilden).—B'HCCl1: [175°]; yellow needles converted by NH, into a greenishblack pp. (Tilden, Z. 1866, 350; Ostermayer, B.

18, 2298).—B'HNO, aq.—B'H,SO, aq. Formate B'H,CO,..—Acetate B'2HOAc.—Butyrate B'C,H,O,..—Isovalerate B'C,H,O,. -Citrate: prepared by adding a solution of

citric acid (1 pt.) in alcohol (71 pts.) to one of caffeine (1 pt.) in chloroform (14 pts.) and evaporating. Semi-crystallins powder, decomposed by most solvents (Lloyd, Ph. [8] 11, 760). According to Tanret (J. Ph. [5] 5, 591) the last five salts are merely mixtures.—Caffsate B'C, H,O, 2aq (Hlasiwetz, A. 142, 226).

Combinations.—B'HgCl₂ (Nicholson, A. 62, 78; Hinterberger, A. 82, 316).—B'HgCy₂ (Kohl

a. Swoboda, A. 83, 341).—B'AgNO₃.

Methylo - chloride B'MeCl aq. 200° it splits up into McCl and cafforne.— (B'McCl).PtOl., Sparingly soluble. Methylo-iodide B'McI aq (Tilden, J. pr.

94, 374; E. Schmidt, A. 217, 286; E. Schmitt a. E. Schilling, A. 228, 141). From Carorne and MeI at 130°. At 100° it loses aq, at 190° it splits up inte casseine and MeI. Triclinic: splits up into casteine and MeI. Triclinia: $a:b:c = .6062:1:4161; a = 91°24'; \beta = 74°; \gamma = 88°.$

B'MeI3

Methylo-hydroxide B'MeOHaq [910], and B'MeOH [138']. From the methyloiodide and Ag.O (Sohmitt a. Schilling, A. 228, 143). Crystals. V. sol. water, alcohol, and chloroform. v. sl. sol. ether or light petroloum. Its solutions are neutral. It is not poisonous. Heated at 200° in the dry state it gives off methylamine while caffeine is also formed. HCl or dilute H.SO, convert only part of it into the corresponding salt, the rest gives methylamine, fermic acid and dimethyldialuric acid, the latter being converted by atmospheric oxygen into amalic acid. Whon HCl is used, caffeine methylocliloride is one of the products. With water at 200° it gives sarcosine, methylamine, formio acid, and CO2. Chromic acid forms cho-

lestrophane, methylamine, formic acid and CO₂.

Ethylo-triiodide BEtI₃. From caffeine and EtI at 130° (Tilden, C. J. 18, 99; 19, 145).—

B'2Et2PtCl6.

Chloro-caffeine C_aH₃ClN₄O₂. [188°]. Formed by passing chlorine into dry caffeine in dry CHCl₄. Crystallised from water. V. sl. sol. cold water and ether, v. sol. strong acids but ppd. by water. Reduced to caffeine by zinc-dust and HCl (Fischer, A. 215, 262; 221, 336).

Bromo-caffeine v. p. 561.

Amido-caffeine C₂H₂(NH₂)N₄O₂. [above 360°].

From bromo-caffeine (2 pts.) and alcoholic NH₄
(20 pts.) by heating for 7 hours at 130° (Fischer, A. 215, 265. Slender needles; may be distilled. V. sl. sol. water and alcohol; sol. cono. HOAc; sol. conc. HClAq, but reppd. by water, being apparently less besic than cassoine.

 $0\,\mathrm{xy}$ -caffeine $C_\mathrm{s}H_\mathrm{s}(\mathrm{OH})\mathrm{N_{\bullet}O_{2}}.$ [o. 345°]. From ethoxy caffoine by heating with dilute HOI (Fischer, A. 215, 268). Mass of white needles (from water). V. sl. sol. alcohol, ether, or cold water. Sol. cono. HCl but reppd. by water. Oxy-caffoïne is an aoid.—NaA' 3aq. Needles.— BaA', 3aq. Redctions .- 1., The silver salt with EtI at 100° gives ethoxy-casselne.—2. PCl, in POCl, gives ohloro-casselne.—3. Cl at a high temperature gives di-methyl-allexan.-4. Cl gas at 0° in a solution of oxy-caffeins in HCl gives apo- and hypo-caffeine.—5. Dry bromins forms an addition compound C₆H₉(OH)N₄O₂Br₂(?) as a red mass, decomposed by water or alcohol, the latter giving diethoxy-oxy-caffeine dihydride. Ethoxy - caffeine O.H. (OEt) N.O. [140

From brome-caffeine and alcoholic KOH (Fis-

cher, A. 215, 266). White needles (from water). Sl. sol. cold alcohol or ether, v. e. sol. hot alco-

action of methyl alcohol on oxy-caffeine bromide. Colourless crystals. Sol. water and alcohol. By HCl it is decomposed into methyl alcohol, methylamine and apo-caffeine (Fischer, B. 14,

642).
Di-ethoxy-oxy-caffeine dihydride
C₂H₂(OEt)₂(OH)N₁O₂ Di-ethyl derivative of trioxy caffeine dihydride. [1950-205"]. From oxycaffe ne Br, and alcohol, as above. Warmed with HCl z gives alcohol, methylamine, apocaffeine, and hypo-caffeine. Funning HI or HI gas passed into chloroform solution reduces it to oxy caffeine. With phosphorus oxychloride it forms a crystalline substance that appears to be C_aH_aN₄O₂(OH)(OEt)Cl. This body is reconverted by alcohol into diethoxy-oxy-caffeine dihydride, but it is decemposed by water, one of the products being di-methyl allexan, although this is not formed from diethoxy-oxy-casseine dihydride by water or acids (Fischer a. Reeso, A. 221, 887).

Allo-caffeine C₈II₉N₃O₅. [198°]. A by-product obtained in the preparation of the preceding body from oxy-caffeine, bromine and alcohol, especially when the latter is wet (92 p.c.). Sandy powder. V. sl. sol. water, sl. sol. boiling alcohol. Decomposed by boiling HCl (Fischer,

A. 215, 276)

Apo-eaffeine C, II, N, O, . [148°].

Formation.—1. From di-ethoxy oxy-caffeine dihydride (5 g.) by evaporating with (20 g.) di-lnte (20 p.c.) HCl at 100° (Fischer, A. 215, 277); the equation is: $C_8H_0(OH)N_1O_2(OEt)$, $\pm 2H_2O = O_1H_1N_2O_2 + MeNH_2 + 2HOEt$, $\pm 2H_2O = O_2H_2O_2O_2 + MeNH_2 + 2HOEt$, $\pm 2H_2O_2O_2 + MeNH_2O_2O_2 + MeNH_2O_2 + MeNH_$ caffeine, HCl and KClO, (Maly a. Andreasch M. 3, 100).

Properties. - Monoclinio crystals (from water). a:b:c = 8025:I: 6976. V. sol. hot water. alcohol or chloroform, sl. sol. cold water, benzene or CS... Boiling water decomposes it into CO2, hypo-caffeine and caffuric acid (q. v.).

Hypo-caffeine C₆H,N₃O₃. [182].

Formation.—1. Formed along with apocaffeine by warming the di-ethyl docivative of tri-oxy-caffeine dihydrido with hydrochloric acid, thus: $C_aH_a(OH)N_4O_2(OEt)_2 + 2HO = C_0H_4N_2O_3 + 2HOEt + NH_4MO + CO_2 - 2$. From oxy-caffcino, HCl and Cl (Fischer, A. 215, 288).

Properties. — Crystallised from water. sol. hot water or aloohol, sl. sol. cold water. May be distilled with but slight decomposition.

Ba(O₂H₂N₃O₃)₃C₂H,N₃O₃: v. sol. water.

Reactions.—Not affected by boiling funing
HNO₃, chlorine- or bromine-water, K₂Cr₂O₃ and dilnte H.SO., HMnO., conc. HCl, fuming HI, Sn and HCl, Ac.O or POCl, and PCl.. Water at 150° completely destroys it. Boiled with baryta

it gives oaffolin (q. v.).

Caffolin C,H,N,O₂. [194°-190°]. Formed by boiling hypo-cafforne with lead sub acetate (Fischer, A. 215, 292). Slender needles (from alcohol) or long prisms (from warm water). V. e. sol. warm water. Sl. sol. alcohol. Does not combine with acids. It is but a feeble acid, for its barium compound is decomposed by CO₂. Foiled with Ag₂O, it forms a crystalline silver of mpound.

Realtions.—1. Cono. H'll at 100° splits it up into CO, NH₃, NMeH₄, &c.—2. Cono. HI forms methylk' area.—3. K₃FeCy₆ gives mothyl-oxamio acid and methyl-u-ea: C₃H₄N₁O₂+O+H₄O=MeNH.CO.CO₂H + MeNH.CO.NH₂.—4. KMnO₄ and KOH give di-methyl-oxamide and ammonia according to the reaction. C.H. N. A. A. H. O. according to the reaction: C, H, N,O2+O+H2O = MeNH.CO.CO.NIIMe + CO_2 + NII_3 . -5. Potassium bichromate and H.SO, give cholestrophane: $C_sH_gN_gO_s + O = C_sH_gN_gO_g + NH_g$, -6. Nitrons acid com ctely destroys it. -7. Boiled with Ac.O it forms the acetyl derivative of acecassiu Call 10 AcN 3()2.

Acconfin CoH₁₁N₃O₂. [110°-112°]. From its acetyl derivative by evaporating with fuming HCl at 100 and decomposing the resulting hydrochlorido by Ag.O (Fischer, A. 215, 300). Trimetric crystals (from benzene). a:b:c=6707:1:1-2445. May be distilled undecomposed.

V. e. sol. water and alcohol.

Acetyl derivative C_eH_{1e}AeN₂O₂... [106°-107°]. From caffolin by boiling with Ae₂O as long as CO₂ comes off (12 hours). Mouoclinio tables (from chloroform mixed with ether).

Caffurio acid CaHaNaO4. [2100-2200]. From apo-caffeine by boiling water (Fischer, A. 215, 280). C₇H₇N₂O₅ + H₂O = C₆H₈N₂O₅ + CO₅. Transparent tables (from alcohol). V. sol. water, sl. sol. cold alcohol, chloroform or ether. Feeble acid, its barium salt being decomposed by CO. Salt. - AgA'. Tables, sl. sol. water.

Reactions.-1. Not affected by chlorine- or bromine-water .- 2. III converts it into hydrocaffuric acid. -3. Warmed with land sub-acetate it gives mesoxalic acid, methyl-urea and methylamine. -4. II A KOH gives off NH Me.

Hydro-caturic acid C₆H₅N₃O₃. [210°-248°]. From catturic acid, uning III and PH₄I (Fischer, A. 215, 285). Colourless prisms (from water). V. sol. hot water, sl. sol. cold water.

Reactions. -1. Gives no pp. when boiled with lead sub-acetate (unlike caffuric acid) .- 2. Gives with ammoniacal AgNO, a mirror in the cold. 4. Chlorine-water oxidises it to caffuric acid. -3. Hot KOH gives off methylamine.-5. Warmed with baryta it forms methylamine and methylhydantoin carboxylic acid, the latter splitting up into CO and methyl-hydantoin.

Metayl caf aric acid C.H., N.O. [167°]. From allocaffeine by boiling with water (Schmidt a. Schilling, A. 228, 172). Needles (from water). V. sol. water, alcohol and chloroform. Basio lead acctate converts it into mesoxalio acid, methylamine and dimetnyl urea.

Amalic acid v. n. 149. Coastitution of Caffeine .- Medicus (A. 175.

MeN = CO250) proposed the formula -NMe while Emil Fischer (A. 215, 314) proposed

CO C.NCH. . Both formulæ readily re->00

Me.N-CH

present the formation by oxidation of dismethylalloxan and methyl urea. According to lischer's formula the derivatives of casseine would be re-presented as follows: Oxy-casseine would be NMe—C(OH) = C – NMe >CO. Ethoxy-oxycaffeine dihydride would bo NMe - C(OH)(OEt) - C(OEt).NMe>CO. Apocaffoine would be Caffurio acid would be HO.C(CO₂H) – NMe SCO Hydro-caffurio acid might be written: HC(CO₂H)-NMe >00. Methyl-hydantoïn earboxylic acid would bo IIC(CO₂H) . NMe _____>CO; whence methyl-hydantoin Hypo-caffeinc may bo >CO, se that apo-caffeine would be its earboxylic acid. Caffolin may then bo | CO; hut the formation of cholestro-HO.CH.NMe CO — NMe | rrom it is in that ease NMo—CO somewhat anomalous. Streckor's caffeïdine (from caffeïno by alkalis)
McHN.CII = C - NMe MeIINC N would be boiling alkalis gives CO2, NII3, 2NMell2, formic acid and sarcosine. • Theobromine will be • CO C. NMe | | >CO | ItN -C = N HN-CII C-NMe as is shown by the formation of hypo-ethyl-theobromine. Somewhat similar formulæ are arrived at if we start from the formula of Medicus. Inasmuck asoaffein methylo-hydroxide differs from caffeine in giving no NH, but only NMeH, in its decom-

positions, we must assume that it has the formula

со о—име —

McN-CO

MeN-CH CO Ö MeN -C-NMe(OH) Its decomposition-product, di-methyl-dialuric acid, should, according to Maly a. Hinterberger (M. 3, 85), be represented by the formula McN-CO CO CH(OH), which agrees better with the for-MeN—CO mula of Medicus. On the other hand, the frequent occurrence of mothyl-urea as well as a di-methyl uren among the decomposition-products of caffeine and its derivatives accords best with Fischer's formula (Schmidt a. Schilling, A. 228, 174). CAFFEOL C₂H₁₀O₂. (196°). Givon off (to the extent of '05 p.o.) on roasting coilee together with caffeine ('18 p.c.), palmitic acid, aoctio acid, CO₂, and traces pyrrol, methylamine, and hydroquinone. It is extracted by other from the liquid distillate (Bernheimer, M. 1, 459). Liquid, smelling like coffec, sl. sol. hot water, v. c. sol. alcohol and ether; v. sl. sol. conc. KOHAq. Fe₂Cl₆ colours its alcoholic solution red. Potash-fusion gives salicylic acid. It is perhaps a methyl derivative of o-oxy-benzyl alcohol. CAFFETANNIC ACID C, H18O8. Occurs in coffee berries to the amount of 3 to 5 p.c. as Ca and Mg salt, and perhaps also as a double salt of K and caffeine (Pfaff, (1830) Scher. 61, 487; Rochleder, A. 59, 300; 63, 193; 66, 35; 82, 196; Liebich, A. 71, 57; Payen, A. Ch. [3] 26, 108). Prepared by mixing an alcoholic infusion of collec with water; liltering from ppd. fatty matter; boiling the filtrate, and ppg. as lead salt by Pv(OAc). Colourless mammellated crystalline groups, v. sol. water, m. sol. alcohol; has an astringent taste; strongly reddens litnus. P.c.Cl, colours its solutions green. It does not ppt. forrous salts, tartar-emetic, or gelatin, but it ppts. quinine and oinohonine. It reduces AgNO3Aq, forming a mirror. Its salts turn green in air. Potash-fusion gives protocatechuic acid. Boiling conc. KOHAq splits it up into caffeio acid and a sugar (Hlasiwetz, A. 142, 220). Salts:-BaA', (at 100°): amorphous, v. sol. water; addition of baryta forms a yellow pp. -PbC₁₅H₁₆O₈.-Pb₂(C₁₅H₁₅O₈)₂.-Pb₂C₁₆H₁₄O₈ (at 100°). Viridic acid. An acid formed by the atmbspheric oxidation of an ammoniacal solution of caffetannio acid. According to Rochleder the green colour of coffee berries is due to calcium viridate. It is ppd. by Pb(OAc)₂. Amorphous brown mass, v. sol. water. Cono. H₂SO₄ forms a crimson solution whence water gives a flocu-

CAFFULIN v. CAFFEINE.
CAFFURIC ACID v. CAFFEINE.
CAIL-CEDRIN. A bitter, neutral, resinous substance present to a minute extent in the bark of the Caul-cedra (Caventou, J. Ph. [8] 16, 855; 83, 123).

1858, 261).

lent blue pp. The aqueous solutions are turned green by alkalis, and give a bluish-green pp. with baryta-water (cf. Ylaanderen a. Mulder, J.

Found in Cainoa root (from Chiococca anguifuga and racemosa) (François, Pelletier, a. Caventou, J. Ph. 16, 465; Liebig, A. Ch. [2] 47, 185; Rochleder a. Hlasiwetz, A. 76, 338; Rochleder, J. pr. 85, 275). The root is exhausted with alcohol and the oaincin ppd. either by milk of limo or Pb(QAo)2. Crystallino flakes, tasteless at first, afterwards very bitter; v. sl. sol. water and ether, v. sol. alcohol; reddens litmus. Boiling alcoholio IICl splits it up into a sugar (C₆H₁₂O₄) and crystalline carnoctin C₂₂H₃₁O₃. Carnoctin is resolved by petash-fusion into butyrio acid and caïne igen in $C_{11}H_{21}O_{2}$ which is possibly related to asseigen in. Camein in dilute alcoholic solution is converted by sodiumamalgam in crystalline $C_{36}H_{58}O_{15}$ whence

fuming HCl forms gelatinous C₁₈H₂₈O₂.

CAJEPUT, OIL OF. A light green oil propared in India by distilling the leaves of Melaleuca leucodendron with water. Its chief constituent is cincol $C_{10}H_{10}O$ (q.v.), which is also called oajoputol. $P_{20}O_{3}$ converts it into terpones (q.v.) which when so obtained may be callod oajéputenes (Schmidt, C. J. 14, 63; Wright a. Lambert, C. J. 27, 619; Histed, Ph. [3] 2, 804; Blanchet, A. 7, 161; Gladstone, C. J. 49, 621).

CALAMUS ROOT. According to Genther (A. 240, 92) the acerin prepared by Thoms (p. 60) from Acorus calamus is not a definite substance, but is separated by alkalis into a neutral amorpheus brown mass (C10 Ha, NO, ?) and an acid (C₁₁II₁₈O₃?). Calamus root after extraction with water still contains a combined acid (C11H18O4?) which may be extracted by adding HCl and shaking with other. When the root is distilled with steam, in thyl alcohol and a mixture of terpenes (q, v) and a compound $C_{10}H_{10}O(?)$ is obtained $(G_{\cdot}; Sokpedermann, A.$ 41, 374; Kurbetow, B. 6, 1210; Gladstone, C. J.

CALCIUM. Ca. At. w. 39.91. Mol. w. nnk. i. wn. Molts at red heat. S.G. 1.57 (Matthiessen, A. 93, 27). S.H. (0'-100') .1686 (Bunsen, P. 141, 1). S.V.S. abt. 25. E.C. (Hg at 0'=1) 12.5 (Matthicssen, P. M. [4] 12, 199; 13, 81). Chief lines in emission-spectrum, 6121-2, 5587-6, 4226.3, 3968, 3932.8.

Occurrence.-Never free. Very widely distributed, and often in largo quantities, as silicato, phosphate, sulphato, caroonate, fluoride, &c. Most natural waters contain Ca salts; phosphato and carbonate of Ca are found in plants and animals. Ca salts occur in the sun, fixed stars, and meteorites. Calcium carbonate and burnt lime have been known from very anciont times. In 1722 Fr. 11offmann showed that lime is a distinct earth; Black (1755) was the first to make a quantitative examination of limestone and burnt lime. In 1808 Davy obtained calcium (impure) by the electrolysis of lime.

Formation.—1. Dry CaI, is heated with Na in an iron orucible with an air-tight cover (Liés-Bodart a. Jobin, A. Ch. [3] 54, 363; Dumas, C. R. 47, 575; Sonstadt, C. N. 9, 140).— 2. Dry fused CaCl (300 parts), Na (100 parts), and pure distilled granulated Zn (400 parts), are heated in a orucible with loosely fitting lid,

CAINCIN C. H. O. Carneic acid. S. 14. possible without volatilisation of much Zn; an according to v. Rath, P. 136, 434). This alloy is heated in a orusible of gas coke until the Zn is a distilled off (Carm, C. R. 48, 450; 50, 547). is electro, ysed, using an amalgamated Pt wire as

negative electrode (Bensen, A. 92, 218).

Preparation.—A mixture of dry CaCl₂ and SrCl2, in the ratio 2CaCl2:SrCl2, mixed with a little NH,Cl, is melted in an open crucible; the current from 3 or 4 Bunson cells is passed through the melten mass, the positive electrode being a stick of earbon, and the negative ariron wire as thick as a knitting needle, drawn out to a fine point. The point of the iron wire is kept just under the surface of the molten mass for a minute or so at a time; the Ca separates in small lumps (Matthiessen, A. 93, 277; 94, 108). Frey obtained lumps of Ca weighing from $2\frac{1}{4}$ to 4 grams (A. 183, 367); he passed the negative electrodo through tho stem of a tobacco pipe with the bewl dipping under the molten mass in the crucible; II was then passed into the pipe; when the pipe and bewl were filled with this gas, the H was stepped, and the current was started; the Ca rese into the bowl of the pipe, and being in centact with II remained quite unoxidised.

Properties .- Lustrous, clear yellowish white, very ductile, but brittle when hammerod out, maller sle; about as hard as calespar. Frey (A. 183, 367) says it is brittle and cannot be hammered out or drawn into wire. Melts at full red heat, and then burns with yellow flame and production of much heat and light; [Ca, O] = 130,930 (Th. 3, 251). Does not oxidise in dry air; but in ordinary air is quickly covered with CaO. Not velatilised at temperature of inflammation (Caron, C. R. 48, 410). Decomposes cold II O rapully;

[Ca, Λq] [Ca, Ω , H^2 , Λq] $-2[H^2, O] = 80,900$ (Th. 3. 251). As no compound of Ca has been gasified, the value to be given to the atomic weight of the metal is decided partly by the S.H. and partly by purely chemical considerations. The mass of Ca that combines with 15.96 (i.e. with 1 atom) O is 39.91, hence the simplest formula for the exide is CaO (Ca = 39.91); the same mass of Ca combines with 2×35.37 Cl, 2×79.75 Br, &c.; the simplest formular for the chloride and bremide are thereforo (iCL and CaBr₂ respectively (Ca = 89.91). These forms a are in keeping with the reactions of the compounds, hence they are adopted. The chief compounds of Ca by analyses of which the value Ca 32 91 has been found are: (1) CaCl₂ (Berzelius, G. A. 57, 451; Dumas, A. Ch. [3] 55, 1901; (2) CaCO, converted into CaO (Dumas, C. R. 14, 537; Erdmann a. Marchand, J. pr. 26, 472). Ca is a strongly positive metal, forming well-marked and stable salts by replacing the light of acids. Salts of Ca derived from almost every acid are known; several of these form double salts; very few basic salts are known. CaO₂H₂ is an alkaline hydroxide; CaO combines with H₂O with produotion of much heat; CaO2H2 is dehydrated to CaO at a high temperature. [CaO, H'O] = 15,540 (Th. 3,251). The heat of neutralisation as high a temperature being maintained as is of CaO,H2Aq is the same as that of KOHAq,

NaOHAq, and BaO,H,Aq, viz. 31,150 for H,SO,Aq and 27,640 for H,Ol,Aq. (is combines with the halogons with production of heat; [Ca,X²] = 169,820 when X = Cl; 140,850 pure HClAq; the solution is evaporated to dryness and Combinations.—1. Wijh water

Reactions and Combinations.—1. Wijh water

forms CaOAq and H .- 2. With acids forms salts, usually evolving H; cone. HNO, Aq only acts at high temperatures.—3. Combines directly, when heated, with many non-metals; especially Cl, Br, I, O, S, P (v. Calcium chlorine, &c.). 4. Forms alloys with soveral metals, by heating the metals together. Alloys with Al, Sb, Pb, Hg, Na, and Zn have been described (Caron, C. R. 48, 440; 50, 547; Wöhler, A. 138, 253). Calcium is usually estimated either as carbonato or sulphate for volumetrically, by K2Mn2O8Aq, after ppn. as CaC₂O₄ and decomposition of this ealt by H2SO,Aq.

Calcium, Alloys of, v. Calcium; Combinations, No. 4.

Calcium, Arsenates of. CaHAsO, and under ARSENIC, Ca₂(AsO₄)₂: v. ARSENATES, ACIDS OF.

Calcium, Arsenites of. Ca₂(AsO₂)₂; Ca(AsO₂)₂; and Ca, As, Os: v. ARSENITES, under ARSENIO, ACIDS OF.

Calcium, Bromide of. CaBr. Mol. w. unknown, as compound has not been gasified. (CaBr²,Aq] = 165,360 (Thomsen).

Formation.-CaBr, is formed by dissolving CaO or CaCOs in HBrAq, evaporating, and orystallising; or by decomposing Fe, Br, Aq by

CaOAq.

Preparation .- 1. 121 parts Br and 1 part amorphous P are allowed to react in presence of H2O; the solution is neutralised by CaCO. or CaO,H., filtered from Ca, 2PO,, evaporated, and crystallised (Klein, A. 128, 237).-2. 20 parts S are dissolved in 210 parts Br, and the liquid is poured into thin milk of lime, centaining 140 parts CaO; CaSO, is ppd. by alcohol, the solution is filtered off, evaporated, and crystallised (Faust, Ar. Ph. [2] 131, 216).

Properties .- White, lustrous, deliqueseent, needles: very soluble in H.O and alcohol. Absorbs NH, forming CaBr., 6NH, (Rannnelsberg, P. 55, 239). CaBr,Aq boiled with CaO.H., and filtered, on cooling jields, crystals of CaBr₂.3CaO.15H₂O.

Combinations. . - With water to form

 $CaBr_26H_2O$. $[CaBr^2,6H^2O] = 25,600$ $[CaBr^2.6H^2O,Aq] = -1690 (Th. 3, 251).$

Calcium Bromide, hydrated, v. CALCIUM, BROMIDE OF, Combinations.

 Cl^2,Aq = 187,230 (Thomsen).

Formation. - In making NH, by the action

into the acid liquid until all Fe and Mn salts are completely oxidised. Milk of lime is added to alkaline reaction, the whole is digested, the liquid is filtered from lime and ppd., oxides of Mg, Fo, and Mn, nentralised by HClAq, aud evaporated as in 1.

Properties and Reactions .- A white, porous vory deliquescent, solid: after melting and cooling it is distinctly crystalline. Absorbs moisture rapidly: hence is much used for drying gases, &c.; if the CaCl, to be used must be free from CaO, e.g. for drying CO2, it should be placed for some time in a stream of CO, and then of dry air at the ordinary temperature. CaCl, is very soluble in water and alcohol, much less soluble in HClAq. It is partly decomposed by heating in air (v. Weber, B. 15, 2316), more completely by heating in O, with production of CaO. Heated with KClO, or KClO, part of it is changed to CaO (Schulze, J. pr. [2] 21, 407). CaOl, Aq is used as a bath for maintaining temperatures above 100°; 50 parts CaCl. in 100 parts H₂O forms a solution boiling at 112°; 100 CaCl. in 100 water, B.P. 128°; 200 CaCl. in 100 H₂O, B.P. 158°; and 325 CaCl. in 100 1H₂O, B.P. 160° (Magnus, P. 112, 408; Willner, P. 110, 564°; Legrand, A. 17, 34).

Combinations.-1. With water to hexagonal crystals of CaCl_x6H_xO [CaCl², 6H²O] = 21,750 (Th. 3,251); best prepared by evaporating a solution of CaCO, or CaO, in HClAq and crystallising. CaCl_6H_O melts at 28° (Tilden, C. J. 45, 268); heated to 200°, or placed in vacuo, the livdrate CaCl 2H2O remains. This hydrate is also produced, according to Ditte (C. R. 92, 212), by saturating HClAq with MaCl. at 12' and cooling. Hamerl (Sitz. W. (2nd part) 72, 667) says that CaCl₂.4II₂O is formed by repeatedly melting and cooling CaCl2.6112O. According to Dibbits (Ar. N. 13, 478) CaCl26II2O loses 4H2O in a current of dry air, and 6H,O in dry air at 80°. S.G. 12° of CaCl_6H_O, 1.612 (Kopp, A. 93, 120). S.H. of CaCl_6H_O (-20° to 2°) 345, (4°-28°) 647; melted (34°-59°) 5601, (34°-99°) 552, (100°-127°) 519 (Person, C. R. 23, 162). C.E. (cubical) for solid CaCl, 6II,0; Vi=Vo $(1 + 000 645 1t - 000 053 77t^2 + 000 001 906 t^2)$ for interval $11^{\circ} - 26^{\circ}$ (Kopp, A. 93, 129). H.F. $[Ca, Cl', SHO] = 191, 980; CaCl_2.6H_2O dissolves$ in tvater with disappearance of much heat $[CaCl^{2}.6H^{2}O,Aq] = -4.340$ (Thomsen). This salt mixed with snow produces great lowering of temperature; for use as a freezing mixture the Calcium, Chlorido of. CaCl. Mol. w. unknown, as compound has not been gasified.

[719°-723°] (Carnelley, C. J. 29, 497). S.(÷. 2°
2206 (Schiff, A. 108, 23). S.H. (23°-99°) 1612; CaCl.,6H,O is thus obtained as a fine dry (Regnault, A. Ch. [3] 1, 129). S. (0°) 49·6; (20°) 74; (30°) 93; (35°) 104; (40°) 110; (50°) 120; (C0°) 129; (70°) 136; (80°) 110; (50°) 120; (20°) 147; (95°) 151; (99°) 154; (Mulder, J. 1866. 66). H.F. [Ca,Cl²]=169,820; (CaCl.,6H,O: 8·45 H,O (as snow). CaCl.,6H,O; S(Cl.,An) = 187,230 (Thomsen). S (0°) 72.8; (13.8°) 80.9; (24.5°) 89.5; (29.5°) 100 (Hamerl, Sitz. W. (2nd part) 72, 287).—2. With

ammonia forms CaCl, 8NH; dissociated by heat into CaCl, and AH; NH, also removed by dissolving in H,O and passing in a ourrent of air Weber, B. 15, 2316). Isambert (C. R. 66, 1259) (Weber, B. 16, 2316). Isambert (C. R. 66, 1259) describes CaCl₂.4NH₃ and CaCl₂.2NH₃; he gives these thermal values $\frac{1}{2}$ [CaCl², 2NH³] = 14,000; $\frac{1}{2}$ [CaCl², 4NH³] = 12,200; $\frac{1}{2}$ [CaCl², 8NH³] = 11,000 (C. R. 86, 968).—8. With alcohol to form CaCl₂.2C₂H₂O; decomposed by H₂O (Chodnew, A. 71, 241; Johnson, J. pr. 62, 264). Forms combinations also with acctone (Illasiwetz, A. 76, 294).—4. With lime to form CaCl₂.3CaO, 15H₂O; prepared by boiling CaCl₂Aq with CaO, H₂, filtering while hot, and allowing to with CaO2H2, filtering while hot, and allowing to root. decomposed by H.O or C.H.O (Beesley, Ph. 9, 56%; Rose, S. 29, 155; Bolley, D. P. J. 153, 202; Crimshaw, C. N. 30, 280).—5. With platinous chloride to form CaCl. PtCl. 8H.O; M.P. = 100° (Nilson, J. pr. [2] 15, 260).

Calcium chloride, hydrated, v. Calcium, CHLORIDE OF; Combinations, No. 1.

Calcium, Cyanide of. Ca(CN)2. Said to be obtained by heating Ca ferrooyanido and dissolving out with water (Schulz, J. pr. 68, 257). v. CYANIDES.

Calcium, Fluoride of. CaF₂. Mol. w. unknown, as compound has not been gasified. [abt. 902°] (Carnelley, C. J. 33, 280). S.G. abt. 150 (Schröder, Dichtigkeitsmessungen (Heidelberg, 1873); Kengott, Sitz. W. 10, 295). S.H. (21°-50°) '209 (Kopp, T. 155, 71); (15° 99°) '2154 (Regnault, A. Ch. [3] 1, 129). Index of refraction at 21° for line B = 1·432; line D = 1.4339; line F = 1.43709.; line G = 1.43982; line H = 1.44204 (Stefan, Sitz. W. 63 (2nd part), 239). S. (15°) 0004 (Wilson, J. 1850, 278). [CaH²O²,2HF] = 66,600 (Guntz, G. R. 97, 1483, 1558; 98, 816).

Occurrence. - As Fluorspan in octahedra, cubes, and other forms of the monometric system, fairly widely distributed in many rocks; in small quantities in many mineral waters, plantash, bones (Lassaigne, S. 52, 141), enamel of teeth, &c.

Preparation .-- 1. As a golatinous mass, by decomposing an aqueous solution of a Ca salt by that of a fluoride. -2. As a granular powder by digesting freshly ppd. CaCO, with HFAq. -3.
In small octahedra by digesting the gelatinous pp. obtained in 1 with dilute HClAq for I0 hours at 240° (Sénarmont, A. Ch. [3] 32, 123; Schieerer

a. Drechsel, J. pr. [2] 7, 63).

Properties and Reactions. — Transparent, colourless crystals, melting without decomposition at about 900°. Forms easily fusible mass with BaSO, SrSO, and many other insoluble compounds; hence much used as a flux. Soluble in aqueous solutions of NH, salts (Rose, P. 79, 112). Not decomposed by fusion with KOH or NaOH, but partially by fusion with excess of alkali carbonates. Docomposed, to also decomposed by hot H₂SO₄Aq, but only very partially by boiling HClAq or HNO₄Aq. Said to be partly decomposed by Al. 3SO, Aq (Friedel, Bl. [2] 21, 241).

Combinations.—With hydrofluoric acid and water to form CaF2.2HF.CH2O; produced in and TFAq by hot water (Fremy, A. Ch. [8] 47

Calcium, Hydrate of, CaO2H2, v. CALCIUM HYDROLIDE OF.

Calcium, Hydrosulphide of, v. Calcium sulphide prate.

Calcium, Hydroxide of, CaO.H. (Slaked lime.) Mol. w. unknown: compound is decomposed by heat. S.G. 2.078 (Filhol, A. Ch. [3] 21, 415). S.G. (crystalline) 2.236 (Lamy, A. Ch. [5] 14, 145). S. (15°) 13; (54°) 103; (100°) 08 (Dalton, New System, 2, 331); S. (18°) 12. (100°) 47 (Ringar, C.B. 41, 509, p. also 13; (100°) C7 (Bineau, C. R. 41, 500, v. also Lamy, C. R. 36, 333). H.F. [Ca,O,H²O] = 146,470; [CaO,H²O] = 15,540 (T. 3, 251).

Preparation. -1. By adding to 1 part II.O 31 parts CaO. -2. By allowing Ca to oxidise in moist air. 3. By adding KOHAq, or NaOHAq, to a cono. aqueous solution of a Ca salt, collecting pp., washing well, and drying at 100°. Gay-Lussac (A. Ch. 1, 334) obtained CaO2H2 in small six-sided plates by evaporating an aqueous

solution over H.SO, in vacuo.

Properties and Reactions.—A white, compact mass; slightly soluble in cold, less soluble in hot, water [CaO'II',Aq] $\approx 2,290$ (Th. 3, 251). Stronglyalkaline reaction. CaO.H.Aquentralises acids with production of same quantity of heat as when 2NaOHAq, or 2KOHAq is used, viz. about 31,000 for H.SO,Aq, and about 27,900 for 2IIC.Aq (Thomsen); also pps. many heavy metals as oxides or hydroxides, and saponifies fats. Moist Call O. absorbs CO2, forming CaCO3 and H₂O. Cali, O. Aq forms insoluble salts when neutralised by H₂SiO₃Aq, H₃BO₃Aq, H₃PO₄Aq, &c. ; pps. are also formed by adding animal char, sand, &c. Call O2 is soluble in solutions

[4] 6, 203; Péligot, A. Ch. [3] 54, 383; Déon, Bl. [2 17, 155; Berthelot, A. Ch. [3] 46, 173). Cah O, is much more soluble in glyceriu than in water. At a bright red heat CaH,O, is decomposed to CaO+H.O. For reaction between Cl and CallyO2 v. Bleaching Powder under Hypo-ORLORITES, under Chlosine, expactes of.

Calcium, Iedide of. Cal. Mol. w. unknown, as compound has not been gasified. [631° (Carnelley, C J. 33, 279). S. (0°) 192; (20°) 204; (40°) 228; (43°) 286; (92°) 455 (Kremers, P. 103, 65). H. F. (C. 12, 107.55) H. F. [Ca, I2] = 107,250; 103, 65). [Ca, I^2 , Aq] = 134,940 (7h. 3, 251).

Formation .- By the action of IIIAq on CaO2H2; or of I on CaS suspended in water (Liès-Bodart a. Jobin, A. Ch. [3] 54, 363).

Preparation. -To 1 part amorphous P and 40 parts H.O. 20 parts I are slowly added the whole is digested at 100°; the colourless liquid CaO and HF, by heating to cednees in steam; is neutralised by milk of lime, and evapotraed in an atmosphere free from CO, (Liebig, A. 121, 222; Wagner, C. C. 1862, 143).

Properties and Reactions .- White, deliques. cent mass; very soluble in water and alcohol; undecomposed when melted out of contact with water to form CaF, 2HF. O; produced in air: melted in air gives CaO and I. Consmall crystals by evaporating a solution of CaO CaL, Aq dissolves I; on evaporation in vacuo in large excess of HFAq; decomposed to CaF, crystals of a periodide are said to be obtained. Absorbs 6NH, (Isambert, C. A. 66, 1259). Horms | HNO,Aq; collecting pp., washing well with cold an easily decomposed double compound with AgI; CaL, 2AgI.6H, O (Simpson, Pr. 27, 129) | water, and heating the CaO, 8H, thus produced in a current of dry air free from CO, to 100° -

Calcium hydroxyhydrosulphide v. post (inder Calcium sulphydrate.

Calcinm, Oxides of. Two oxides are known; CaO a strongly basic compound, and CaO, which acts as a peroxide. CaO, cannot be formed by

the action of O on CaO (comp. BaO₂).

I. Calcium monoxide. CaO (Lime, burnt lime). Mol. w. unknown, as compound has not been gasified. S.G. 3.15 (Schröder, P. Jubelbd. 452); S.G. (crystalline, by heating Ca2NO₃) 3.251 (Brügelmann, W. 2, 466; 4, 277). S. variablo according to state of aggregation of the CaO &c. Lamy (A. Ch - 5] 14, 145) gives the following numbers representing grams of CaO in 1000 grame of solution; CaO being made (1) by heating Ca2NO₃, (2) by heating CaCO₃, (3) by heating CaO₂II₂:—

Temp. 1·430 1.362 1.381 1.342 1.384 10 1.311 15 1.277 1.299 1.348 1.162 1.195 30 1.142 45 0.996 1.005 1.033 60 0.844 0.868 0.885 0.576 0.584100 0.562H. F. [Ca, O] = 130,930; [Ca, O, Aq] = 149,260

(Th. 3, 251).

Preparation .- Pure marble, or Iceland spar, ie strongly heated in a crucible with a hold in the bottom to allow escape of CO2; or a piece of charcoal is placed in the crucible beneath the marble, CO is thus formed and sweeps out tho CO2 with it. CaCO3 is not completely decomposed when heated in an atmosphere of CO2; v. CALCIUM OARBONATE, undor CARBONATES. Sestini (Fr. 4, 51) strongly heats powdered marble with eugar, washes with H.O. dissolves in HNO₃Aq, pps. CaCO₃ by (NH₄)₂ CO₃Aq, and etrongly heats the dried pp. By strongly heating Ca2NO₄, in quantities of about 15-20 grams at a timo, in a porcelain flask, Brügelmann (W. 2, 466; 4, 277) obtained cubical crystals of CaO; semitransparont, harder than the amorphous form, and less easily acted on by H.O and CO...

Properties and Reactions .- White, amorphous (or crystalline v. supra), powder: does not fuse at full white heat. Strongly basic; reacts with most soids to form salts. CaO is decomposed by heating to whiteness with K; heated in Cl, CaCl₂ is formed. CaS is produced by heating with S, and CaS and CaCO₃ by heating in CS... Ca6 does not combine with O (v. Courcy, C. J. [2] 11, 809).

Combinations .- With carbon dioxide to form CaCO, (but dry CaO doos not react with CO,: Scheibler, B. 19, 1973); combination bogins at about 400° (v. Birnbaum a. Mahn, B. 12, 1547); [CaO, CO²] = 42,520 (Th. 3, 251). Heated with silica or eilioates, silicates of Ca aro formed, which in contact with water set to a hard compact maes (hydraulic mortars). With water, CaO.II, is formed with production of much heat [CaO, \tilde{H}^2O] = 15,540 (Thomson); the lime is said to be slaked.

II. CALCIUM DIOXIDE, CaO, (Calcium perdide). Mol. w. unknown. Prepared by adding pure H.O.Aq to excess of CaOAq, or by adding excees of CaOAq to Na₂O₂Aq containing some exception of CaF₂ all the salts are more or less

120°. Forme a enow-white cryetalline powder; doos not melt at red heat, but gives off O and forms CaO. The hydrate CaO, 8H,O is slightly soluble in H2O, in contact with H2O it slowly decomposes to CaO₂H₂Aq and H; soluble in NH₄ClAq, but not in NH₃Aq; dissolves easily in dilute acids, oven in H.C₂H₃O₂Aq, without evolution of O. It forms prismatio dimetric crystals, isomorphous with BaO₂.8H₂O and SrO₂.8H₂O (Schono) (Thénard, A. Ch. [2] 8, 306; Conroy, C. J. [2] 11, 808; Sohöno, B. 6, 1172).

Calcium oxide, hydrated, CaO.H., v. Cit.

CIUM, HYDROXIDE OF.

Calcinm, Oxybromide of, CaBr2.3CaO.15H2O v. Calcium, Bromide of; Properties.

Calcium, Oxychloride of, CaCl...3CaO.15II.O, v. Calcium, chiloride of ; Combinations, No. 4. Calcium, Oxysulphides of, v. Calcium Poly-

SULPHIDES; under CALCIUM, SULPHIDES OF.
Calcium, Phosphide of. When Ca and P are
heated under rock oil, and the unacted-on P is dissolved out by CS2, a black powder remains which is acted on by H₂O and acids with production of PHa; this black powder is said by Vigier to be Ca phosphide (Bl. 1861. 5). By strongly heating CaO in P vapour, a brown, amorphous mass is obtained; when heated with conc. HClAq, non-inflammable PH, is evolved, but with dilute HClAq the gas evolved takes fire. Probably in the first case liquid PH, is formed and at once decomposed to gaseous PII. and solid P.H; in the second case the decomposition of PII, proceeds more clowly, so that some is carried into the air with the PH3 and causes the combustion (Thénard, A. Ch. [3] 14, 12). The brown substance got by heating CaO in P vapour is sell to be a mixture of CaP and Ca₂P₂O₄ (Thénard, l.c.): this brown substance is described by Thénard as a very hard solid; unchanged in dry air; deliquescent in moist air; burns when heated in air; acted on by water free from air gives CaOAq and PH₂, PH₂ decomposes to PH₄ and P₂H₄ and the P₂H is decomposes posed by the CaOAq to Ca(H2PO2)2Aq and H.

Calcium, Salts of. Compounds obtained by replacing H of acids by Ca. These salts form one series CaX_2 where $X_2 = Cl_2$, $(NO_3)_2$, SO_4 , CO_3 , $\frac{3}{3}$ PO_4 , A_2 . They are generally formed by the action of CaO or CaO_2H_2 on the acids in agneous solution, or by the decomposition of salts of the heavier metals by CaO2H2Aq. none of the Ca salts has been gasified, the formule are based partly on similarities between these salts and those of analogous metals which form gasifiable compounds, chiefly Zn, Cd, and Ilg. and partly on the fact that the general formula CaX2 is the simplest that can be given, provided the atomic weight of Ca is about 40 (this has been ostablished by analyses of CaCl2 CaCO, &c. and by determinations of the S.H. of the metal; v. Calcium). Salts of Ca derived from a great many acids are known; they are well marked stable bodies; many form double ealts; fow basic ealts are known. Moet of the Ca salts are soluble in water; the more insoluble are the areenite, carbonate, fluoride, oxalate, phosphate, eulphate, and enlphite. With the soluble in dilute acids. The Ca calte of non-volatile acids are generally undecomposed by heat. Ca salts derived from a great many acids are known (v. Berates, Carbonatee, Phosphatee, Bulphates, &o., &c.).

Calcium, Selenide of. CaSe. Mel. w. unknown. White solid, rapidly changing in air, prepared by heating CaSeo, to dull redness in H; [Ca,Se] = 78,000 (Fabre, C. R. 102, 1469). Calcium, Seleniocyanide of. (? CaSe₂(CN)₂).

Probably existe. Data very meagro (Crockes,

J. pr. 53, 161).
Calcium, Sulphides of. One calcium sulphides of solid: solutions phide, CaS, is knewn as a solid; solutions whir' most probably contain CaS, and CaS, respectively, have been prepared. The sulphides of Ca are decidedly less basic than those of Ba, e.g. they do not react with the sulphides of the negative metals As and Sb to form thic-

I. CALCIUM MONOSULPHIDE. CaS. Mol. w. unknown. H.F. solid, from solid materials: [Ca,S] = 92,000 (Sabatier, A. Ch. [5] 22, 598).

Preparation.—I. 11,8 is passed over CaO.II. kept at about 60°; the sele products are CaS and H2O. If the reacting bodies are perfectly dry the change does not occur (Veley, C. J. 47, 478).—2. By gently heating crystals of CaS2H2.6H2O (q.v.) in H2S; the product contains some CaO,II, (Divers a. Shimidzu, C. J. 45, 270). Schöne's method, heating CaCO, in a mixture of CO2 and H2S (P. 112, 193) is said by Divers to yield a mixture of CaS and CaO in the ratio I1CaS:5CaO (C. J. 45, 282).

Properties and Reactions .- A white amorphous solid; soluble in water with gradual decomposition, giving H2S, and solution of Cn.SH.OH (q. v.) which then slowly decomposes in air forning CaS.O. Aq and CaS. Aq (Divers a. Shimidzu, l.c.). The impure CaS produced by heating CaO with CS., or CaSO. with C, is not soluble in, although it is partially decomposed by, water. Perfectly dry Cas does not absorb CS2; but in presence of H2O a basic calcium thiocarbonate, 2Cao, II, CaCS, 1011.O, is produced (Veley, C. J. 47, 486). Salatier (A. Ch. [5] 22, 598) gives the thermal value [CoC] and 200 [5] 1.3 (200 [6] 1.3 (2 [Ca,S] = 92,000; [CaS, Aq] = 6,010 (? puro materials).

When CaS II. CALCIUM POLYSULPHIDES. (prepared by heating CaO in CS₂ and CO₂ and therefore containing some CaO) is beiled with S and H.O, it dissolves, forming an orange-red liquid: the quantity of S which goes into solution corresponds with that required to form CaS, and CaS,; if more S is used it is deposited on cooling the liquid; if less S than S, to CaS is used, some of the CaS remains undissolved. Both solutions are decomposed on concentration with ppn. of CaO, H, and S, and evolution of H,S (v. Schöne, P. 117, 58). Warm CaS, H,Aq dissolves S very readily, forming a solution of CaS, and evolving H2S; the solution is completely decomposed (if cold and dilute) by H.S forming CaS.H.Aq with ppn. of S (Divers a. Shimidzn, C. J. 45, 270).

CaS, Aq is decomposed in centact with air. By boiling 3 parte CaO, 1 part S, and 20 parts HO for some time, and allowing to stand for several days, orange-red needles are obtained of

Schön, P. 117, 58), 2CaO.CaS, 10 or 11 H.O according to Geuther (A. 224, 178). If CaS (prepared by action of CS, and CO, on CaO) is boiled with much water and filtered hot, CaSO, is said to separate out and then yellow needless of 5CeO.CaS, 20H,O (II. Reso, P. 55, 433), or 4C O.CaS, 18H,O (Schöne, P. 117, 58), or 3CeO.CaS, 14 or 15 H,O (Churches-3CaO.CaS, 14 or 15 H.O (Geuther, A. 178). These exysulphides are easily decomposed. Calcinm, Sulphocyanide of. Ca(CNS). By saturating HCNSAq with CaCO, v. SULPHO CYANIDES, under CYANIDES.

Calcium Sulphydrate (or hydrosulphide); and Calcium hydroxy-sulphydrate (or hydroxyhydrosulphide). CaS.H. 611.0, and CaSH.OH.3H.O. By passing H. By passing H.S solution of CaO containing solid CaII, O2, CaS,H...6II.O is formed: 1 part CaO is added to 3-4 parts wem water; when cold, H2S is passed into the semi-solid substance until all has dissolved; more CaO is added, little by little, the whole being surrounded by ice, and H.S ie passed in until a little CaO remains nndissolved; the liquid is quickly decanted into a tubo kept in ice; the crystals which separate are drained and a current of dry ILS is swept ever them at 0'. Air must be excluded during the entire operation (Divers a. Shimidzu, C. J. 45, 270; Veley, C. J. 47, 478). CaS₂II₂·6H₂O forms colonrless prismatic crystals, which melt in their water of crystallisation, giving off H2S and corming Ca.Sil.OllAq and CaO2H2. At about 15°-19°, ILS is evolved even in an atmosphere of ILS. CaS,IL,GHO is very soluble in water and alcohol. CaS,IL,Aq is slowly exidised in contact with air, giving a little CaS2O3Aq and CaS Aq. Thomsen (Th. 3, 251) gives the thermal value [Ca,S²,H²,Aq] = H_{2}^{2} 50.

References.—Pelonze, C. R. 62, 408; H. Rose, P. 55, 133 ; Berzelius, S. 34, I2; P. 6, 442;

Böttger, A. 29, 79; 33, 344.

When a stream of 11 is passed through an i.e-cold solution of CaS.H., crystals of Ca.S.H.OH.3H.O are formed, and H.S is evolved. The same compound is formed by the combination of H₂O with CaS, as in the interior of heaps of soda-waste; and by the mutual action of CaO.II. and H.S. as in the purification of coal gas. Calcium hydroxysulphydrate crystallises in colourless four-sidea prisms; it is soluble in water with decomposition into CaS_H_Aq and water with decomposition into CaS.H.Aq and CaO.H.; insoluble in, but slowly decomposed by, alcoluble [CaS.H. goes into solution and CaO.H. remains; Divers a. Shimidzu, C. J. 45, 270]. It absorbs CS. forming a basic thiocarbonate 2CaO.H. CaCS. [10] [AO; it is the active agent for absorbing CS. in gas-purification (Veley, C. J. 47, 478).

M. M. P. M. M. M. P. M. (Veley, C. J. 47, 478)..

CALLUTANNIC ACID CHILLO, Occurs in Calluna vulga, is, the common Ling. The green parts are extracted with alcohol, water is added, and from the filtrate the lead salt is ppd. by Pb(OAo). Amber-colouled mass. Its colution in alkalis rapidly absorbs oxygen from the air. Boduces AgNO₂Aq. Fe₂Cl₂ gives a green colour.

Dyes mordanted wool sulphur - yellow. —

Salts. — (PbC, H₁₂O₂)₂(PbO)₃ aq(?)—

(PbC₁,II₁,C₀)₂(PbO)₂ 2aq(?).

Bn(C₁,II₁,C₀)₂(SnO₂)₂ 2aq(?).

Boiling dilute mineral acids convert callutamite \$CaO.CaS, 12H,O (Herschel, N. Ed. P. J. 1, 8; | acid into calluxanthin C14H10O,, a yellow floceu-

lent pp., sl. sol. cold water, v. sol. hot water and alcohol. Ite alkaline solutions rapidly absorb oxygen from the air (Rochleder, A. 84, 354). CALMUS v. CALMUS.

CALOMEL. Merculous ohloride (HgC). V.

MEROURY, OHLORIDES OF.
CALOPHYLLUM RESIN 3, H, O,. S.G. 112. A resin from Calophyllium calaba or longifoium of South America. Said to give butyrio soid on oxidation (Levy, C. R. 18, 242).

CALORIMETER. Instrument for measuring quantities of heat. V. Physical methods, Sect. THERMAL

CALYCIN C₁₈H₁₂O₅ [240° uncor.]. Occurs in a yellow lichen, Calveium c. rysoccphalum, from which is is extracted by boiling ligroin (Hesse, B. 13, 1816). Sublimable. Yollow needles or prisms. S. sol. cold petroleum spirit, petrolemm ether, ethor, alcohol, and acetic acid, more easily in the hot solvents. By strong aqueous KOH it is split up into oxalio and phenyl-acetio acids. Carbonated alkalis give salts of calycie acid.

CAMELLIN C₃₂H₃₄O₁₀. A glucoside occurring in the seeds of Camellia japonica (Katzujama, Ar. Ph. [3] 13, 334). Extracted by alcohol, and ppd. by lead acetate. White powder with bitter tasto, insol. water. Somewhat resombles digi-

CAMPHANIC ACID C, H, O, i.e.

Oxy-camphoric anhydride.

From bromo-camphoric anhydride, the product of the action of bromine on camphoric anhydride, by treatment with water (Wroden, A. 163, 330; Woringer, A. 227, 3). From campholic acid and bromine (Kachler, A. 162, 264). Formed also as a by-product in the preparation of camphoric acid by exidation of camplior with 11NO, (Roser, B. 18, 3112). According to Fittig (A. 172, 151) it is a lactor's acid, formed vid $C_s II_{12} Br < CO > O$ and $C_s H_{13} (OH) (CO_2 H)_2$.

Properties .- Feathery crystals or Lisms (from water). Monoclinic, a:b:c = 1.2723:1:1.522. À=66°34′.

Salt.-BaA', 3, aq

Reaction .-- 1. On distillation camphanic acid

gives CO_2 , campholactono C_bH_{14} \subset , and lauro-

nolic acid C₅H₁₅CO₂H.—2. K₂Cr₂O₇ and H₂SO₄ oxidise it to camphoronic acid C₉H₁₄O₆ (Bredt, B. 18, 2989).

CAMPHENE v. TERPENES.

CAMPHENOL v. Bo. NEOL and CINEOL. CAMPHENYL-p-TOLYE-AMIDINE

C.H.,C(NH₂):NC,H, [115°]. Fr.10 white glistening needles (from ligroin). Formed by heating campholenonitrile C.H.,CN with p-toluidine hydrochloride at 250° (Goldschmidt a. Koreff, **B**. 18, 1633).

CAMPHIC.ACID C10H1502. S. .14 at 100. [a]n = 15°45' (in alcoholic solution). Formed together with camphoric acid by passing air through a boiling solution of sodium eamphor, CteHisNaO in xylene. Thick mass, v. sol. alco-his and ether. KMnO, oxidiscs it to camphoric acid. The calcium salt distilled with calcium formate gives camphor and camphrene CaH14O

(c. 288°) (Montgolfier, 4. Oh. [19 14, 70; C. R

CAMPHILENE &. TERPENES.

CAMPHIMIDE C, HISN or CaHI

Formed together with dicamphorilimide by distilling the hydrochlor le of amido-camphor (v. CAMPHOR) with steam (Schiff, B. 13, 1405).

CAMPHINE v. TERPENES. CAMPHO-CARBOXYLIC ACID v. CAMPHOB

CARBOXYLIC ACID.,
(a)-CAMPHOGLYCURONIC ACID C10H21O. [130°]. S. 5. [a]_p = -33°. Occurs, together with uramido-camphoglycuronic acidhe urine of dogs that have taken camphor (Schmiedeberg a. Meyer, H. 3, 422). Small thin laminæ (containing aq); v. c. sol. alcohol and hot water, insol. ether. Boiling dilute HCl splits it up into glycuronio acid CaH10O7 and crystalline oampherol C₁₀II₁₀O₂ [198°]. HNO₃ oxidises it to camphoric acid.—BaA".—BaA" 2aq.—AgHA" 2nq.

(8) Camphoglycnronic acid C_{1s}H₂₁O₅. An amorphous modification of the preceding, formed by warming it with baryta.—AgHA"3aq: crystals, more soluble than the Ag salt of the

CAMPHOL a name for Borneol (q.v.).

CAMPHOLACTONE [50°].

(230°-235°). From camphanic acid by distillation, together with lauronolic acid (Woringer, A. 227, 10). Slender needles (from water). Has a pungent odour of camplior. Like other lactones, its solution becomes cloudy when gently heated, but the oily drops afterwards dissolve up again. Volatile with steam. K₂CO₂: separates it than its aqueous solution. When boiled with baryta the salt of the corresponding;

oxy-acil, C.H., (011)CO.H. is formed.

CAMPHOLENE C.H. (136°). V.D. 4·35.

Prepared by the action of dehydrating agents on campholic acid (Delalande, A. 38, 340) and by distilling potassium campholate with soda-lime (Kachler, A. 162, 266). Probably identical with the hydrocarbon got by distilling the calcium salt of campholenic acid (Goldschmidt, B. 20, 483). The name campholone has also been given to C,H, (o. 123°) obtained by the action of dehydrating agents on camphoric acid

camphoric soid and its smiles (Ballo, B. 12, 324).

CAMPHOLENIC ACID C₁₀H₁₀O₂ i.c.

C₃I1₁, CO₂:1 or C₈I1₁₃(CO₄H):CH₂. Oxy-camphor.

(c. 260°). Colourless oil. Formed by saponification of its nitrile which is obtained by heating eamphoroxim with acetyl chloride. Formed also by treating an alcoholic solution of (8)-di-bromo-camphor with sodium-amalgam (Goldschmidt a. Zürrer, B. 17, 2069; Kachler a. Spitzer, B. 17, 2400: M. 3, 216; 4, 643). The Ca salt on dry distillation yields C.H., possibly campholene (Goldschmidt, B. 20, 483). Oxidising agents give oxy-camphoronic acid. At 250° the NH₄ salt gives the amide [127°].—NH₄A'.—BaA'₂4ac.

Anide C₂H₁₅ CONH₂. Isocamphoronim

[125°]. Glistening plates; sol. alcohol, ether, and cono. acids. Formed by heating the nitrile with alcoholic KOH, or by heating the ammonium ealt of the acid to 250°. By distillation tory brneols (Montgolfier, C. R. 88, 915; A. Ch. with P.S., it yields the nitrile (Nägeli, B. 17, [5] 14, 20). 805; Goldschmidt a. Zürrer, B. 17, 2069).

Nitrite.—C.H., CN. (216°). Formation.—
1. By heating camphoroxim with Acci which removes H₂O.—2. By distilling campholenamide (ieocamphoroxim) with P.S. Reactions. -1. By heating with alcoholic KOH it is converted into campholenamide. By Eng boiling with alcoholic KOH it yields campliolenic acid.-2. By heating with hydroxylamine it gives an amidoxim C10H18N2O which crystallises in white plates melting at [101°].—3. Reduced in alcoholic colution by Zn and HCl to the amine C_pH_CH₂NH₂ (Goldschmidt a. Zörrer, B. 17. 2069; Goldschmidt a. Koreff, B. 18, 1634).—4. Successivo treatment with sodium amalgam and HCl yields CheHa, NaCl, the hydrochloride of camplyl-di-phenyl-hydrazinamino C₄H₁₄(CH₂NH₂)N₂H₂Ph. G. 17, 155). [157°] (Balbiano,

Nitro - eampholenic acid $C_{10}H_{15}(NO.)O_{e}$ tro - oxy - camphore [164°] (Z.); [170 $^{\circ}$] Nitro - oxy - camphor (K. a. S.). Formed by nitration of campholenic acid (Zürrer, B. 18, 2228; Kachler a. Spitzer, M. 4, 643; B. 15, 2336; Swarts, B. 15, 2135). Monoclinic pyramids, a:b:c=.76:1:.13; $\beta=$ 89° 18'. Sol. hot alcohol and other. Reduced by tin and HCl to amido-campholenic acid whose hydrochlorido crystallises in laminæ [250°].

CAMPHOLIC ACID $C_{16}II_{18}O_{2}$. Mol. w. 170. [95°] (K.); [106°] (M.). $[\alpha]_{j} = 50$ ° (in alcohol). Formation.—1. By passing camphor-vapour

over nearly red-hot notash-lime (Delalande, A. Ch. [3] 1, 120).-2. By adding potassium in small pieces to a solution of camplior (1 pt.) in boiling petroloum (3 pts.) at 120° (Malin, A. 145, 201).—3. By boiling camphor with alcoholic KOH (Kachler, A. 162, 259).—4. By hoating camphor with Na at 280 Montgolfier, A. Ch. [5] 14, 99).

Properties .- Monoclinio prisms (from diluto alcohol) or nodular groups of lanima (from ether-alcohol). V. sl. sol. water; volatile with

Reactions.-1. HNO3 gives first camphoric and then camphoronic neids. - 2. Moist Br gives at first camphoric acid, then bromocamphoric anhydride, and lastly oxy-camphorio anhydride C10 II,104. - 3. P2O, gives campholone; red-hot soda-lime acts similarly. •

Salts. - KA'2aq: lamino. - CaA', aq. AgA'.

Chloride (224°) (Kaehler, A. 162, 265).

CAMPHOR C₁₆H₁₆O. Mol. w. 152. [175°]. (204°). S.G. 12 992. S. 1. S. (alcohol of S.G. 806) 120. V.D. 5.32. R₀₀ 78.11 (in a 32.3 p.c. bonzens solution) (Kanonnikoff). [a]_D = 55.4 - 1372 q (where q = no, of grass, of alcohol in 100 grms, of solution).

Occurs in the wood and bark of Laurus camphora, from which it is extracted by distillation with steam followed by sublimation. Varieties of camphor occur also in several essential oile (v. infra). Campho may be recovered from its brome-derivative by the action of nascent H or of alcoholic KOH (Schiff, B. 13, 1407; 14, 1377). Camphor is also formed by dietilling calcium camphato with calcium formate and by oxidicing dextro- and levo- rota-

Properties.-Hexagonal prieme, terminated by hexagonal pyramids (Descloizeaux, A. Ch., [3] 56, 219; Cazeneuve a. Morel, C. R. 101, 438). Tough, with peculiar odour; eublimes at ordinary temperatures. Small pieces rotate upon puro water. V. sl. sol. water, v. eol. ordinary solvents. Camphor is dextro-rotatory, the rotation varying greatly with the nature and strength of the solvent (Arndtsen, A. Ch. [3] 54, 403; Landolt, A. 189, 334). Ite refractive power is that of a saturated compound (Gladstone, C. J. 49, 621).

Reactions. 1. Camphor (5 kilos.) gives, when oxidised by HNO3, (1.7 kilossef) purs camphoric acid insol. cold water, and (18 kilos. of) erndo camphoronic acid. Besides camphoronic acid the solublo portion contains (1 kilo. of) dinitroheptoic acid, and (2 kilo, of) soide $C_{s}II_{14}O_{s}$ (hydro-oxyeauphoronic acid), $C_{s}H_{12}O_{s}$, $C_{r}H_{12}O_{s}$ (?) [145°], and another acid. A very small quantity (2 g.) of mesocamphorie acid, C₁₀H₁₄O₄, is also got. This forms woolly nssdles, soluble in cold water [120°]. 2. By oxidation with CrO, it gives camphoronic acid CoH12O, and hydro-oxy-camphoronic Call 140 but not adipic acid (Kachler, B. 13, 487; cf. Ballo, B. 12, 1597). Alkalinc KMnO, gives camphoric acid (Grosser, B. 14, 2507).—3. The chist products of the dry distillation of camphor with ZnC, (2 pts.) are m-methyl-isopropyl-benzene (m-cymene) and (1:2:4)-di-methyl-cthyl-benzene (laurene), together with smaller quantities of (1:2:3:5) - tetra - methyl - benzeno (isodurene), carvaerol, camphorone, and various other bodiss (Armstrong a. Miller, B. 16, 2255) such as CH, benzene, tolnene, xylene, and y-comene (Fittig. A. 145, 129; Rommier, Bl. 12, 383; Lippmann a. Longuin ne, A [2] 5, 413; Montgolfier, A. Ch. [5] 14, 87). 4. By the action of iodine it yields a hydrocarbon C₁₀H_a, carvaerol, (1:2:4)-dimetayl-ethyl-benzene, (1:2:3:5)-tetra-methyl-benzene, and traces of ordinary cymene (A. a. M. : c. Armstrong a. Easkell, B. 11, 151; Rayman a Preis, B. 13, 346).-5. By treatmont with P2O5 ordinary cymeno is formed, which is also the chief product of the action of P.S. but accompanied in the latter case by small quantities of m-methyl-isopropyl-benzene and tetramethyl-benzene (Delalande, A. Ch. [3] 1,268; Armstrong a. Miller, B. 16, 2255). - 6. By distillation over red hot zinc-dust a mixture is formed of toluene, p-xylene, cymeno, and a little benzene (Sehrötter, B. 13, 1621)!-7. Cone. H.SO, forms can here or camphorphorone C.H.O (Chautard, C. R. 44, 66; Schwanert, A. 123, 298).—8. Boiling alcoholic KOH forme campholic acid and borneol (Berthelot, A. Ch. [3] 56, 94; Id. [2] 17, 890; Montgolfier, Bl. [2] 18, 114; 25, 12; Wheeler, A. 146, 84; Kachler, A. 162, 268). Campholic acid is also formed by passing camphor-vapour ever heated soda lime. 9. Cl has no action, but in presence of alcohol or PCl, chlorination ensues (Claus, J. pr. 25, 257).—10. HClO forms chloro-camphor.—11. Br forms $O_{10}H_{16}OBr_2$ which readily splits up into HBr and bromo-camphor.—12. ICl at 250° forms CCl₄, C₂Cl₅, and C₅Cl₅.—13. PCl₅ form C₁₀lI₁₀Cl₂.—14. Camphor absorbs HCl (Bineau, A. Ch. [3] 24, 328). Aqueoue HCl at 170°

splits it up into H₂O and cymene (Alesejeff, J. R. 12, 187).—15. Camphor absorbs SO₂, becoming liquid.—16. It also absorbs NO₂.— 17. Cono. HIAq at 200° forms C_{1e}H_{1s} (163°), C_eH_{1e} (135°-140°), find C_{1e}H_{2e} (170°-175°) (Weyl, Z. [2] 4, 496; B. 1, 96).—18. Solium amalgam has no action.—19. Na acting (# 90° on a solution of camphor in toluenc forms sodium camphor and sodium borneol (Baubigny, Z. [2] 2, 408; 4, 298) (cf. p. 672). -20. CO. gas passed into the product of the action of Na on camphor in toluene forms the carboxylic acids of camphor and of borneol (Baubigny, Z. [2] 4, 482, 647). Air passed into the same mixture forms camphoric acid (Montgolfer, A. Ch. [5] 14, 75).-21 Camphor does not combine with NaHSO₃ (Fittig a. Tollens, A. 129, 371).— 22. AcCl has no action .- 23. Converted in the animal economy (of a dog) into camphoglycuronio acid (q. v.).—24. Melted camphor absorbs BF₃ forming C₁₀H₁₆OBF₃ [70°]; when this is heated for 24 hours there is formed cymene and its polymerides, C_0H_{10} , $C.H_{12}$, and other hydrocarbons (Landolph, C.R. 86, 539).—25. Chloral hydrate forms an unstable molecular compound with camphor. It is a viscous liquid, sol. alcohol and CHCl3, insol. water (Cazenchye a. Imbert, Bl. [2] 34, 209; Zeidler, J. 1878, 645). -26. By heating with ammonium formate at 220°-240° it yields formyl-bornylamine

C₈H₁ CH₂ (Leuchart a. Bach, A 20,

104).—27. It does not react with phenyl-cyanate (L. a. B.).—28. Hydroxylamine forms an oxim, v. Camphoroxim.

Phenyl-hyd razide C₁₀H₁₆:N_.HPh. (233°) at 10 mm. From camphor and phenyl-hydrazine. Oil. Split up by dilute HCl into camphor and phenyl-hydrazine, and by 1ry HCl into aniline and the norice of campholenic acid_(Balbiano, G. 16, 132).

Constitution.—The action of hydroxylamine on camphor coupled with the fact that camphor does not combine with NaIISO₃ indicates that it is a ketone. The ready formation of benzene derivatives indicates a six-carbon ring. Its optical properties are those of a saturated compound, v. also Terrenes.

The two following formulæ amongst others have been proposed for camphor:

CH₂.C(C₃H₇).CH₂

| | | (Schiff, A.226, 249; Kanon-CH₂C(CH₃). CO nikoff, J. pr. [2] 32, 511; cf. Kekulé, B. 6, 931); CH₂CH₂CH. CH₂

CH₂.CH₂.CH . CH₂
(Armstrong a. Miller, B.

CH.CH.CH. CMe.CO
16, 2255). V. also TERPENES.

Chloro - campher C₁₀H_{1,5}ClO. [95°]. From camphor and conc. HClOAq (Wheeler, Am. S. [2] 45, 48; A. 146, 81). Crystalline powder (from alcohol); at 200° it gives off HCl. Decomposed by alcoholic AgNO₂.

(a)-Chloro-camphor C₁₀H₁₃ClO. [84°]. (C.); [93°] (B.). (246°). [a]₁ = 90°. Formed by passing dry Cl into a solution of camphor (760g.) in absolute alcohol (230g.) (Cazeneuve, C. R. 94, 1530; Bl. 138, 9; 44, 161). Formed also by heating chloro-camphor carboxylic acid (Schiff a. Puliti, B. 16, 887; Balbiano, G. 17, 95). Hard, brittle,

monoclinic needles; smelling like samphor; rolatile with steam. Not decomposed by alcoholic AgNO₂. Sodium-amalgam reduces it to camphor; the copper-zine couple, and hot sodalime, act similarly. Alcoholic KOH at 180° gives borneol. Phenyl hydrazine forms C₁₀H₁₀(N₂HPh) [56°] (B.).

C₁₀H₁₃(N₂HPh)(N₂H₂Ph) [56°] (B₃).

(B)-Chlero-camphor C₁₀H₁₃ClO. [100°]. (246°).

[a]₁ = 57°. Deposited from the mother-liquor after the preceding has separated (Cazeneuve, C. R. 95, 1358; Bl. [2] 30, 116). Soft, minute, needles, more soluble than the preceding; miscible with boiling alcohol. Not decomposed by alcoholic AgNO₃, but converted by boiling alcoholic KOH into the preceding body. Phonyllydruzine produces the same compound [56°] as

with the preceding (B.).

(a).Di-chlore.campher C₁₀H₁₁Cl₂O. [96°]. S.G. 4·2. [a]_j = 57·3° (in alcohol or chloroform). Formed by passing dry chlorine for soveral hours into camphor dissolved in absolute alcohol at 80°-90°. The product is ppd. by water and caystallised from alcohol (Cazeneuve, C. R. 94, 730, 1058; Bl. [2] 37, 454). Trimetric prisms (from alcohol); a:b:c=1·8358:1:1·4820; sl. sol. cold, v. sol. hot, alcohol; almost insol. water but rotates upon it. Above 150° it blackens, giving off HCl. Insol. HOAc (difference from camphor). It forms a liquid combination with aldeliyde.

(β)-Di-chlore-campher C₁₀II₁₁Cl₂O. [77°]. [a]j = 57·4° (in alcohol); 60·6° (in CHCl₂). Ppd. by adding water to the mother-liquor from which the preceding has crystallised (Cazeneuve, C. R. 94, 1360; Bl. [2] 38; 8). Crystallises from alcohol with difficulty; v. e. sol. alcohol, ether, and chloroform; liquefied by ohloral hydrate

(difference from the preceding).

Tri-chlore-camphor $C_{10}\Pi_{12}Cl_2O$. [54°]. [a], = 64° (in alcolar). Formed by saturating chlore-camphor with chlorine at 100° (Cazencuvo, C. R. 99, 609). Minute crystals, insol. water, sol. other menstrua. Gives off HCl when heated.

Brome-campher $C_{10}H_{15}BrO$. [76°]. (274°). S.G. 1'44 R_{∞} 88°5 (in a 7'37 p.c. alcoholic solution) (Kanonnikoff, J. pr. [2] 32, 504). [α] $_{\rm b} = 139$ °. Formed by heating camphor dibromide at 100° (Perkin, C. J. 18, 92; Maisch, C. C. 1873, 437). Monoclinic prisms (from alcohol); m. sol. alcohol, v. sol. CHCl₃ and benzene; may be sublimed (Montgolfier, BL [2] 23, 253).

Reactions.—1. Sodium-amalgam reduces it, in alcoholic solution, to camphor; alcoholic KOII also gives camphor. Sodium added to its solution in toluene gives sodium camphor (R. Schiff, B. 13, 1407).—2. PCl, has no action even at 100° (Schiff, B. 14, 1378; Kachler a. Spitzer, M. 3, 205).—3. Heating with ZnCl, at 160° gives a mixture of p-xylene hexahydride and a phenol C₁₀H, O apparently identical with the carvacrol obtained by the action of I on camphor (R. Schiff, S. 13, 1407).—4. Nitric acid (Armstrong, B. 12, 1358; R. Schiff a. Maisson, C. 10, 317).—5. Phenyl - hydrazine forms C₁₀H₁₃(N₂PhH)(N₂PhH₂), [56°] (Balbiano, G. 17, 35, 155).

(a)-Di-brome-camphor C₁₀H₁₄Br₂O. [115°].

(a)-Di-breme-camphor $C_{10}H_{14}Br_2O$. [115°]. Formed, together with the preceding, by heating brome-camphor (1 mel.) with Br (2 mels.) for 11

hours at 180° (K.a. S.; cf. Swarts, Z. [2] 2, 205; B. 15, 1622). Jormed also by lieating (B)-dibromo-camphof with gaseous HBr at 130° (Swarts, B. 15, 2135). Trimetric crystals; a:b:c=95:1:52; sl. sol. alcohol, ether, and petroleum. Less volatile with steam than the preceding.

Reactions .- 1. Sodium-amalgam gives camphor and oampholenio noid .- 2. Conc. IINO,

gives di-bromo-nitro-camphor [130°].

(β)-Di-bromo-camphor C₁₆H₁₁Br₂O. [61°]. Formed, together with its isomeride, by the action of Br (1 mol.) on bromo-camphor (1 mol.) for 7 hours at 120° (Kachler a Spitzer, M. 3, 208; Zepharovich, M. 3, 231; cf. R. Schiff, G. 11, 178; Mongolfier, Bl. [2] 23, 253). Trimetric orystals, a:b:c=2.0685:1:1.5778 (Cazencuve a. Morel, Bl. [2] 44, 161) = 1.944:1:1.558 (K. a. S.). V. sol. alcohol, ether, and petroleum.

Reactions. -1. Alcoholio KOII, or sodiumamalgam, reduces it to brome-campher, and finally to camphor .- 2. Sodium and CO2 form camphor carboxylic acid.-3. Conc. HNO gives camphoric, and hydro-oxy-camphoric, acids, together with bromo-di-ritro-methane (Kaehler

a. Spitzer, M. 4, 554).

(a)-Chloro-bromo-camphor C ... H ... ClBrO. [98°]. [a] = 78°. Formed by heating chlorocamphor [84°] with bromine in scaled tubes for 5 honrs at 100°. White needles. Insol. water, sol. hot alcohol, other, and CHCl₃ (Cazeneuve,

Bl. [2] 44, 115; C. R. 100, 802).

(B)-Chloro-bromo-camphor C, II, ClBrO. [51.5°], [a] = 51°. Prepared by heating oldorocamphor (1 mol.) with bromine (2 mols.) in sealed tubes for one hour at 100° C. Hard, trimetric crystals, a:b:c=1:914::1:1:5395. Insol. water, v. sol. alcohol, v. sol. cun. s, CHCl., C.H. and CS. (Cazcueuve, Bl. [2] 44, 115; C. R. 100, 859). Decomposed by Ling AgOAcAq difference from preceding). (difference from preceding).

Iodo-camphor C10H13IO. [440]. Formed, together with NaCy and NaI, by the actiof of ICy on sodium-borneol dissolved in benzene (Haller, C. R. 87, 695). Monoclinic crystals, insol.

water, sol. alcohol. Decomposes at about 150°.
Nitro-camphor C₁₀H₁₅NO₄ [83°]. Prepared by the action of alcoholic KOH on brone nitrocamphor (Schiff, B. 13, 1402; G. 10, 330; 11, 21). Dissolves in aqueous alkalis. Gives a red colouration with Fe₂Cl₆. HNO₂ gives a nitrosocompound. On oxidation with INO3 it gives camphorio acid. By reduction it gives amidocamphor. Bromino forms $C_{a_0}H_{11}N_{a_0}Br_{a_0}I_{12}$ [95°]. Cl forms similarly $C_{a_0}H_{11}N_{a_0}Cl_{20}I_{12}$ [110°]. Steam-distillation gives camphoric acid and anhydride and NH_a. This substance is probably a mixture of the two following.

(a)-Nitro-camphor C10H15(NO2)O. [a]_j (19978 p.c. in benzene) -98? (3:33 p.c. in alcohol) -7.5°. Formed, together with its (3)-isomeride, by the action of Zn, Cu, Fe, or alkalis on either chloro-nitro camphor dissolved in alcohol. Best prepared by using the copperzino couple. The resulting zinc-sult of nitro-oamphor is decomposed by HCl. The (a), compound is the less soluble in cold alcohol (Cazeneuve, C. R. 103, 275; 104, 1522; Bt. [2] 47, 920). Trimetric prisms. Decomposes at 160°. Leevorotatory. Its rotation varies with concentration of the solution. It forms a com-

pound with benzene. It reddens litmus, and decomposes carbonates. Fe₂Ol_a colours its alcoholic solution blood-red.

(\$\beta\$)-Nitro-camphor $C_{10}H_{11}(NO_9)O$. [98°]. [a], (3°33 p.c. in benzen,) -75° ; (3°33 p.c. in alcoho) $+7^\circ$ 5°. Prepared as above. Soft, fornlike, crystals (from alcohol). Insol. water, sol. other menstrua. Less stable than the (a)-isomeride. Fe_xCl_s colours its solutions red. Its salts are more soluble than those of the (d)- compound.—NaA' — ZnA'2: sol. water.

(a) - Chloro - nitro - camphor C10H11Cl(NO2)O. [95°]. [a]₁ = -6·2°. From chloro-camphor [93°] [1 pt.] and funing HNO₃ (1 pts.). Large trime-tric prisms (from alcohol); $a:b:c=2\cdot022:1:1\cdot475$ (the author does not say whather these numbers belong to this or to the following body). Insol. water, m. sol. cold aloohol. Decomposes above 100°. Reduced by nascent hydrogen to nitrocamphor (Cazencuve, C. R. 96, 589; Bl. [2] 39,

503).

 (β) -Chloro-nitro-camphor C₁₀II₁₄Cl(NO₂)O.
 [98°]. [a]_j = 10·5° (in alcohol). Occurs in the mother-liquor from which the preceding has separated. Soft crystals, v. sol. cold alcohol. Reduction gives nitro-camphor. Less stable than the (a)-isomeride, for alkalis remove Cl forming nitro-camphor even in the cold. Not decomposed by alcoholie AgNO₃ (Cazenouve, C. R. 98, 306; Bl. [2] 41, 285; 41, 161; 47.

(i) Brome - nitro - camphor C₁₀ll₁₁BrNO₃. [105°]. [a]_i = -27°. Prepared by nitration of bromo-camphor. Trimetric crystals, a:btc=2-0851:1:1-5423. Nearly insol. cold alcohol. By the action of alcoholic KOII or mascent II. it gives nitro-camphor (Schiff, G. 10, 324; B.

13, 1402; 11, 1377).

Di-bromo-nitro-camphor C10H13Br (NO2)O. [130°]. From (1)-di-bromo-camphor by nitration (Kaehl . a. Spinser, M. 4, 554). Trimetric prisms or readles, a:bic=1.76:1:1:49: Reduced by tir and HOAc to anido-camphor.

Amido camphor GiuHi.NO. (217°). Waxy solid. Strong base of alkaline reaction. Prepared by reduction of nitro camphor with sodium-amalgam in alkaline solution. It reduces Fehling's solution, AgNO₃, and HgCl₂. With HNO2 it produces oxy-camphor (Schiff, B. 13, 1404). On distillation of the hydrochleride of amido camphor with steam, 'dicamphylamine' C_aM₁NO₂ passes over and 'camph-imido C_aM₁No₂ passes over and 'camph-imido C_aM₁No₂ passes over and 'camph-phylamin' [160] crystallises in needles, insol. acids, volatile with steam. Camphimide forms crystalline takes, soluble in acids; nitrous acid converts its hydrochloride into 'diazo-camphor' C₁₀H ₄N 0 [71°] which may be reduced by Zn and HOAc to amido-camphor. Diazo-camphor is converted by heat into 'de-hydro-camphor' C₁₀H_{1,0} [160'] (R. Schiff, G. 10, 362; 11, 171; B. 14, 1375).

Oxy-camphor (?) C₁₀'I₁₀O₂ [155']. Prepared by the action of IINO₂ on amido-camphor (?) C₁₀'I₁₀O₂ [155'].

(Schiff, B. 13, 1404). Colourless ci stals. Volatile with steam.

Wheeler's chloro-oamphor (q. v.) gave with

alcoholic KOH an 'oxy-camphor' [137°].

The acctyl derivative [69°] of an 'oxy-camphor' [249°] is formed by oxidising acceptborneol.

An 'oxy-camphor' [61°] is formed by oxi-dising camphone (v. Terrenes) (Kachler a. Spitzer, A. 200, 358).

V. also Campholmnio acid. .

'Nitro-oxy-camphor,' v. NITRO-OAMPHOLENIC

Cyano-camphor v. Nitri'e of Camphon car-BOXYLIC ACID.

Ethyl-camphor C₁₀H₁₅EtO. (228°). 8.G. 22 946. [a]_j = 161°. From sodium-camphor and EtI (Baubigny, Z. [2] 4, 481). Oil. Isoamyl-camphor C₁₀II₁₃(C₈II₁₄)O. (278° cor.).

 $[a]_j = 59.4^\circ$. From sodium-camphor and isoanyl iodide (B.).

'Nitrohexcic acid' C.II. (NO.)O. i.e. Mc.CH(NO.).CMe. CO.II(?) [115]. From dinitroheptoic acid and sodium-amalgam (Kullhem, A. 167, 45; Kachler, A. 191, 157). Rectaugular four-sided columns (from water). Monoelinic; b:c=1: 6115; $\beta=83$ 30'. After several fusions it melts at III 5°. Quickly heated, it explodes. Reactions.—1. Dissolved in a little aqueous

KOH, mixed with KNO, and dilute H.SO, a fine blue colour is formed. This colour is taken up blue colour is formed. by ether; hence the body is probably akin to pseudo-nitroles.-2. Sn and HCl produce methylisopropyl ketone, hydroxylamine, and CO2.-3. Baryta in scaled tubes at 95° does not decom-

pose it, but forms the basic salt C_o||J_oBaNO_o.

Dinitrohexoic acid C_o||I_o(NO_o)_oO_o? i.e.

MeC(N_oO_o).CMo_oCO_oH(?) [215°]. Got by Kullhem by treating the residues in the preparation of camphoric acid with strong IiNO, (A. 163, 231; Kachler, A. 191, 155). Monoclinic plates; $a:b:c = .5735:1; .6024; \beta = 70° 42'.$ M. sol. cold water, v. sol. hot water. Explodes when rapidly heated. Its anumonium salt gives with cupric acetate a bluish pp. sol. excess of the acetate (difference from camphoric acid). The free acid does not pp. caprig or lead acctate. The neutral salts give a fraculent pp. with lead acctate. .

Salt.—BaA', 3aq. Needles.

Reactions.—1. Reduced in alcoholic solution by sodium amalgam to mononitroheptois acid. 2. Reduced by Sn and HCl to methyl iso propyl ketone and hydroxylamine. -3. Potash and baryta heated with the aqueous acid in scaled tubes produce both nitro-heptoic acid and methyl isopropyl ketone.

Hydro-oxycamphoronic acid C₉H₁₄O₆. [164.5°] (Rachler, A. 191, 148). Needles. Sol. cold water. Tribasic acid. Gives no pp. with BaCl, or CaCl, and NH, even on boiling (difference from camphoronie acid). Cupric acetate gives no pp. until boiled when a bluish-green cupric salt is ppd. Lead acctate gives a white pp. seluble in excess. AgNO, gives a white pp. soluble in hot water.

Salts.-NH,II2A".- CallA" 2aq.-CasA"2. -Ba₂A'''₂.-Cu₃A'''₂.-Ag₃A'''.

ISOMERIDES OF CAMPHOR.

Levorotatory camplor $C_{16}H_{16}O$. [172° cor.] (R.); [175°] (C.). (204°). S.G. 18. 9853 (C.). [a]₁ = -47° (C.); -42° (H.). Occurs together with a terpene in the ethereal oil obtained from the leaves of fever-few, Matricaria Parthenium (Chautard, C. R. 87, 166). Apparently formed also by oxidising the levorotatory terpene obtained by treating with alcoholio KOH the pro-

duot of the action of HCl on laworotatory oil of terpentine (Riban, Bl. 24, 19). Obtained also by oxidising the borneols of madder, valerian, Ngai, and Bang-Phien (Haller, C. R. 103, 64; 104, 66). HNO, gives lavorotatory camphoric acid [a], = -46°. The corresponding bromocomphor is also levorotatory, $[a]! = -128^\circ$.

Inactive comphor $C_{10}H_{10}O$. [1739]. Got by oxidising inactive berseel (q,v) with cold HNO.

and then adding water (Armstrong a. Tilden, C. J. 35, 752). Also by oxidising inactive camphene with H₂SO₄ and K₂Cr₂O₇. Heated with HNO, it Yorms a camphorio acid, [203°] and giving when heated alone an anhydride, [2235].

Inactive camphor From Oil of Says. $O_{1a}H_{1e}O$. [174]. (205° uncor.). When oil of sage is distilled, the fraction 205°-208° deposits this camphor. It apparently only differs from ordinary camphor in being inactive, for:-1. PCl, gives an oil which is converted by water into a wax-like solid, [80°] whence Na forms a white solid.—2. Boiled with HNO₃ (2:1) it forms inactive camphoric acid [186°].—3. Dissolved in toluene and treated with Na and CO, it forms inactive borncol, [200°] .- 4. Distilled with P2Ss

it forms cymene (M. M. P. Muir, C. J. 37, 685). 'Racemic' camphor $C_{20}H_{32}O_2$ (?). [179°]. This name is given to the product of the oxidation of a mixture of equivalent quantities of levo- and dextro- rotatory borncol, and is therefore inactive by compensation, as racemic acid is (Haller, C. R. 105, 66). It gives a bromoderivative [51°] and a camphoric acid [205°]. The 'racemic' camphor, bromo-camphor, and camphoric acid were also prepared by mixture and found to be identical with the above. They differ in solubility as well as in melting-point from the active compounds. The borneol of amber, and consequently the camphor derived therefrom, appear to be a mixture of dextroand hevo-rotatory varieties in uncqual proportions.

Camphors, or bodies resembling camphor, have been found in many essential oils, e.g. oils of alant (p. 94), absinthe (p. 2), chamomile, eucalyptus, lavender, nutmeg, rosemary, &c. When a camphor is converted into borneol, the rotatory power of the resulting borneol varies with each operation, but the camphor regenerated by oxidising the borneol has in each case the rotatory power of the original camphor (Montgolfier). According to Haller (C. R. 105, 228) this may be explained by supposing that the resulting borneol is always a mixture of a stable borneol rotating in the same direction as the original campnor and of an unstable

borncol votating in the opposite direction.

CAMPHOR-CARBOXYLIC ACID C. H., O., [129°]. Formed as a by-product in the preparation of borneal from camphor by the action of Na on a solution of camphor in toluene, the product beings treated with CO2. It is produced by the union of CO, with sodium-camphor (Baubigny, Z. [2] 4, 481, 647; A. Ch. [4] 19, 22I; Kachler a. Spitzer, B. 13, 1412; M. 2, 233). Long colourless monoclinic pyramids. Sol. water. Decomposes below 100° into CO, and camphor.

Reactions .-- 1. Boiling AeCl forms Cr. H.O. [196°], crystallising in needles.—2. P.O., acting

on its solution in CFOL, forms C₂:H₃₀C₃ [265°]. 3. POL forms C₂:H₃₀OL [45°] which separates from ether-alcohol in triclinic crystals a:b:c= 1: .804 : .47.

Salts.—NaA'.—BaA'.—PbA'. Ethyl ether EtA': (276° uncor.); S.G. 11. 1052; colourless fluid (Roser, B. 18, 3113).

Nitrile C.H_{1.}CyC. Cyano-camphor. [126]. (250°). Formed by passing cyanogon into a mixture of camphor and codium-camphor dissolved in hot toluene; extraoted by shaking with aqueous NaOH and ppg. by HOAc. Rect. angular prisms (from ether); sol. alcohol, ether, and HOAc. Contains an atom of hydrogen displaceable by Na or K, forming unstable salts. Conc. HCl at 100° converts it into camphor carboxylic acid. Oxidising agents give IICy and camphoric acid. Alcoholio NaOll slowly converts it into the ethor of camphor carboxylio acid (Haller, C. R. 87, 843; 93, 72; 102, 1477).

Chlero-campher carboxylic acid C11I1, ClO3. Formed by passing Cl into a solution of sodium oamphor, carboxylato (Schiff a. Puliti, B. 16, 887). Flocculent pp.; decomposes on fusion into CO2 and chloro-camp'ior.

Bremo-campher carboxylic acid C11H1,BrO, [110°]. From camphor carboxylic acid and Br (Silva, B. 6, 1092). Both the acid and its salts readily decompose into CO2 and brome camphor.—BaA'..— $\Lambda g \Lambda'$.

Oxy - camphor carboxylic acid $C_{11}H_{1}$, O_{1} . O_{2} . Formed by boiling the nitrile of camphor carboxylic acid with aqueous KOH (Haller, C. R. 87, 929). Nodales (from ether).—CaA" 6aq.—BaA" 6aq.

CAMPHOR DICHLORIDE C16 II 16 C12. [155°]. Prepared by the action of PCl, on complior in the oold: $O_{10}H_{10}O + POI_{5} = POOI_{5} + C_{10}H_{10}O$ (Spitzer, B. 11, 363, 1819; M. 1, 319). Cormed also by chlorinating bornyl chlorido (h. hler a. Spitzor, A. 200, 361). Feathery trimetric needles; a:b:c = .917:1:1.686.Easily soluble in alcolud and ether. Easily splits off HCl.

CAMPHORIC ACID C₁₀11₆O, i. CH₂-CPr-CO_.11 $C_{\bullet}H_{1\bullet}(CC_{2}H)_{2}$ or (Schiff)

CH₂—CMo—CO₂11 or Pr.OH(OO,H).CH,.CH:CMe.CO,H (W. Roser, **4. 220**, 278). Mol. w. 200. [186°]. S.G. 1·19. S. 625 at 12°. R $_{\infty}$ 83·14 (in a 1 p.c. aqueous solution) (Kanonnikoff, J. pr. [2] 34, 349). [a]_b=46°. Formed by boiling camphor or cam-Formed by boiling camphor or campholic acid with conc. HNO₂ (Kosegarten (1785); Laurent, A. Ch. 63, 207; Malaguti, A. Ch. 64, 151; A. 22, 50; Wreden, A. 163, 323; V. Meyer, B. 3, 116; Kachler, A. 162, 262). It is best to use the mixture of camphor and borncol obtained by the action of Na on camphor (Maissen, G. 10, 280). Formed also by the oxidation of chloroor bromo- camphor with alkaline permanganate (Balbiano, G. 17, 240). Monoclinic orystals; the rotation in alkaline solution has been studied by Thomsen (J. pr. [2] 35, 157). The refractive power indicates a double union, which does not agree with Schiff's formula.

Reactions.—1. Heat splits it up into water and an anhydridc.—2. Water at 180° changes it into meso-camphorio acid.—3. Fuming HCl at 200° forms C, H, and C, H, (Wreden, A. 187, 189).—4. Cono. HIAq at 200° gives xylene tetra-Yop. I.

hydride and hexahydride (W.). -5. By the action of ZnOl₂ xylene tetrahydride O_3H_{14} is produced: $O_6H_{14}(CO_2H)_2 = O_6H_{14} + CO_2 + OO + H_2O$.

The same hydrocarbon is produced by the action of ZnCl2 on ammonium emphoramic acid:

 $O_{10}H_{10}O_{10}O_{10}H_{1$ 324).-6. The anunonium salt distilled with P₂O₂ gives a terpene C₂₀Π₂₂ (Balko, A. 197, 329).
7. Conc. II.SO₄ forms CO and 'sulphocamphoric acid.'—8. Conc. IINO₃ gives camphoronic acid. -9. Potash-fusion gives pimelio acid Pr.CH(CO H).CH2.CO II and an acid C10H16O (Illasiwetz a. Crabowsky, A. 145, 205). 10. Distillation with soda-line gives camphoric anhydride and phorone C.H., O (Mayer, B. 3, 117). Distillation of camphorates gives similar results.

Salts.—(Igemper, Ar. Ph. [2] 110, 106; 117, 23). NH₄HA" xaq.—(NH₄),A".—Li₄A".—Na₄A".—K₄A": deliquescent.—MgA" 7½aq. S. 40 at 20°.—MgA"12aq.—MgA"13½aq.—CaH₂A"₂.—CaA'₄A"₄ aq. —CaA'₄A"₄, 8aq. —BaH.A'₂2aq.—BaA" aq: needles and feathers (Kingzett, C. J. 45, 93).—BaA" 4½aq.—ZnA".—PhA".—ChA".—AA". PbA". CnA".—Ag₂A".

Ethyl-ammonium salt (NII,Et),A": small needles (from alcohol). Convorted by PCl₅ into camphoric di-ethyl-imidino.

Mono-methyl ether MelIA". $[\alpha]_i = 51.4^{\circ}$. Trimetric prisms (from ether); gives camphoric anhydride when distilled. sl. sol. water (Loir, A. Ch. [3] 38, 483).

Mono-ethyl other EtHA". S.G. 201 1:095.

Di-ethyl ether Et, A". (286°). S.G. 161.029. Formed, together with camphorio anhydride, by the distillation of the mono-ethyl ether (Malaguti, A. Ch. [2] 64, 152; 70, 360; Meyer, B. 3, 118). Liquid. Cal., ga produces a tetra-chloro-derivative (C₁H₃Cl₂)₂A". S.G. 14 1:386.

Chloride C10 II14O2Cl2 Heavy oil, docomposing at 200° (Moitessier, A. 120, 252).

Anhydride C, H, O,. [217°]. (above 270°). S.G. 20' 1 194. $[a]_D = -7.7^\circ$ (in bonzene). Formed by heating camphoric acid or its salts (Bouillon-Lagrange, A. Ch. 23, 153; Laurent, A. Ch. 63, 207; Malaguti, A. Ch. 64, 151; Blumenau, A. 67, 119; Monoyer, J. Ph. [3] 45, 177). Formed also from camphoric acid by the action of (1 1.101. of) PCl. (Gorbardt a. Chiozza, A. 87, 294), r. couc. H. SO, of Ac.O., or of AcCle (Anschütz, B. 10, 1881). Long trimetric prisms (from alcohol) (Montgolfier, A. Ch. [5] 14, 5). V. sl. sol. water, v. sol. alcohol, v. e. sol. ethor. Slowly converted by boiling water into camphoric acid. The statement of Brodie (Pr. 9, 361; 12, 655) that parium peroxide formed camphoric peroxide has been denied by Kingzett (C. J. 45, 93).

Amide C₈H₁₁(CON11₂)₂. Amorphous mass (Morcessier, A. 120, 253).

Imide C₆H₁₄:C₂O₂:NH. [180°] (in sealed tubes, Ballo, A. 197, 332). Formed by heating ammonium camphoramate at 160° (Laurent, Compt. chim. 1845, 147; A. 60, 327). Lamins. May be distilled.

Ethylimide C.H.4:O.O.:NEt. [275°). Colourless orystals. Prepared by [50°]. tilling ethylamine camphorate (Wallach a.

Kamenski, B. 14, 164; A. 214, 248).

Allyl-imids C.H.,:C.O.:NO.H.. [49].

Formed by heating camphoric acid with allyl thiocarbimide. In ol. water, sol. alcohol and ether.

Phenyl-imide C, K.14: C,O,:NPh, Formed, together with phonyl-camphoramic acid, by warming camphoric anhydride with aniline (Gerhardt a. Laurent, A. 68, 35). Needles (from ether); insol. cold water.

Di-ethyl-imido-imidine C, H, N,O i.e. O=NC,II, C.H. NC.H. (286°), . S.G. 15 1.018.

Liquid; y. cl. sol. water. Pps. salts of Cu and Fe. Propared by the action of PCl, on ethylamine-camphorate or by the action of cthyl-.CCl₂ >NEt) obtained .CO

amine on the chloride (C.H.

from PCl, and camphoric-ethyl-imide. By HCl at 200° it is decomposed into ethylamine and eamphoric-ethyl-imide. Salts.—B'HCl: deliquescent orystals.—B'HI: sparingly soluble needles.—(B'HCl) PtCl. Its ethylo-iodide B'EtI: [245°]; forms long colourless prisms (Wallach a. Kamenski, B. 14, 162; A. 214, 242)

Nitrile CaH, (CN)2. Formed, in small quantity, together with hydrocarbons C, II, and C20 H32 by distilling ammonium camphoramate with P.O. (Ballo, A. 197, 334). Crystalline;

insol, water.

Camphoramic acid C₂II₁₁(CONH₂)(CO₂II). So-called 'amido-camphoric acid.' The aminonium salt is formed by the action of NII, on an alcoholic solution of camphoric anhydride Trimetric crystals; m. sol. hot water, v. sol. alcohol.—NH, A' aq. (200). When heated with dry ZaCl₂ it gives xylene tetrahydride and a terpene (Ballo, B. 12, 324).—PbA'₂.—AgA'. Phenyl-camphoramic acid

C₃H₁₄(CONPhH)(CO₂H). Formed by boiling the phenyl-imide of camphoric acid with alcoholic NH, or camphoric anhydride with aniliue. Needles (from alcohol); v. sl. sol. boiling water. -AgA' (Laurent a. Gerhardt, A. 68, 36).

Brome-campheric anhydride

C₈H₁₃Br<CO>O. From camphoric anhydride (10 g.) and Br (15 g.) at 130° (Woringer, A. 227, 3); an additive compound $C_{\nu}^{\prime}H_{14}O_{3}Br_{\mu}$ appears to be first formed (Wreden, A. 163, 330). metric crystals (from chloroform). a:b:c=
*8866:1: 5766. NII, gives the imide of oxya:b:c=camphoric acid.

(a)-0xy-camphoric acid O10H15O5. Formed, together with pimelic acid, by fusing camphor with KOH (Hlasiwetz a. Grabowski, A. 145, 212). Thick liquid; the salts are amorphous.

(β)-0xy-camph ric acid C10H16O4 Anhydride C., H.O. i.e. C., H. (CH): C.O.: O (?) Camphanic acid. [201°].

Formed by boiling bromo-camphoric anhydrido with water (Kachler, A. 162, 264). Monoclinic prisms (containing aq or 2aq) (Grünling, A.227, A). Sublimes at 110°. Decomposed on distillation giving CO₂, lauronolic acid C₂H₁₁O₂, and pampho-lactons C₂H₁₄O₂. Water at 180°

splits it up into CO₂ and C₃H₁₄ (119°).— Ba(C₁₀H₁₄O₄)₂ 1³₂aq.—Cd(C₁₀H₁₅O₄)₂ ³₂aq. Ethyl ether EtC₁₀H₁₅O₅. [63°]. From brome-camphoric anhydride and alcohol at 150°.

Prisms.

Imide $C_{10}H_{15}NO_{0}$ i.e. $C_{n}H_{10}(OH):C_{2}O_{2}:NH$ (?) [208°]. Amido camphoric anhydride. From bromo-camphoric anhydride and cone. NH,Aq at 150° (Wreden, A. 163, 339). Long needles (from alcohol). Sublimes at 150°. Converted by nitrous acid into the anhydride.

Oxy-camphoramic acid C₁₀H₁₀NO₄ i.e. C₂H₁₂(OH)(GONH₂)(CO₂H). 'Amido-camphoric acid.' [160']. Formed by boiling the imide of oxy-camphoric acid with dilute KOH (Wreden, A. 163, 540). Prisms containing aq (from alcolol). On fusion it is converted into the parent imide. Converted into oxy-camphoric anhydride by HNO2, by conc. HClAq, or by H2SO4. -

CaA' 2aq.
Sulpho - camphoric acid' so - called. [160°-165°]. Sulphocamphylic CoH10SO. acid. Formed, together with CO, by heating camphorio acid or anhydride with cone. ILSO, (Walter, A. Ch. [3] 9, 177; Kachler, A. 169, 179). Triclinic prisms (containing 2aq); a:b:c=8515:1:7590; $\alpha=82^\circ$ 39'; $\beta=121^\circ$ 10'; $\gamma=111^\circ$ 36' (Zepharovich, Sitz. B. 73, 7). V. e. sol. water, alcohol, and ether. IINO₂ (S.G. 1·25) converts it into C,II₁₂SO₂. Potash-fusion gives C,H₁₂O₂ [148], insol. cold water, but separating from alcohol in monoclinic crystals.

Salts. — (NII.) _A" aq. — K.A". — CaA". — BaA". — PbH_A". _4aq: trimetric. — PbA". — BaCuA". _ Ag.A".

ISOMERIDES OF CAMPHORIC ACID.

Levo rotatory camphoric acid $C_{10}H_{16}O_4$. [186°]. [α]_j = -46.3° (in alcohol). Formed by the oxidation of lawo-rotatory ([a]₁ = -38°) borneoper the corresponding camphor, by heating for soveral hours with a large excess of 11.70₃ (S.G. 1-27) (Chautard, C. R. 37, 166; Haller, C. R. 103, 64). Resembles camphoric acid in all respects oxcept that its rotation though equal is opposite.

Inactive campheric acid $(C_{10}\Pi_{16}O_4)_2$. [204°]. S. 1; S. (alcohol) 33; S. (ether) 28 (C.). Formed by heating inactive camphor with HNO, at 100°, or by mixing equal weights of dextroand levo-camphoric acids (Chautard, C. R. 56. 698; Armstrong a. Tilden, C. J. 35, 757.
Haller, C. R. 105, 66). Less soluble than its

Diethyl ether Et.A". (270°-275°). S.G. 15 ko3 (C.).

isomerides.

Anhy dride $(O_{10}H_{14}O_3)_2$. [223°] (A. a. T.). (chloroform) 25; S. (ether) 4; S. (alcohol) 1.5 (C.). Formed by hoating the acid.

An inactive campheric acid [186°] was obtained by Muir (C. J. 37, 685) by oxidising the camphor of oil of sage

Mesc-camphoric soid C₁₀H₁₈O₄. [113°]. Formation.—1. Formed by heating dextrooamphoria acid (5 g.) with (20 c.c.) fuming HGl at 140° for 30 hours (Wreden, Z. [2] 7, 419; A. 163, 328; B. 6, 565).—2. By heating dextrocamphoric acid (5 g.) with HI (30 c.c. of S.G. 1·6) at 160°.—3. By heating camphoric acid with water at 200° (Jungfleisch, B. 6, 268, 680). 4. Together with 'sulphocamphoric acid' by the action of H₂SO₄ on camphor. Occurs also in small quantity in the preparation of camphoric sold from camphor and HNO₂ (Kachler, A. 169,

179; 191, 146).

Properties.—Crystalline, but separates from alcohol and ether as an oil. More soluble than ordinary camphoric acid. Conc. H.SO, converts it on warming into 'sulphocamphoric acid.' When heated it gives the anhydride of ordinary camphorio acid. Boiling dilute HCl changes it into inactive camphoric acid.

CAMPHORIDE C16H12O6. [222°]. Occurs together with galangin and alpinin in the galanga root (Alpinia officinarum). Flat yellow needles (containing aq), sublimable. Sol. hot alcohol, ether, and acctic acid, sl. sol. chloroform and benzeno, msol. water. Dissolves in alkalis. On oxidation with diluto HNO, it produces anisio and oxalic acids.

Salts .- A"Pb: yellow amorphous pp .-A"Pb₂O.—A"Ba 2aq: orango pp.

Di-acelyl derivative C1. H1. O1 (OAc)... [189°]. Colourless crystals. Insol. water. Si. sol. alcohol.

Di-benzoyl derivative C, H10O4(OBz). [186°]. Fino white needles. Scarcely sol. alcohol, insol. water.

Di-bromo-derivative C16H10Br.O6. [225°]. Yellow needles. Sl. sol. alcohol (Jahns, B. 14,

. Characteristates $C_{14}H_{21}NO_{4}$ i.e. $C_{8}H_{11} < C_{CO}^{CO} > N.Cll_{2}.CO_{2}Et.$ [86°]. Formed by adding a solution of CH2Cl.CO2Et to a solution of sodio-eamphorimide in absoluto alcohol. Large transparent crystals (from alcohol), sol. ether (Haller a. Arth, C. R. 105,

CAMPHOR OIL. An oil oblained, together with camphor, by distilling the wood of Laurus camphora with water. It con. 's chiefly of camphorogenel' but contains also several

terpenes (q. v.).

Camphorogenel $C_{10}H_{16}O_2$ or $C_{10}H_{18}O_2$ or $C_{10}H_{18}O_2$ or $C_{10}H_{16}O_3$ or $C_{10}H_{16}O_$ it forming a small quantity of camphor. CrO, acts similarly. Excess of conc. IINO, gives camphoric acid [185°], [a]j - 40.3°. Ac.O has no action. Sodium roduces it, in alcoholic solution, to borneol, [1987], (2127), [a] = 22.97. With ZnCl₂ it yiolds cymene (Yoshida, C. J. 47, 785; Oishi, C. N. 50, 275; Wallach, A. 227, 296; Lallemand, A. Ch. [3] 57, 404).

CAMPHOR-PHORONE v. PHORONE.

CAMPHORONIC ACID C, II, O. Mol. w.

218. [137°] Formed by oxidising campholic or camphagic acids (Bredt, B. 18, 2989).

Preparation.-From camphor and IINO. Present in the mother liquor from which camiphoronic acid has separated. Obtained by means of the barium salt (Kachler) and purified by de-composing this with HCl, extracting with ether, boiling off the ether, dissolving in water, neutralising with lime and boiling. The pure calcium salt then separates (Bredt, A. 226, 251; cf. Kachler, B. 7, 1728; A. 159, 286; 162, 262; Kachler a. Spitzer, M. 6, 173).

Properties. - Crystalline aggregates of needles, v. e. sol. water, alcohol, and wet ether, sl. sol. butyric acid and the anhydride C.H.2O. [135] which forms trimetric crystals, a.b.c = 96:1: 82; sol. water, alcohol and ether. This anhydride forms with NH, the compound C.H. (NH,)O. [c. 128°], v. sol. water

Reactions.-1. AcCl gives CallingOn and then the anhydride C18H22O8 [176°]; crystals, insol. cold alcohol and other; reconverted into camphoronic acid by boiling alkalis. -2. Br at 130° givos oxy-camphoronic acid .- 3. Potash-fusion gives isobutyrio acid.— 4. Aqua-regia forms two

gives isobutyrio acid.—4. Aqua-regia forms two acids C₉II₁₂O₅.—5. KMnO₁ gives HOAc and an acid C₁II₁₂O₄. (Kachler, M. 5, 415).—6. The Casalt distilled with lime gives a ketone C₁H₁₂O.

Salts.—NI₁Il₂A": [128'].—(NII₁)₂HA": [148'].—K₂IIO" aq.—C²₁A", 12'aq.—BaHA" aq: m.sol.water.—Ba₃A", 2'aq.—Ba₃A", 2'aq.—Sa₄A", 2'aq.—CliHA" 6aq.—Pb A', 2'aq.—Cn₃A", 2aq.—CliHA" 2, 2aq.—Cu₃A', 2. Ag, A", —Ag, HA" aq.

Mono-ethyl ether.—The anhydride (302') (or anhydrides liquid and solid (67'). Hight

(or anhydrides liquid and solid [67°], Hjelt, B. 13, 797) CollinEtO, of this ether is formed together with alcohol by distilling the diethyl ether.

Di-cthyl cther Et.HA". From the acid. alcohol, and HCl.

Tri-cthyl ether Et, A". (302°). From

Ag₃A''' and Etf. Liquid. Chloride C₃H₁₁OCl. [I31°]. Needles, sl. sol. water, sol. alcohol and other.

Mono-amic acid CoH11 (CONH2) (CO2H); Anhydride CaH13NO. [212]. From liquid mono-cthyl camphoronate and alcoholic NH. (Hjelt, B. 13, 798). By the same treatment the solid ethyl camphoronate gives a compound Callia N.O. (? di-amio acid) crystallising with HOEt. It melts at [115], and is converted by boiling HClAq into camphoronic acid.

Diamic ac. d C₀H₁₁N₁O₁ t.c. C₀H₁₁(CONH₂)₂(1) t.2. [c. 160°]. From diethyl camphoronato and NH₁ at 120° (H.). HClAq converts it into a compound CoH13NO [212°].

Constitution .- Camphoronic acid appears to contain 3 carboxyls: C₀ll₁₁(CO₂H)₃, as shown by the salts and others. AcCl gives no acetyl derivative. The formation of an anhydride by distilling the cther does not prove it to be lactonic. Potash-fusion produces iso-butyric acid, hence it contains isopropyl. Since it does not split off CO, on distillation, the carboxyls must be attached to different carbon atoms. Hence

it is is Progyl-tricarballylic acid, Cll_(CO_II).CPr(CO_II).CH_r.CO_H or CH_2(CO_II).CHICO_II).CHIPr.CO_H.

Oxy - camphoronic acid Call 20. [c. 210] Formed by heating can phoronic acid CoH14O (I mol.) with Br (I n.ol.) for two hours at 130° (Kachler, A. 152, 296). Monocliuic crystals (containing a 1), a:b:c=1.1918:1:9808; $\beta=86^{\circ}50'$. According to Zepharovich (J. 1877, 641) they aro dimorphous. V. sol. water, alcohol, and bether; may be distilled.

Salts.-KIIA" aq . crystals.- K2A": gummy. -BaA" aq: pearly plates. -I'b, (C, H,O,), 2aq. -Ag₂A".

Hydrc-exy-campherenic acid v. Самрнов. CAMPHOROXIM C1011, NO i.e. C10H16: N.OH. [115°]. (c. 250°). Formed by the action of pure ether. Produces on distillation CO, iso hydroxylamiue on camphor (Nägeli, B. 16, 498). Long needles. Smells like camphor and rotates on water. Sol. alcohol, ether, acids and alkalis.

Reactions .- 1. Hydroxylamine is not split off by heating with aqueoue HOl even at 120°.—2. By heating with acetyl chloride it loses H2O yielding the nitrile of (campholenic acid C1. H1. N (Goldschmidt a. Zürrer, B. 17, 2069).—3. Is reduced in alcoholic colution by metallic Godium

CH.NII. to bornylamine CaH,

anhydride is reduced to the isomeric camphylamine (Leuchart a. Bach, B. 20, 111)

Hydrochloride C10H16NOII, HCl: white powder, sl. sol. water, v. sol. alcohol and acids.

Sodium salt.—C10II,8N(ONa): white powder, v. sol. ot ater and hot alcohol.

Ethylether Ciellia N(OEt): (209°); mebile

Anhydride C,0 II,N: (217°), liquid; formed by heating camphoroxim with acctyl chloride. Is the nitrile of Campholenic acid (q. v.) (Nägeli, B. 16, 2981).

Isocamphor-exim is the amide of Campno-

LENIO ACID (q. v.).

CAMPHO-TERPENE v. TERPENES.

CAMPHRENE v. PHORONE.

CAMPHRESIC ACID or CAMPHRETIC ACID so called by Schwanert (A. 128, 77) has been shown by Kachler (A. 191, 143) to be a mixture of camphoric and camphoronic acids.

CAMPHYLAMINE C10H19N possibly C₈H₁₃(CH₂NH₂):CH₂ (195°). Prepared by adding metallic sodium to an alceholic solution of campholenic nitrile (camphor-oxim-anhydride). Colourless liquid. Volatile with steam. Readily absorbs CO, from the air and solidifies to a crystalline carbonate. Primary base.

Salte.-B',11,Cl,PtCl,: glistening golden plates, nearly insol. water.—B'HCHIgCl₂: colourless orthorhombic plates; sol. hot water.—B'C₂H₂O₄; aq: [194°], Göörrless orthorhombic glistening crystals, y, sc. hot water.—B'₂H₂O₄q: long rhombic prisms, m col. cold water.—B'₂H₂Cr₂O₇; orange-red plates.—The pierate forms fine vollow needles [140°-104°] pierate forms fine yellow needles, [190"-1940].

Benzoyl derivative C₁₀II₁₈NBz: [770],

colourless prisms (Goldschmidt a. Schulhof, B.

18, 3297; 19, 708; 20, 483).

Isomeride v. Bornylamine CAMPHYL-PHENYL-THIO-UREA

SC(NHC₆H₃)(NHC₁₀H₁₇). [118°]. Formed by combination of phenyl-thiocarbimide and camphylamine. Short colour ss prism. I. V. sol. alcohol and benzene, el. sol. ether, v. eol. ligroin (Goldschmidt a. Schulhof, B. 19, 712)

CAMPHYL - DI - THIO - CARBAMIC ACID C₁₀H₁₇.NH.CS.SH. The camphylamine salt, C₁₀H₁₁, NH.CS.S '11' (C₁₀H₁₇), is formed by mixing camphylamine with CS₂. White powder, [110°-116°], sol. benzene. The sodium ealt C10H17.NH.CS.SNa forms white glistening platee, sol. cold, decomposed by hot, water

(Goldschmidt, B. 19, *12).
CANADA BALSAM Exudes from incisione in the bark of Abies balsamea. Transparent thick liquid with refractive index (1.532) nearly the came ac that of crown glase. Dextrorotatory. Steam-distillation separatee a lavorotatory terpene (167°), which forms a crystalthe compound with HCl (Donastre, J. Ph.

8, 572; Caillot, J. Ph. 16, 436; Wirzen, Dissertation, Helsingfore, 1849).

CANADOL. A term applied by Vohl (D. P. J. 172, 319) to that portion of the volatile hydrocarbons of Canadian and Penneylvanian petroleum which boile at 60° and has a S.G. 65 to ·70. It is also called petroleum-ether or ligroïn. It consists chiefly of n-hexane.

CANANOA OIL. Alan-gilan. From Can-

anga odorata. Neatxal oil (170°-290°). It contains benzoyl and acetyl derivatives, a compound that unites with NaHSO,, and probably a phenol

(Flückiger, Ph. [3] 11, 934).
CANARIUM. The fixed oil of Canarium commune contains 51 p.c. clein and 49 p.c. etearin and myristin (Oudemans, J. pr. 99, 407).

CANAÜBA WAX v. CARNAÜBA WAX.

CANE SUGAR v. SUGAR. CANNABIS INDICA. Indian hemp when distilled with steam yielde an essential oil C_{1,1}H_{2,1} (257°); V.D. 7·1; S.G. § 93; [a]_p = 10·81 at 25·5° (in chloroform). The oil resinifies on exposure (Valenta, G. 10, 479; 11, 196; for Martius, C. C. 1856, 225; Personne, J. Ph. [3] 31, 46). HNO, (S.G. 1.32 to 1.42) acting on the resinous extract of Indian hemp forms 'oxycannahene' C₂₀11₂₀N₂O₄ (Bolas a. Francie, C. J. 22, 417; C. N. 24, 77). This separates from methylated spirit in flat yellow prisme [176°], insol. water, sl. sol. alcohol. Indian hemp, and ite alcoholic extract, contain a poisonous resin (T. a. H. Smith, Ph. 6, 127, 171; Martius). Hay (Ph. [3] 13, 998) has extracted a crystalline alkaloid 'tetano-cannabine' which produces tetanus in frogs. The fixed oil from hemp-seed (Cannabis sativa) is probably a fatty cil, though Lefort (C. R. 35, 734) gives it the formula $C_{11}H_{22}O_2$ and describes $C_{11}H_{20}Cl_2O_2$ and C₁₁H₂₂O₂ and describes C₁₁H₂₀Cl₂(C₁₁H₂₀Dr₂O₂ as preducts of substitution.

CANNON-METAL v. Copper, alloye of. CANTHAREME C.H., i.e. C.H.(CH.), [1:2].

(134°). o Lylene-di-hydride.

Formation .- 1. By heating cantharic acid with fused KOH.—2. By heating cantharic acid or cantharidin with water at 300°, CO, being split off.—3. In a pure state by boiling with cone. aqueous KOH, the product C₁₀H₁₂O₃I₂, obtained together with cantharic acid by the action of Ill upon cantharldine (Piccard, B. 12, 577; 19, 1404).

Properties.—Liquid, smelling like turpentine and camphor. Absorbs oxygen with avidity. Dilute HNC, oxidises it to o-toluic and phthalic

acids.

CANTHARIC ACID C₁₀H₁₂O₄ i.e. (C,H₁₁O)CO.CO₂11. [278° cor.]. S. 85 at 15°; 8.5° at 150°. Propared by heating 1 pt. of cantharidine with 4 pts. of III (1.96 S.G.) for 2½ hours et 100°. Trimetric crystals from water); v. e. sol. alcohol, v. sl. sol. ether. Distilled with lime it gives cantharene, a little xylene, butyric acid, and di-methyl-benzoic acid. It is an a-ketonic acid, for on heating with di-methylaniline and ZnC! it evolves CO, and yielde a condensation product C₂₂H₃₂ON₂; the latter is converted into a green colouring-matter by MnO2, into a violet colouring-matter by chloranil or arsenic acid. -A'Ag: white pp.-KA': elender needles .- PbA'2xaq.

Methyl ether A'Me: (210°-220°) at 50

mm.; colourless liquid.

Bthyl ether A'Et (c. 800°). Oxim C₁=12O₂(NOH): [175°-180°]; colourless four-sided plates (Piccard, B. 10, 1504; 11, 2121; Homolka, B. •19, 1086).

CANTHARIDIC ACID C10H14O, i.e. (C3H14O2).CO.CO3H. The alkaline salts are formed by heating cantharidin with aqueous alkalis. When a cold solution of the salts is treated with acids, the free cantharidio acid appears to be formed, but on warming the solntion it loses H2O and cantharidin is pro-With hydroxylamine it gives an oxim, from the salts of which acids liberate the oxim of cantharidine.—Ag.A."aq.—Ag.A."2aq. (NH.).A."aq.—K.CuA", 2aq. (NH.).A."aq.—K.CuA", 2aq. Di-methylether A."Mc.: [91°]; large flat glistening prisms; sol. alcohol, ethor, and hot

water, sl. sol. cold water (Homolka, B. 19, 1082; Dragendorff a. Masing, Z. 1867, 464; Masing,

J. 1872, 841).

CANTHARIDIN C₁₀H₁₂O₄. Lactone of cantharidic acid. [218° gor.]. S. 02 at 15°; 29° at 100°; S. (alcohol) 2·1 at 78°; ·13 at 15°; S. (benzene) 3·38 at 80°, ·51 at 15° (llennard); S. (ether) ·11 at 18°; S. (CS₂) ·06 at 18°; S. (COC) ·12 at 18°; S.

(CHCl.) 1.2 at 18° (Bluhm).

Occurrence.-In Spanish flies (Lytta vesicatoria) and many other insects (Thierry, A. 15, 815; J. Ph. 21, 44; Robiquet, A. Ch. 76, 302; Gössinann, A. 86, 317; Pocklington, Ph. [3] 3, 681; Regnault, A. Ch. [2] 68, 159; Warner, Am. J. Ph. 28, 193; Ferror, J. 1860, 597; Mortreux, J. Ph. [3] 46, 33; Fumouze, J. Ph. [4] 6, 161; Bluhm, Z. [2] 1, 675; Dragendorif, Z. [2] 3, 187, 464; 4, 308; Rennard, C. C. 1872, 568; Wolff, Ar. Ph. [3] 10, 22; Piccard, B. 10, 1504).

Preparation .- 1. Powdered cantharides are extracted with chloroform or ether, the solvent is evaperated and the residue freed from fat by washing with $CS_2 - 2$. Cantharides are mixed with water and MgO, dried, treated with dilute H2SO, and then shakon with other.

Properties .- Trimetrie plates. Blisters the

skin. Sublimes readily at 85° (Blyth).

Reactions .- 1. HI forms cantharic acid. --2. By distillation with P.S. it gives o-xyleno (Piccard, B. 12, 580).—3. By heating with alkalis it is converted into salts of cantharidio aoid C10H14O5, from whose hot solutions cantharidin is re-precipitated on the addition of acids.

Oxim C₁₀H₁₂O₂(NOH): [166°]; splendid long glistening prisms; v. o. sol. alcohol and ether, v. sol. hot water, sl. sol. cold. By conc. IICl at 150° it is split up into its constitution. ents.—C₁₀H₁₀O₄(NOAg): four-sided prisms.— C₁₀H₁₂O₄(NOMb): [134°]; large colourless prisms; v. 6, sol. alcohol and ether, v. sol. hot, sl. sol. cold, water (Homolka, B. 19, 1082).

Compound C10H12O3I2. 'Cantharidin iodide' is formed as a by-product (5-8 p.c.) in the preparation of cantharic acid by the action of HI (196 S.G.) upon cantharidin at 85°. Crystalline solid. V.sol. benzone and chloroform, sl. sol. alcohol, insol. water. On boiling with cono. KOH it is converted into pure cantharene (o-xylene-di-hydride) C₈H₆(CH₄)₂ (Piccard, B. 19,

various trees belonging to several natural orders. The sap, which is obtained by making an incision in the bark of the tree, is a white oreamy

liquid with a sp. gr. 1012.

The caoutchouc exists in the sap in the form of minute globules, and is consolidated in various ways, then by heating over a smoky fire which produces the brown colour of the commercial

·Caoutchoue is colourless when puro, it is a bad conductor of heat and a non-conductor of electricity. S.G. about 925. At ordinary temperatures it is soft, flexible, and very elastic, but at about 10' it begins to lose its elasticity, and at 0° becom@hard and rigid. When heated it loses it: clasticity and becomes soft, slowly resuming its original properties when cooled; if heated to 150°-200° it melts, and after this it remains semi-liquid and sticky on cooling. burns readily with a smoky flame, leaving little or no ash.

Exposure to air in the absence of light produces little effect on caoutchouc, but light and air together causo it to loso its clusticity and become glutinous, due to the absorption of oxygen (Spiller, C. J. 18, 44; Miller, ibid. p. 273).

Caoutchouc is insoluble in water, but when immersod in it becomes white and increases in bulk, absorbing about 25 p.c. of its weight of water, which is given up again on exposure to air. Alcohol acts upon it in a similar way.

Diluto acids do not affect it, but it is attacked by strong nitric or sulphurio acid. Chlorine renders it hard and brittle. Alkalis' produce

little offect.

Ether, benzene, mineral oil, sulphide of carbon, chloroform, oil of turpentine, oil of caontelioue, and many essential and fixed oils, act upon cacatelous, causing it to swell greatly and become gelatinous and soft. The action of these solvents appears to be to dissolve one constituent part of the enoutehoue, leaving the less soluble part in a disintegrated condition.

Accoding to Payen, sulphido of carbon with about 5 p.c. of absolute alcohol is the best

solvent.

Caoutchouo is composed of carbon and hydrogen. The proportions vary in different analyses C. 86:1-90:61.c.; II. 10-12:8 p.c. It appears to consist chiefly of two hydrocarbons, which can be partly separated by the prolonged action of a solvent, but the proportion of these constituents obtained varies according to the solvent employed. The more soluble part is soft and ductile, while the less soluble is tenacions and elastic.

When caoutchouc & subjected to dry distillation an oil consisting of a mixture of various hydrocarbons is obtained. This is

called oil of caovielione.

Among the constituents of this oil are isopene C₁H_e (37°-38°) S.G. 682; caoutchene C₁₀H₁₆ (171°) S.G. 842; and hevene (315°) S.G. 921 (Hinly, A. Ch. 27, 41; Gregory, bid. 16, 61; G. Williams, Pr. 10, 517; Bouchardat, J. Ph. 1837, 454; Bl. 24, 108; C. R. 89, 361).

When isopreno is actod on by strong hydric 1404).

CAOUTCHOUC. India rubber. This subchloride a mixture of the mono and di hydrachlorides, together with a solid substance, is
stance is obtained from the milky sap of ties with caoutchous (Bouchardat, C. R. 89.

Vulcanised caoutchous.—When caoutchous is heated to about 115° in contact with sulphur, it absorbs some of the catter and becomes vulcanised. The introduction of the sulphar can be attained in many ways, immersion in a mixture of carbon disulphide and chloride of snlphur, of in a solution of polysulphide of calcium, &c.

About 2 p.c. of sulphur appears to enter into combination with the caoutehouc. If more than this quantity is introduced the excess remains mixed with the rubber and can be dissolved out by the ordinary solve its of sulphur, while the combined sulphur cannot be so extracted. An excess of sulphur renders the caoutcheuo less durable. Vulcanised caoutchouo does not lose its elasticity at a low Comperature and does not soften so easily with heat as ordinary rubber. It is less affected by solvents than pure caoutcheuo.

The ordinary vulcanised rubber, besides containing an excess of sulphur, is often adulterated

with 40-60 p.e. of mineral matter.

Ebenite.—When caoutchoue is heated with half its weight of sulphur, with or without the addition of some mineral matter, a hard dark substance which can be polished is obtained. This is much used for insulating purposes, but according to Wright (Am. S. [3] 4, 29) it becomes hygroscopic when exposed to the action of ocone owing to the formation of H.SO. Ebonite is little affected by the solvents of caoutchouc.

CAPILLARITY v. PHYSICAL METHODS, Sect. MISCELLANEOUS

CAPRAMIDE the Amide of Decord Acid (q.v.). The name has also been applied to the amides of Octoro ACID (q.v.) and Handic Acid (q.v.).

CAPRAMIDOXIM v. HEXAMIDOXIM. CAPRIC ACID v. DESOIC ACID.

CAPRIC ALDEHYDE v. DECOIC ALDEHYDE. CAPRILAMIDE v. Amide of Octoic ACID.

CAPRILIC ACID v. Octolo acid.

CAPRILONE v. DI-HEPTYL-RETONE. CAPRILONITRILE v. Nitrile of Octoic ACID.

CAPRINONE v. DI-KNNYL-KETONE. CAPRO-AMIDE v. Amide of HEXOIC ACID.

CAPRO-ANILIDE v. Anilide of Hexoic acid. CAPROIC ACID v. HEXOIC ACID.

CAPROIC ALDEHYDE v. HEXEIC ALDEHYDE. CAPRO-LACTONE v. Lactone of Oxy-nexole ACID

CAPRONE v. DI-AMYL-RETONE.

CAPRONITRILE v. Nitrile of HEXOIC ACID. CAPROYL = HEXOX.

CAPROYL AMIDE v. Amide of Hexore acid. CAPROYL CHLORIDE v. Chloride of Hexoic

CAPRYL ALCOHOL v. OCTYL ALCOHOL.

CAPRYL-AMIDE v. Amide of Octoic Acid. CAPRYLAMINE v. OCTYLAMINE.

CAPRYL CHLORIDE v. Chloride of Decoro ACID; also OCTYL CHLORIDE.

CAPRYL-BENZENE v. OCTYL-DENZENE.

CAPRYLENE v. OCTYLENE.

CAPRYLENE HYDRATE v. OCTYL ALCOHOL. CAPRYLIC ACID v. OCTOIC ACID.

CAPRYLIC ALCOHOL v. OCTYL ALCOHOL. CAPRYLIC ALDEHYDE v. OCTOIO ALDEHYDE. CAPRYLIDENE v. OCTINENE.

Caprylidene tetrabromide v. TETRA-BROMO-COTANE

CAPRYLONE v. DI-HEPTYL-RETONE.

CAPRYLONITRILE v. Nitrile of Octoic Acid. CAPRYL-PHENYL-AMINE v. p-AMIDO

PHENYL OCTANE, p. 178.

CAPSAICIN C. H. 10. [59°].

Preparation. — Nowdered cayenne pepper (Capsicum fastigiatum) is extracted with ether, the extract is evaporated, dissolved in het alcoholic KOH, diluted with water, ppd. by BaCl,, and the dried pp. treated with ether. On evaporating the extract, an oily red liquid remains, which is disselved in diluto potash, and ppd. by addition of ammonium chloride.

Properties.—Colourless prismatic crystals, insol. water, sol. alcohol. Begins to volatilise at 100°. Powerful irritant. The pungent taste is removed by heating with potassium biohremate and diluto sulphuric acid. BaCl, and CaCl₂ in alcohelic solution give a pp. sol. other; AgNO₃ a pp. sol. ammonie; Fe₂Cl₃ a red pp. when warned (Thresh, Ph. [3] 7, 21, 259, 473).

CAPSICINE. An alkaloid which may be extracted by benzeno from the fruit of Capsicum fastigiatum. The benzene is evaporated, and the residue dissolved in ether, from which the alkaloid is obtained by shaking with dilute ILSO, (Thresh, Ph. [3] 6, 941). Needles; insol. water, v. sol. alcohol and other; may be sublimed. Volatile with steam. It is not pun-The hydrochloride crystallises in oubes and tetrahedra, the sulphate in prisms.

CAPSULÆSCIC ACID C13H12O8. An aoid obtained from the husks of the horse-chestnut (Rochleder, Z. 1867, 83). Crystals; may bo sublinied. Fe'Cl, turns its solution greenish-

blue

CARAGHEEN MOSS. Irish pearl moss. gelatinous Scaweed (Chondres crispus) & Swells up in cold water, almost entirely dissolves in hot water. Ppd. by Pb(OAc)₂. Appears to be chiefly composed of a carbohydrate, which is insel. Schweizer's solution, and not turned blue by H₂SO₄ and I (Schmidt, A. 51, 56; Flückiger a. Obermayer, N. R. P. 1868, 350). Caragheen moss gives galactose when boiled with dilute H.SO. (Haedicke, Bauer, a. Tollens, A. 238, 302).

CARAJURA. A red dye, probably identical with chica red. Insol. water, sol. alcohol and dilute alkalis, reppd. by acids (Viroy, J. Ph.

1844, 151).

CARAMEL. A black substance obtained by heating cane-sugar at c. 200°. It is said to be a mixture of caramelan $C_{12}H_{18}O_{2}$, caramelen $C_{36}H_{36}O_{23}$, and caramelin $C_{96}H_{192}O_{51}$. They all reduce Felling's solution. Dilute (84 p.c.) alcohol extracts caramclan, cold water then dissolves caramelen, leaving caramelin. Caramelan is carameten, teaving carametin. Carametan is a colourless, brittle, deliquescent resin.—
C₁₂BaH₁₆O₃BaO.—C₁₂IbH₁₆O₅.—C₁₂PbH₁₆O₅PbO.
Carameten is a mahogany-coloured solid.—
C_{3c}H₄₆BaO₅.—C_{3c}H₄₆PbO₂₅. Carametin is a glittering black solid, sol. boiling water.—
C_{3c}H₁₀₉BaO₅₁.—C_{3c}H₁₀₉BaO₅₁BaO.—C_{3c}H₁₀₉PbO₅₁
(Gélis, A. Ch. [3] 52, 352). Corametan and carameten are crystallides carametin is oaramelen are crystalloids, caramelin is a colloid. The formulæ and purity of these bodies are, of course, very doubtful; other

observers have arrived at somewhat different results, indeed the nature of the resulting products depends upon the temperature used in preparing them (Péligot, A. Ch. [2] 67, 172; Völckel, A. 85, 59; Maumene, C. R. 39, 422; Graham, C. J. 16, 258; Thomson a. Sherlock, C. N. 85, 242, 263. C. N. 25, 242, 282).

CARAWAY OIL. Oil of caraway contains a terpene (q. v.) C₁₀H₁₀ identical with citrene, and

carvol (q. v.) C₁₀H₁₄O.

CARB-ACETO-ACETIC ETHER is mesitence-

lactone carboxylic acid, p. 20.
CARBACETOXYLIC ACID C.JI.O. A syrupy acid, said to be formed by the action of moist Ag2O on β-chloro-propionic acid and on aα-dichloro-propionie ether. Reduced by sodiumamalgam to glycerie acid, and by HII to pyruvic acid (Wiehelhaus, A. 143, 7; 144, 351; Klimenko, B. 3, 468; 5, 477; 7, 1406; cf. Beckurts a. Otto, B. 10, 2039).

TRI-CARBALLYLIC ACID C.H.O. i.e. CO2H.CH(CH2.CO2H)2. s-Propane tri-carboxylic

 acid. Mol. w. 176. [158]. S. 40.5 at 14.
 Formation.—1. In the preparation of sugar from beet-root (Lippmann, B. 11, 707; 12, 1619; Weyer, C. J. 38, 864) .- 2. By the saponification of its nitrile which is prepared by the action of KCy on s-tri-bromo-propano in alcohol (M. Simpson, Pr. 12, 236; 14, 77; C. J. 18, 331). 3. By reducing aconitie acid or its ether with sodium-amalgam (Dessaignes, C. R. 55, 510; Wichelhaus, A. 132, 61; Hlasiwetz, Z. 1861, 734). -4. By the action of potash on the product of the action of KCy on \$-chloro-isocrotonic ether (obtained from aceto-acetic ether und PCl,) (Claus a. Lischke, B. 14, 1089).-5. In the same way from a chloro-crotonic ether or from di chloro propylene (phichlorhydrin) (Claus, B. 5, 358; 9, 223; A. 170, 131; 191, 63).—6. Appears to be formal by the action of HC and KClO₃ on gallic acid Chreder, A. of HC and KClO, on gallic acid "chreder, A. 177, 292).—7. Acetyl-succinic other is converted by Na into acctyl-tri-carballyho other CH.CO.C(CH..CO.Et)...CO.Et whence alcoholic KOH or baryta-water produce tri-carballylic acid (Miehle, A. 190, 322).—8. By the oxidation of di-allyl-acetic acid by dilute HNO₃ (Wolff, A. 201, 53) .- 9. By boiling citraconic acid with zinc and HCl (Beltmanu a. Hofmann, B. 17, 2692).—10. From propane tetra-carboxylic acid (CO₂H.CH₂)₂C(CO₂H)₂ by heat (Bischoff, A. 214, Ċ6).

Properties .- Hard short trimetric prisms (from water); v. sol. water and alcohel, st. sol. ether. The ammonium salt gives with BaCl, or CaCl₂ no pp., even on adding NII₃. • 'b(OAc)₂

or CaCl₂ no pp., even on adding NII., a b(OAC), gives a whito pp. Fe_Cl_a gives a red pp.

Salts. — Na,HA''' 2aq(?) — Ki_A'''. —
Ca,A'''_2 4aq.—BaHA'''.—Ba,A'''_2 6aq.—Pb,A'''_2—
Cu,A'''_2.—Ag,A'''.

Tri-ethyl ether Et,A''' (c. 300°).

Tri-isoamylether(C,H₁₁),A''' (above 360°).

CARBAMIC ACID CH,NO₂ i.e. 'iI_.CO_II.

Anido-formic acid. Anide of carbonic acid.

Not known in the free stato. The am mon rum alt is formed by the union of dry or moist salt is formed by the union of dry or moist CO₂ (1 vol.) with gaseous NH₂ (2 vols.) (J. Davy, N. Ed. P. J. 16, 345; Rose, P. 46, 352; 2l. 30, 47). Formed also by sublimation of neutral ammonium carbonate, and therefore occurs in commercial ammonium carbonate. Formed also

by oxidising glycocoll, leucine, syrosine, and albumen, with alkaline KMnO, (Drechsel, J pe. [2] 12, 417; cf. Hofmeister, J. pr. [2] 14, 173). It may be conveniently prepared by digesting commorcial ammonium cerbonato with saturated aqueous NII, for 30 or 40 hours at 20°-25° (Divers, C. J. 23, 215; cf. Kolbe a. Basaroff, C. J. 11, 194).

Reactions. -1. Acids decempose carbamates with formation of CO, and NII, -2. Boiling unter converts carbamates into carbonates .-3. Strongly heating converts the Na salt into sodium cyanate and H.O (Drechsel, J. pr. [2] 16.

199).

Salts.—The carbamates are soluble in water (difference from most carbonates). -NH.A' (v. supra). Deliquescent plates. Its aqueous solution quickly changes to carbonate, but it is stable in presence of excess of NH, in the cold. At 60° it is completely split up into CO, and NH, (Naumann, A. 160, 1; B. 18, 1157; Horstmann, A. 187, 48; Erckmann, B. 18, 1154). In a sealed tubo at 140° it forms urea.—NaA'xaq: formed by adding NaOEt to an alcoholic solution of the ammonium salt; prisms.-KA': deliquescent. —CaA'₂ aq: ppd. by adding lime and alcohol to a solution of NII₄A' at 0°; crystalline powder, sol. water, the solution quickly deposits CaCO₂. When strongly heated it leaves calcium cyanamide. -- SrA'₂. -- BaA'₂. -- Chloride OC(NII.)Cl. [c. 50°]. (62°). Pre-

pared by passing a stream of dry COCl, into NH Cl heated to about 400°. Long broad needles. Strong odour. On keeping it slowly changes into evamelide with evolution of HCl. By water it is decomposed into NH,Cl and CO2. On vaporisation it probably dissociates into cyanic acid and HCl, which again recombine on cooling. By CaO it is convorted into oyanic acid. With aromatic hydrocarbons in presenct of Al₂Cl₆ i. gives amides of aromatic acids (Gattermann a. Schmidt, B. 20, 858).

Carbamic ethers. Srethancs.

Preparation .- 1. From ehloro-formic ethers and NH₃-2. From eyanic acid and alcohols.-3. From cyanogen chloride and alcohols.-4. By heating alcohols with urea nitrate.

Properties .- Solid substances, sl. sol. water, v. sol. alcohol and ether; may be distilled.

Reactions .- 1. Heating with NII, gives ures. 2. P2O5 gives cyanates. - 3. Alcoholie KOII acts upon carbamic ethers of the fatty series according to the equation: NiI, $OO_2C_1H_{2n_1} + KOH = Kh.CO + C_6I_{2n_1}OIH + H_2O$ (Arth, Bl. [27 45, 702; A.Ch. [6] 8, 428). Bornyl and menthyl carbamates act similarly.

Methul ether MeA'. [529]. 217 at 110°; S. (alcohol) 73 at 15° (Echevarria,

A. 79, 110).

Ethyl ethere Eth. Urethane. Mol. w. 89. [c. 50°]. (c. 182°). Formed by the above methods (Dumas, A. Ch. [2] 54, 233; A. 10, metnoas (Dumas, A. Ch. [2] 54, 238; A. 10, 281; Liebig a. Wöhler A. 54, 370; 58, 260; Wurtz, A. 79, 280; C. R. 22, 503; Bunte, Z. [2] 6, 96; A. 151, 181). Also from carbonic cther and NH₃ (Cahours, C. R. 21, 629; A. 56, 200).

Reactions .- Alcoholic potash, at the ordinary temperature, gives large crystals of potassium cyanate. In this case NH, CO, K is not formed as an intermediate product. A solution of urethane in absolute ether, treated with K or Na givos the derivatives KNH.CO₂Et, and NaNH.CO. Et. Of these, the Na derivative is sufficiently stable for analysis. It is v. sol. absoluto alcohol, insol. absoluto ether. With alcoholic potash containing water K.O. is obtained. The body HgN.CO.OEt is obtained by mixing alcoholio solutions of urethane, HgCl2, and KHO (Mulder, R. T. C. B, 170).

Acetyl derivative NHAc.CO.Et.

(Conrad a. Salomon, J. pr. [2] 10, 23). Chloro ethyl ether H.N.CO.C.H.Cl. [76°]. From NH, and the chloro-ethyl ether of chloroformic acid. Prisms. V. vol. water, aleohol and ether (Nemirowsky, J. pr. [2] 31, 174).
n-Propyl ether PrA'. [53°]. (195°) (Cahours, J. 1873, 748; Roemer, B. 6, 1102).

Long prisms. Isobutul ether C,H,A'. [55°]. (207°) (Mylius, B. 5, 973; Humann, A. Ch. [3] 44,

840; A. 95, 372).

Isoamyl ether C, H, A'. [60°]. (Medlock, A. 71, 106; Wurtz, J. Ph. [3] 20, 22). Noedles.

Octylether C,H,A'. [55°]. (135°) at 25 mm.; (231°) at 760 mm. On distillation it is partially converted into cyanuric acid (Arth, O. R. 102, 977).

Bornyl carbamate v. p. 523.

Menthylcarbamate v. Menthon.

CARBAMIDE v. UREA.

CARBAMIDO- v. URAMIDO-.

CARBAMINES. Carbylamines. Iso-nitiles. Compounds of the formula R.N:C.

Formation.-1. By distilling primary monamines with ohloroform and alcoholio potasli: $RNH_2 + CHCl_3 + 3KOH = 3KCl + RNC + 3H_2O$ (Hofmann, A. 144, 114; 116, 107). - 2. By treating an alkyl iodido (I mol.) with silver cyanide (2 mols.) a double salt RNCAgCy is formed; on distilling this compound with conc. aqueous KCy there is formed KCyAgCy and the carb-amine passes over (Gathier, A. 146, 119; 149, 29, 155; 151, 239). HgCy₂ and ZnCy₂ may also be used in preparing carbanines (Calmels, BL, [2] 48, 82).-3. In small quantity in preparing nitriles by distilling potassium alkyl sulphates with potassium cyanide.-4. By distilling the compounds of thio-carbimides with tri-ethylphosphine (Hofmann, B. 3, 766; Z. 7, 29).

Properties .- Volatile stinking poisonous oils. Reactions.—1. Alkalis have no action.—2. Mineral acids instantly convert them into alkylamines and formic acid: RNC+2H_O= RNH2+H.CO.H. Water at 180° acts similarly. 8. Dry HCl forms a compound, quickly decomposed by water as in 2 .- 4. Organic acids form alkyl-formamides .- 5. EtI forms a compound (difference from nitriles).-6. HgO oxidises them to alkyl cyanates R.N.CO, alkyl-formamides being also formed (Gautier, A. 149, 811)

CARBAMINE-CYAMIDE or CARBAMINE-CYANAMIDE so called is described as Amoor discrementations. Its derivatives are described as ETHYL-CARBIMIDO-UREA, CARBIMIDO-ETHYL-UREA,

CARBANIL v. PHENYL OYANATE. OARBANIL10 ACID v. PHENYL-CARBAMIO

CARBANILIDE v. 8-DI-PHENYL-UREA.

CARBANILIDO- p. PHENYL-URAMIDOv. PHENNA.

ClaHaN i.ec. C.H. ND.

C.H. ND. 5-86

V.D. 5-86 CARBAZOLE [236°]. Mol. w. 167. (oalo. 585). S. (alcohol .92 at 14°; 8.88 at 78°.

S. (toluene) .55 at 16.5; 5.46 at 100° (Bechi, B. 12, 1978).

Occurrence. - Among the products of the distillation of coal tax; hence it occurs in orude anthracene (Graebe a. Glasor, B. 5, 12, 376; A. 163, 513; 167, 125; 174, 180; 202, 21; Zeidler, A. 191, 297).

Formation. - 1. By passing vapour of aniline or diphonylamine through a red-hot tube .-2. From imido-di-phenyl snlphide (thio-diphonylamine) by boiling $HN < \stackrel{\circ}{C_0} \stackrel{\circ}{I_1} > S$ with freshly reduced copper for 2 or 3 hours; the yield is about 60 p.c. (Goske, B. 20, 233).

Properties.—White laminæ or tables. Easily

sublimes. A solution in conc. H2SO4 is turned greon by HNO3. May be distilled over red-hot tine-dust without change. Although an imide, it forms a compound with picrio acid and its acetyl derivative is obtained with difficulty.

Reactions. — I. It is not affected by cono. HClAq or alcoholio KOH even at 300°. Cold conc. II, SO, dissolves it without ohange, but at 100° a di-sulphonic acid results. - 2. HNO. forms nitro- compounds .- 3. Sodium-amalgam does not reduce it in alcoholio solution, but HI and P at 210° reduce it to carbazoline C12H11N .--4. By exhaustive chlorination with SbCl, it yields por - chloro - diphenyl or por - chlorobenzene according to circumstances as yet undetermined (Merz a. Weith, B. 16, 2875).— 5. By heating with oxalic acid the compound $C_{27}H_{23}N_6O$ or $HO.C(C_0H_2 < N_{C_0H_2})_3$ is obtained. It forms minute C_7 ystals which very readily yield bluo solutions (carbazolo blue) on or Mation (Suida, B. 12, 1403; Bamberger a. Müller, B. 20, 1905).

Potassium derivative C₁₂H₈NK. From carbazole and KOH at 230°. Piorio aoid compound C_{1.}H₈N C₂H₈(NO₂)₂OH. [182°]. From carbazole (1 pt.) and pierio acid (1 pts.) in tolueuc. Red prisms; v. sl. sol. cold bonzeno or alcohol. Decomposed by a large quantity of alcohol, by water, and by alkalia.

Nitrosamine C.H.N.NO. [82°]. Nitrons acid in an atsoliolic solution of oarbazole forms mono- and di-nitro-carbazole. If carbazole (3 g.) be mixed with acetic acid (60 g. of S.G. 1.04) and other (60 g.) be poured in, on adding KNO, the nitrosamine is dissolved in the ether as fast as it is formed, and crystallisos out on evaporation. Long flat golden neodies. Solublo in ether, CS₂, chloroform, glacial acetic acid and benzene. It is decomposed if heated with alcohol mixed with an acid, carbazole being regenerated. Alcoholic KOII turns it blood-red. Reducing \ gents regenerate carbazole. Cono. II,SO, gives a dark-green colour (Zeidler, A. 191, 305).

Acetyl derivative CuH, NAc. (above 360°). From carbacole and Ao O at 250°. Slender needlos (from water); v. sl. sol. water, v. e. sol. alochol. Erdmann's solution does not turn it gresn. Its pioric acid compound

is orange.

References. - BROMO-, CHLORO-, NITRO-, METHYL-, and ETHYL-, CARBAZOLE

CARBAZORE TETRAHYDRIDE C12H13N. [120°]. (c. 328°). Formed, together with hydrogen, by heating oarbazoline hydrochloride at 800°. Crystallises from alcohol. V. e. sol. alcohol, insol. water. Does not combine with acids. Reduced by HI and P to carbazoline. The piorio acid compound C₁₂H₁₃NC₂H₂(NO₂),OH forms brown lamins.

CARBAZOLE v-CARBOXYLIC ACID C₁₃H_eNO₂ i.e. C₁₂H_eN.CO₂H. $[272^{\circ}].$ potassium carbazole and CO, at 270°. Micaceous soales or flattoned prisms with faint blue fluor-

escence: insol. water, sl. sol. cold alcohol (Ciamioian c. Silbor, G. 12, 272).

CARBAZOLINE C₁₂H₁₅N. Carbazole hexahydride. [99°]. (297° i. V.). V.D. 6·13 [5·99 calc.). Formod by heating carbazole (3 pts.) with (12 pts. of) HIAq (127°) and amorphous P (1 pt.) at 220°. White needlos (from alcohol); may be sublimed; volatile with steam; v. e. sol. alcohol and ether, v. sl. sol. water. H1 and 1' at 339° reduce it to diphenyl decahydride C121I20. Does not combine with pieric acid. Salts.—B'HCl: v. e. sol. water.—B'11Br:

tables.—B'HI.

Acetyl derivative C12H14NAo. From carbazoline and Ac.O at 110°. Needles

(from alcohol).

· CARBIDES. Compounds of carbon with one other more positive etement. A carbide of iron Fe₂C probably exists in cold rolled steel; other carbidos of this metal are described, but their existence is doubtful. Silver is said to form three carbides, Ag,C, Ag,C, and AgC. Nickel takes up a small quantity of carbon when strongly heated with chareer but no definite compound has yet been prepared. We have very little definite information regarding this class of compounds (v. Iridium, Thon, Nickel, Palladiem, Platinum, and Silven). M. M. P. M. CARBIMIDE v. CYANIC ACID.

CARBIMIDO-ALLYL-THIO-UREA C,H,N,S i.e. SC<\br/>NII >C:NH or C_111,N11.CS.N:C:NH. Allyl-thio-carbamine-cyamide. Formed, as tho crystalline sodium salt, by mixing allyl-thiocarbimide and sodium cyanamide. Decomposed by acids into its constituents (Wunderlich, B.

19, 448). CARBIMID - AMIDO - BENZOYC ACID is

Guanido-di-bonzoic acid v. p. 157. CARBIMIDAMIDO-BENZOYL v. OXY-QUIN-

AZOLINE and p. 155. CARBIMIDO-CYANAMIDE v. AMILO-DICYANIO

AOID, p. 163.

CARBIMIDO-ETHYL-THIO-UREA C.11, NoS i.e. SC<NH or EtHN.CS.N:C:NH. Ethyl-thio-carbamine-cyamide. Formed, as the crystalline sodium salt, by mixing ethyl-thio-19, 448)

CARBIMIDO-ETHYL-UREA C.H,N,O i.e. OC NEt C:NH or EtHN.CO.N:C:NH. Ethylcarbamine-cyamide. Formed, as the orystalline

stituents by acids. Forms a green crystallitie copper compound (Wunderlich, B. 19, 448).
CARBIMIDO-METHYL-THIO-URKA

C₂H₃N₂S i.e. SC<NM >C:NH or

MeHN.CS.N:C:N11. Milhyl-thio-carbamine-cy amide. Formed, as the crystalline sodium salt, by pixing methyl-thiocarbinide and sodium cyanamide. Decomposed by acids into its constituonts (Wunderlich, B. 19, 448).

CARBIMIDO-PHENYL-THIO-UREA

C_sH₇N₃S i.e. SC<NII or

PhNH.CS.N.T.NH. Phenyl-thio-carbaminecyamide. Formed, as the crystalline sodium salt, by mixit phenyl-thiocarbimide and sodium cyanamide. Decomposed by heids into its constituents (Wunderlich, B. 19, 448)

DI-CARBIN-TETRA-CARBOXÝLIC ACID v.

ETUYLENE TETRA-CARBOXYLIC ACID.

CARBINOL. A name given by Kolbe to methyl alcohol, but used only in describing alcohols derived therefrom by displacement of hydrogen of its methyl by one or more alkyls. Cf. Alconols.

CARBINYL. The corresponding term for the alcohol radicles of the alkyl-carbinols; thus, Me₃C may be called tri-methyl-carbinyl.

CARBO-ACETO-ACETIC ETHER v. p. 20. CARBO-ALLYL-PHENYL-AMIDE v. Phenyl-

ALLYL-CYANAMIDE CARBO-DI-BUTYL-DI-PHENYL-IMIDE v.

DIBUTYL-DI-PHENYL-CYANAMIDE.

CARBO-ISO-BUTYRALDINE C,II, N2S2 i.e. (NH₂)CS.S.N(C₄H₈). [91°]. From iso-butyralde-hyde, CS₂, and aqueous NH₃. Prisms, insol. water, sol. alcohol (Pfeiffer, B. 5, 701).

CARBO-CAPRO-LACTONIC ACID v. Lactone

OXY-PROPYL-SUCCINIC ACID.

Di-carbo-capro-lactonio acid v. Lactone of

OXY-PENTA OR TO CURROXYLIC ACID. CARBOJINCHOMERONIC ACID is PYRIDINE

TRI-CAUBOAYIAC ACED. . CARBO-GLUCONIC ACID C.H. O. amorphous acid whose NII, salt is obtained by treat a glucose or cane-sugar with aqueous HCN (Schützenberger, Bl. [2] 36, 144).

CARBO-DI-GLYCOLLIC ETHER v. GLYCOLLIO

ACID. CARBOHOMOPYRROLIC ACID v. METHYL-

PYRROL-CARBOXYLIC M.i.D.

CARBOHYDRATES. A term applied to compour ds which may be represented by the formula $C_X(H_*C)_Y$ where x is 5, 6, or 12, and y is 5, 6 or 11, and to compounds derived from several such molecules by abstraction of water. They are non-volatile solids, and the non-saccharine members of the grou; may be converted by boiling dilute acids kato a sugar, usually glucose (dextrose). Tray contain hydroxyl. On oxidation they frequently give rise to oxalio, racemio, saccharic, and mucic acids. Most of them are orytainine and sodium cyanamide. Decomposed, optically active. Celluluse is insoluble in water; by soids into its constituents (Wunderlich, B. the gums dissolve, or at least swolf up, in water, but are ppd. by alcohol. Sugars are soluble in water, are not reppd. by alcohol, and have a sweet taste. Iodine turns starch blue, and affects ocllulose in the same way after it has been treated with a delaydrating agent. The sodium salt, by mixing ethyl cyanate and carbohydrates vary also in their behaviour to-sodium cyanamide, Decomposed into its con-wards alkaline coppor solutions, and as regards carbohydrates vary also in their behaviour to-

the articles Arabio acid, Cellulose, Dextrin, STAROH, SUCAR, ETC.

CARBOLIC ACID v. PHENOL.

CARBO-MESYL v. METHYL-OXINDOLE CARBO-DI-NAPHTHYL-IMIDE v. DI-NAPH-

THYL-CYANAMIDE.

CARBON GROUP OF ELEMENTS .- Carbon and Silicon. Of these elements, carbon occurs in the free state in the forms of diamond, graphite, and amorphous carbon; silicon is not known at such in naturo, but combined with oxygen it is one of the most widely-distributed elements. Diamond was regarded by Newton as a combustible body because of its great refractive power: in 1694 the Florentine academicions succeeded in burning small pieces of diamond; and in the early years of this century Davy proved it to be pure carbon. In early times graphite was thought to be very similar to lead; hence the

fermentation by yeast. They are described in name plumbago; for a time it was confused with molybdenum-glance, but in 1799 Scheele proved it to be closely related to coal in its composition. It is only in somewhat recent times that approximately pure graphite has been obtained. Char-coal is the commonest form of impure amorphous carbon; this modification of carbon can be obtained approximately pure only with considerable difficulty.

After the earthst had been proved to be metallic oxides in 1807, it was generally supposed that the common earth-like body silica would also be found to contain oxygen and a In 1823 Berzelius decomposed silica metal. and obtained the non-metal silicon, in the form of a brown amorphous powder. Argeod many years later Deville prepared crystallised silicon in two forms, one more or less resembling diamond. and the other, graphite. The leading properties of the two elements are as follows:-

	CARBON.	Silicon.	
Atomic weights	11.97	28	

Many compounds of each element have been gasified. Molecular weights unknown: (?) probably greater than C2 and Si2.

Melting-points Specific gravities (approximate)

Specific heats

Does not melt at any temperature hitherto attained. Diamond 3.5; graphite 2.25; amorphous 1.5-1.9.

0.46 (at about 1000°).

1,100°-1,300° (uncortain).

Graphitoidal 2.2-2.5 (doubtful).

0.203 (at about 250°)

The specific heat of either element increases rapidly as temporature increases from -50° : the rate of this increase is showever, very small after about 500° for earbon and about 150° for silicon. The specific heats of diamond and graphito vary considerably at temperatures below about 600°, but from this point upwards the values are practically identical.

Atomic weight Spec. grav. (approximate)

11.2

· Heats of formation of various compounds (Thomsen, Berthelot, &c.). (Generally from amorphous Carbon or Silicon.)

[M,H']	21,750	24,800
[M,Cl]	21,00	157,600 (product liquid)
$[M,O^2]$	96,960	219,200
[M,S ²]	-26,000	40,000
Change of amor-		
phous M to crys-	3,000	8,100
tolling M		1

Heats of neutralisation of aqueous solutions of CO2 and SiO2 (Thomsen).

n [CO ² Aq, nN ₁ OHAq] 1 11,000 diff. = 9,150 2 20,150	;	n in Line 2	[SiO ² Aq, n NaOHAq] 3,240 4,315 4,730 5,230 5 410
•		1.4	5,410

Silicic acid shows no constant neutralisation point. The quantity of heat produced is a hyperbolic function of the quantity of soda added, and approaches a probable maximum of 6,300 gram-units for one formula-weight of SiO2 (v. SILICATES).

Physical properties

Diamond: hardest known substance; crystallises in regular forms, octahedral predominating; bad condnctor of electricity; refractive index large ($\mu_D = 2.430$); lustre very marked; usually colonrless and transparent, but sometimes green, brown, or yellow,

Adamantine: very hard, scratches glass, darkiron-greycolour, reddish by reflected light; crystallises in forms derived from a rhombio octahedron.

Graphitoidal: softer than adamantine but scratches glass; may be pulverised; metal-like lustre, leadenTable -cont.

CARBON.

SILICON.

Graphite: cryetallises in hoxagonal forms; good conductor of electricity; tough and difficult to preverise; grey, metal-like appearance

Amorphous: black powder; very porous, absorbs large quantities of gases and of many colouring matters from solutions.

Occurrence and preparation

The three forme occur in nature, but neither graphite nor amorphous pure; constituent element of all animal and vegetable matter; carbonates very widely distributed; graphite prepared by dissolving amorphous in molten iron, or by decomposing by heat the CN compounds in the mother liquor of eoda manufacture, &c.: approximately pure amorphous, prepared by washing eugar-charcoalin acid, alkali, and water, and strongly heating in chlorine; or by decomposing CO₂ by Na, &c.

Ohemical properties.

Allotropy marked. Diamond heated by powerful battery in absence of oxygen gets grey-black and cokoliko bnt does not volatilise; heated in air combustion begins at 950'-1000°. Graphite not affected at any te aperaturo in absence of oxygen; oxidised by repeated treatment with KClO3 and IINO3 to graphitic acid C₁₁H₄O₅ (or O₆), a yellow sold, sl. sol. water, acting towards alkaline basee like a feeble acid. Amorphous burns easily in air; combines with H at a very high temperature to for a C2H2; also combines directly with S to form CS, with O to form CO and CO2, and under special conditions with N to form C2N2; compounds with halogens formed indirectly; combines directly with Ir, Fe, Ni, Pd, Pt, Ag, and perhaps some other metals. Carbon a negative olement; does not form salts by replacing II of acids; CO2 an anhydride; an aqueous solution of CO, probably contains the dibasic acid H.CO, salts of this acid well marked; H2CS, prepared. Atom of C is tetravalent; Catoms tend to combine with each other; vast number of compounds produced by addition of other atoms to groups of C atoms.

grey colour; gryetalliese in leaflets composed of betahedra; good conductor of electricity. Amorphous: brown powder; heated

out of coutact with air to high emperature it contracts and becomes erystalline; bad conductor of electricity; dissolves in molton Al or Zn and crystallizes out on cooling.

Very widely distributed as silicates of the Market Alectric amorphous

Very widely distributed as silicates of Ca, Mg, Fe, Al, & a.; amorphous obtained by action of K on hot SiCl., SiF, or K, SiFe; graphitoidal cobtained by melting Al with K,SiO, and cryolite, or by decomposing SiCl, at a high temporature by Na; adamantine obtained by melting Zn with K,SiFe and Na.

Allotropy marked. Amorphous Si burns easily in air to SiO,; graphitoidal does not oxidise when heated; adamantine not even at a white h at in oxygen; adamantine Si oxidised at red heat in CO, (giving CO + SiO2), also by strongly heating with K.CO, or Na.CO, (giving CO+SiO,+C), but not changed by molten KHSO, or by heating with KNO, if temp, at which that ealt decomposes is not reached. Amorphous Si soluble in HFAq giving ILSiF, Aq and hydrogen, also in strong hot potash loy giving K.SiO, Aq and hydrogen; adamantino Si insoluble in IIFAq and hot alkali solutions. Si does not directly combine with II, SiH, produced by action of HClAq on compound of Si and Mg; combinee with S at high temperature to form SiS,; with O to form SiO2; with Cl, Br, or I, to form SiCl., SiBr., or Sil,; and with N at white heat to form Si₂N₃; combines directly with Al, Cu, Fe. Mg. Mn, Ni, Pt, and perhaps some other metals. SiO, an anhydride; probable existence of deveral silicic acids; SiO(OH), probably present in solution obtained by neutralising K.SiO.Aq by HClAq and dialysing; this colution very readily gelatinises. Atom of Si is tetravalent, and, to some extent at least, atoms of Si tend to combino together and form groups which combine with other atome, forming molecules eimilar to thoes of the organic compounds.

General formula and characters of compounds. MO, MO, MS, (SiO unknown, ?CS and C,S,); MO,H, (neither known except (?) in aqueous solution, v. Carbon and Silicon), CS,H,; aqueous sointond, v. Carbon and Silicon, Os_1A_2 ; MH_4 ; C_1H_{2n-1} ; C_2H_{2n-1} , C_2H_{2n-2} , &co., &co., and a vast Lumber of derivatives; MX, $(X=Cl, Br, I, or in case of Si also = F), <math>M_1X_6(X=Cl, Br, or I where M=Si, X=Ul or Br wheres <math>M=Cl$, C_2Cl , C_2Br , &c.; SiF_1H_2 ; CH_2Cl , CH_2Cl , CHCl, SiHCl, $SiHI_3$, &c.; C_2N_4 , CN_4 , CN_4 , CN_4 , CN_5 and salts, C,N,NOFeH, and salts, &o., &o.; Si,N, &o. The compounds of C and Si exhibit oonsiderable differences in their properties; CO and CO. are gases, SiO, is a very fixed solid; CS. is a liquid, SiS, a solid; CCl, is not acted on by water, SiCl, is at once decomposed into SiO, and HCl; Si readily forms a fluoride and also a donble fluoride with hydrogen, no coffesponding compounds of C are known; CH, is a stable gas, SiH, is oxidised by mero contact with air and is easily decomposed by heat (at 400); Si (amorphous) dissolves in potash evolving hydrogen and forming a silicate, carbon is unacted on by alkalis. Both elements form many compounds with H and O (alcohols, others, acids &o.), the composition of which is similar, in some cases the properties of the Si compounds closely resemble those of C, e.g. C(C,H,),H and Si(C2H3)3H, C(C2H3)3OH and Si(C2H)3OH; but in other cases the properties of the two classes of compounds differ much, e.g. CII, CQ.H and C.H. CO.H are liquids soluble in water, but CH, SiO II and Cail SiO II are amorphous solids insoluble in water. Many silicates and carbonates are isomorphous. cates, except those of the alkali metals, are insoluble in water, and most of them are with difficulty decomposed by acids; the normal carbonates of the alkali metals are soluble in water, other normal carbonates are insoluble; aqueous solutions of heid carbonates are generally easily decomposed by heat yielding either normal or basic carbonates; the normal carbonates of the alkali metals are not droomposed by heat alone, other normal carbonates are decomposed into metallic oxide and CO.,

Group IV. of the elements, as the elements are classified by the application of the periodic law, contains the following:—

The metals titanium, zircouium, and germanium show considerable analogies with tin; cerium and thorium are usually classed together among the rarer earth-metals, and lead is generally considered apart from other metals: nevertheless, there are gwell-marked analogies between all the elements which comprise Group IV. of the periodic system. Titanium is an amorphous body closely resembling amorphous silicon; it forms the compounds TiF., TiCl., TiBr., TiI., TiCl., TiO., (probably TiO(dH) and Ti(OH).), Ti.O., Ti.N., &c.; titanates are known [M.180.], many of them isomorphous with sili-

C or Si, it forms a sulphate Ti(SO4), and other salts wherein the hydrogen of acids is replaced by titanium. Zircchium again is more decidedly metallic than titenium; it forms a series of well-marked salts Zr(SO)2, Zr4NO3, &o., &o. On the other hand Zr rosembles C and Si in that it has been obtained both as an amorphous powder, and also in crystals which resemble Si in their behaviour (owards acids; zirconates (M.ZrO.) are also known. Germanium forms oxides, chlorides, and sulphides, &c. (GeX and GeX₂, $X = O = S = Cl_2$), resembling those of Sn; it is, however, more markedly non-metallic in its chemical functions than Sn; physically Ge is decidedly metallic. Cerium forms two oxides Ce.O, and CeO.; the former dissolves in acids forming a series of ealts of which Ce. 3SO. is a type; CeO, is a peroxide, it dissolves in HCl with evolution of Cl and formation of CeCl, but a sulphate Ce(SO₄)₂ is known corresponding to the sulphates of Ti and Zr. Ce also forms a fluoride CcF, a double fluoride 3KF.2CeF, and a chloride Ce, Cl,. Thorium again approaches more closely than cerium to Zr and Ti; it is a dark-coloured amorphous powdor resembling Si, but more soluble in acids than Si, Zr, or Ti; it forms the compounds Thel, ThF, K2ThF6 ThO, &c.; the sulphate is Th(SO,), and other analogous salts are known. Tin forms the two oxides and ohlorides SnO and SnO2, SnCl2 and SnCl,; the hydrates of SnO, are feebloacids, producing stannates (M.SnO3) and metastannates (M.Sn.O.1), both of which are easily de-composed by diluto acids or by heat. Both the stannous salts e.g. SnSO, and the stannic salts e.g. Sn(SO_d), are well-marked compounds. Lead is decidedly metallic in its character; it forms four oxides PbO, Pb3O4, Pb2O3, and PbO2; the last is a peroxide, and it may also be regarded as an anhydride inasmuch as plumbates (M.PbOa) wist, but these salts are unstable and easily decomposed. The best-marked salts of lead are derived from the oxide PbO, e.g. PbCl, Pb2NO3, PbSO4, &c.; PbCl, has not been obtained pure, but this series of salts is represented by the tetramethide Pb(CH3)4 which is stable as a gas. The atoms of all the elements of Group IV., so far as evidence has been obtained, are tetravalent. Looking at the properties of these elements as a whole, it may be said that carbon is to a considerable extent set apart from the others, but that it is more closely allied to silicon than to any other member of the group; that titanium and ziroonium are closely related; and that tin and lead, while showing distinct analogies with the rest of the group, are yet each characterised by properties which mark them off from the other elements. Not much can yet be said regarding cerium and thorium; they require further study.

For more details and descriptions of the various elements see the articles on these elements; also v. Titarium oroup of elements; also v. Carronares, Nitraates, Sculphates, &c. In some of their physical properties carbon and silicon, especially the latter, resemble boron, but boron must be classed with those elements the atoms of which are trivalent: v. Boron.

M. M. P. M.

(M.T.Co.), many of them isomorphous with silisates and carbonates. Ti is more metallic than element has not been gasified. S.G. diamond

38° 8.514 (Schrötter, Sitz. W. 63, (2nd pt.) 462), 2 3-518 (Banmhauer, Ar. N. 8, 1). graphite 2 11 to 2 26 (Kenngott, Brodie, Mêne; Sitz. W. 13, 469; A. 114, 7 P. C. R. 64, 104). S.G. amorphons charcoal 140 to 1.7 (v. Vic-5.07. almorphis of other of 1.74 (b. Violette, A. Oh. [3] 93, 291). S.C. hard gas-coke 2.356 (Marohand a. Meyer). S.H. about 5 at 1000° (v. infra). C.E. (diamond, linear at 40°) 00000118; (diamond, cub. at 49°) 00000354; (graphito, linear at 40°) 00000786; (Fizeau, C. R. 62, 1133; 68, 1125). $\mu_{\rm h} = 2.16$, $\mu_{\rm h} = 2.479$, for diamond (Schrauf, P. 112, 588). E.C. graphito, 138; (Hg at 0°) 15 (review linear) for different control of the control of th phito, 082 (Hg at 0°=1) [varies much for different specimens] (Mnraoka, W. 13, 307). E.C. hard gas-coke, 01 (Hg at 0°=1) (Mnraoka, Lc.). Crystallino Jam; diamond, regular octahedra and forces derived therefrom; graphite, hexago-nal forms chiefly rhombohedral (Kenngott, Sitz. W. 13, 469); Nordenskiold (P. 96, 100) observed monoclinio crystals in graphite from Finland. **H.C.** $[C,O^{2}] = 96,960$ for amorphous C (Th. 1, 411); 93,350 for diamond, and 93,560 for graphite (Favre a. Silbermann, A. Ch. [3] 33, 411). Emission-spectrum observed by passing sparks through pure CO or CO, is characterised by a double line 6583 and 6577.5, three sharp lines **5150·5, 5144·2, 5133, and a** band **1**266 (Ångström a. Thalen, Nov. Act. Ups. 9 [1875]). Besides these, and many other less marked, lines, Liveing a. Dewar describe the arc-spectrum as showing the following marked lines, 3919-3. 2837-2, 2836-3, 2511-9, 2509-9, 2296-5 (Pr. 30, 152, 494; 33, 403; 34, 123, 418). A very different spectrum—the band-spectrum is observed at the base of a candle or gas flame, also in oyanogon burnt in O, or by passing sparks through CN, CO at increased pressure, CS., &c.; the most characteristic band are 5633, 5164, and 4736. There has been much discussion as and 4730. There has been much ensusation as to whether this spectrum is that of C or of a hydrendon (v. B. A. 1880, 254). Three allotropic forms of carbon are known; diamond, graphite, and amorphous carbon.

The diamond was regarded by Newton as a

combistible body because of its high refractive power; in 1691 diamond was burnt by the Florentine Academicians; Lavoisier found that CO2 is produced when diamond is burnt, and Davy showed that diamond is pure carbon. Lavoisier, about 1780, recognised that carbonie acid (thon called fixed air) was a compound of O and the element which is the essectial element of coal; to this element he gave the name car-Graphite was long considered to be a kind of lead; Scheele, in 1799, showed it to be closely related to coal; he regarded it as a compound of iron and carbon, but Kastner proved that the iron found in graphite was only an impurity, and that pure graphite is a form of

carbon.

Occurrence.-Carbon occues as diamond and graphite, the former is pure, the latter sometimes approximately pure, carbon; many compounds of Coccur in nature; the chief are CO2 in the air and all waters, mineral carbonates e.g. of Ca and Mg, and compounds with H, O, N, and sometimes P and S, in all animal and vogetable organisms. Diamonds are found in India, Borneo, Brazil, the Cape, &c.; graphitc, in Cumberland, California, Siberia, &c. Berthelot

(C. R. 73, 494) found graphite in a meteorite which fell near Melbourne (Australia); and Fletcher found a cubic form of graphite in a meteorite from Western Anstralia (Mineralog. Mag., Jan. 1887). Graph te is found both amorphous and foliated. Coal, anthraoite, poat, &c., contain from 50 to 95 p.c. of carbon.

Formation.—Many attempts have been made to forif diamond; none has been certainly successful (v. Liebig, Agriculturchemic [1840] 285; Wilson, J. 1850. 697; Favre, J. 1856. 828 [from CCl.]; Despretz, C. R. 37, 369 [elootrio current for a month from Pt to C pole]; Siminler, P. 105, 466 [crystalli ation from liquid CO.]; Lionnet, C. R. 69, 213 [from CS.]; Chancourtois, C. C. 1866. 1037 [oxidation of hydrocarbon]; Rossi, C. R. 63, 408; Hannay, Pr. 30, 188 a. 450 [action of Mg, and Li, on gascous hydrocarbons mixed with N-containing compounds at very high temperatures and pressures]; Marsden, Pr. E. 11, 20 [by dissolving amorphous C in molten Ag]). Graphite is formed: -1. By heating charcoal with molten iron, and dissolving out the Fe by HCl and HNO, Aq. -2. By the slow decomposition of HCNAq, and boiling the product with HNO, Aq (Wagner, J. C. T. 1869. 230). -- 3. By evaporating the mother liquors obtained in making soda; these contain CN compounds which are decomposed at a certain concentration of the liquid with formation of NII, and graphite (Pauli, D. P. J. 161, 129; Schaffner, W. J. 1869, 250).-4. By leading CO over Fe.O. at 800?—400° (Grüner, C. R. 73, 28; Stingl, B. 6, 392). Amorphous C is also formed (Berthelet, C. R. 73, 494).—5. By the decomposition of CS, at high temperatures. 6. By leading CCl, over molten pig-iron (Deville, A. Ch. [3] 49, 72). Amorphous carbon is formed in many ways:—1. By heating wood, coal, or almost any animal or vegetable matter, out of contact with air, to a high temperature. -2. By the incomplete combustion of wax, tallow, oil, or other combustible compounds of C and H.-3. By decomposing, at a very high temperature and out of contact with air, the gaseous C compounds obtained in the production of gas from eoal: the earbon thus obtained is very hard (v. Proper-

Preparation. Pure graphite is obtained by Intimately mixing 14 parts of finely powdered foliated graphite with 1 part KClO, and 2 parts conc. H.SO, heating on the water-bath so long as Ci comes off, washing repeatedly with hot water drying and heating to remove H₂SO₂: if the graphite contains silica it is treated with Naf and H.SO, besides treatment with KClO, and H.SO, (Brodio, T. 1860.1; v. also Winckler, J. pr. 98, 243; Stin. 1, B. 6, 391).

Amorphous carbon is prepared approximately pure by strongly heating cane sugar in a closed I't crucible, boiling the charcoal thus produced with (1) conc. HClAq, (2) KOHAq, (3) water, drying, heating to full redness in a stream of dry Cl and allowing to spol in the same; H is removed as HCl, O as CO, also traces of SiO₂, Fe₂O₃, &c. as SiCl₄, Fe₂Cl₅, Al₂Cd₅, &c. The soot from semi-burnt turpentine oil, after treat-ment with ether, and heating to a high temperature in a closed vessel, is approximately pure oarbon. It seems to be impossible to obtain finely divided amorphous O quite free from gases such as H, O, or Cl; even when purified as described it retains traces of Cl, this may be removed by strongly heating in connection with a Sprengel pump, but on exposure to the air considerable quantities of O, CO, &c. are quickly absorbed. The absorbed gases cannot bo removed by hoading at ordinary pressures; Erdmann a. Marchand (J. pr. 23, 169) found 2 p.o. H and 5 p.c. O in sugar-oharcon which had been heated nearly to whiteness for 3 hours. According to Porcher (C. N. 44, 203) amorphous C free from H, O, and N is obtained by passing CCl, vapour over hot pure Na in a hard glass tube, and then heating the C obtained to a little under the temperature at which burning begins. A very hard kind of amorphous carbon is formed by placing wood (box; ash, elder, lilac, or oak), or flax, hemp, cotton, paper, or silk, in a porcelain tube, driving out all air by CS, vapour and then gradually heating to redness for an hour (Sidot, C. R. 70, 605). The harder the wood and the higher the temperature to which it is heated, the harder and denser is the carbon produced. Various materials consisting mainly of carbon are prepared for industrial use; charcoal, by partially burning piles of wood covered with turf or earth, or by the dry distillation of wood; colic, by heating coal in iron retorts arranged so that the liquid and gaseous products may be separated from the residual carbonaceons matter; lamp black, by partially burning tallow, turpentine, &c., and condensing the soot on cold surfaces; animal char (which however contains only about 10 20 p.o. C) by heating bonng in closed vessels.

Properties .- Unchanged by action of acids;

has not been melted or vaporised.

Diamond is a colontless, transparent, very refractive and dispersive, crystalline, solid; some diamonds are coloured yellow, brown, blue, or black. Diamond is the hardest substance known, but rather brittle; very bad conductor of electricity and heat C.E. small, especially at low temperatures, at $-42^{\circ} = 6^{\circ}$ Unohanged by heating out of contact with air to 1300°-1100°; but placed between the carbon poles of a powerful battory it glows brilliantly, swells up, splits, and after cooling the surface resembles coke from bituminous coal (comp. Rose, P. 168, 497; v. Schrötter, Sitz. W. [2] 63, 462; Morres., C. R. 70, 990; Jacquelain, A. 61, 256; Gassiot, Ph. C. 1850, 893; Baumhauer, Ar. N. 8, 1). Unchanged when heated to whiteness in watervapour (Baunihauer, l.c.). Strongly heated in a stream of O, diamond is completely burnt to CO,; it may also be burnt by heating with molten KNO₃; or, very slowly, by powdering finely and heating with K.Cr.O., ILSO₄, and a little H.O (Rogers, J. pr. 50, 411).

Graphite occurs nativo both crystallino (foliated) and amorphous; it forms a grey, motal-like, hard, opaque, solid; fair conductor of electricity, especially after purification by KClO₂, &c. (n supra), fair conductor of heat; is not changed by heating out of contact vith air; burns in O to CO₂ at a high temperature, but more slowly than diamond; burnt to CO2, more oasily than diamond, by molten KNO, or by K₂Or₂O₂ and H₂SO₄Aq; also by heating with various metallic oxides. When graphite is heated with KClO, and HNO, Aq a compound of (l.c.); 201 R. (l.c.).

C. H. and O is formed, called by Brodie graphitie acid (probably C, H,O,); this body is not obtained from diamond or amorphous carbon (v.

Reactions, No. 9.

The graphits-like form of coke which is formed in the upper parts of the retorts in which ooal is hoated for gas-making, or is obtained by passing hydrocarbon vapours through red-hot porcelain or iron tubos, is an extremely hard, metal-like, lustrous, sonorous solid; S.G. (2.356) nearly same as that of graphite; it is a good conductor of electricity and a fair conductor of heat; burns with difficulty; it contains no H, and leaves only from '2 to '3 p.c. ash (Marchand a. Meyer).

Amorphous carbon (sugar-charcoal; lamp-black) is a denso, black, powler; it is extreinely slowly acted on by any reagonts, even energetic exidisers; non-conductor of electricity. The harder forms of amorphous carbon, obtained by calcining hard woods at high temperatures ont of contact with air, somowhat resemble graphite in appearance, they are more or loss lustrous, conduct electricity fairly woll, and burn slowly when heated in air or O. Ordinary amorphous C, or ordinary wood charcoal, absorbs large volumes of gases: Sanssure (G. A. 47, 113) gives the following volumes absorbed 47, 113) gives the following volumes absorbed by 1 vol. box-charceal at 12° and 724 mm.: NH₃ 90, IICl 85, SO₂ 65, H₂S 55, N₂O 40, CO₂ 35, CO 9·4, C₂H, 35, O 9·2, N 7·5, II 1·75. Hunter (P. M. [4] 29, 116; C. J. [2] 3, 285; 5, 160; 6, 186; 8, 73; 9, 76; 10, 649) gives these numbers for 1 volume coco-nut charcoal at 0° and 760 mm.: NH, 171-7, CN 107-5, NO 80-3, CH₂Cl 70-4, (CH₂) O 76-2, C₂H, 74-7, N₂O 70-5, PH, 69-1, CO, 67-7, CO 21-2, O 17-9. According to Angus Smith (Pr. 28, 322) absorption of gases by charcoal takes place in definite volumes; thus if the vol. of H absorbed under definite conditions is 1, the vol. of O=8, CO=6, $CO_2=2$ N=4 66. Chemical reaction some times occurs between gases absorbed by chareo.f.; thus, HCl is produced by leading H over charcoal which has absorbed Cl, and SO, Cl, by leading SO, over charcoal under the same conditions. The absorbed gases are removed in vacuo. Recently heated porons wood charcoal removes many colouring matters, c.g. indigo, from solutions; it also removes fusel oil from weak alcohol, alkaloids from aqueous solutions, man; metallic salts from solutions. &c.; in some cases chemical change is produced, e.g. CuSO, Aq and AgNO, Aq are reduced with pps. of Cu and Ag (Monde, J. pr. 67, 255; v. also Graham a. Hoimann, A. 83, 39; Graham, P. 19, 139; Weppen, A. 55, 241; 59, 351; Favro, A. Ch. [5] 1, 209; Guthe a. Harms, Ar. Ph. 69, 121; Stenhouse, A. 90, 186).

Specific heat of carbon.—The following numbers summarise the chief determinations exclusive of those of Weber: the temporature-

interval is about 35°-55°:

Diamond: 142 Bettendorff a. Wüllner (P. 133, 293); 147 Regnault (A. Ch. [3] 1, 202); 366 [20°-1,000°] Dewar (P. M. [4] 44, 461).

Gas carbon: 165 Kopp (A. 126, 362;

*Surpible 8, 1 a. 299); 186 B. a. W. (l.c.); 197 R. (l.c.); 32 [20°-1,000°] D. (l.c.). Graphite: 174 Kopp (l.c.); 188 B. a. W.

Wood charcoat: 24 R. (Lc.).
In 1874 Websr made essent determinations of the S.H. of the different forms of carbon at different temperatures; he used (1) diamond, (2) native graphite, (3) porous wood charcoal in a slender filament strongly heated in dry Cl and sealed at oncs in a glass tube. His chief results wers as follows (v. P.M. [4] 49, 161 a. 276):—

Diamoud.

+10° \$5° 250° 606° 985° Temp. -50° 8.H. 0635·1128 ·1765 ·3026 ·4408 ·4529 S.H.

Graphite.
Temp. -50° +10° 61° 201° 250° 641° 978° 8.H. • 1138 1604 199 2966 325 4454 467

Wood Charcoal.

Tomp. 0°-23° 0°-99° $0^{\circ} - 223^{\circ}$ ·2385 ·1935 ·1653

These numbers show that the S.H. increases as temp. increases, but that the rate of this inorease is much smaller at high than at low temperatures. From 600° onwards the S.H. of diamond is the same as that of graphite; as the values for wood charcoal are nearly the same as those for graphits for the same temperatureintervals, the conclusions may fairly be drawn that at temperatures above 600° the different forms of carbon have all the same S.H., and that at lower temperatures there are two values for the S.H., one belonging to graphite and amorphous C, the other to diamond.

Allotropy of carbon. Carbon exhibits allotropic changes in a marked way; diamond may be, superficially at any rate, changed to graphito; amorphous C may also be changed to graphito; each of the three varieties is charactorised by special properties. The S.G. of each is characteristic. The hears of combustion (v. supra) are different. The S.H.s are not the same; but Weber's results and to show that, as reserds S.H., there is but one form of C existing at emperatures above 600°. Amorphous C remained unchanged when subjected to a pressure of 6,000 7,000 atmos. (Spring, A. Ch. [5] 22, 170). The three forms are clearly distinguished, chemically, by their reactions with KClO, and HNO, (v. Reactions, No. 9).
Atomic weight.—Determined (1) by burn-

ing diamond in O and weighing the CO. produced (Dumas a. Stas, A. Ch. [3] 1, 5; Erdmann a. Marchand, J. pr. 23, 159; Roscoe, A. Ch. [5] 26, 136; Friedel, Bl. [2] 41, 100) ≥ (2) by heating silver acetate and weighing the Ag (Marignae, A. 59, 287); (3) by heating Ag salts (oxalate and acetate) and weighing the Ag and CO₂ formed (Maumené, A. Ch. [3] 18, 41). The mean of all the (closely agreeing) results is 11.97 (0 = 15.96).

Chemical properties .- The atom of C is tetravalent in gascous molecules (CII, CCl, CBr., &c.). The atomicity of the molecule of C is unknown, as the element has not been gasified; certain considerations, e.g. the increase in S.H. as temperature increases; and perhaps the character of the spectruma, seem to indicate that the molecule of C is probably composed of several atoms.

Carbon is distinctly a non-metallic element; it doss not replace the H of acids to form salts; it forms stable, but easily gasified, compounds flask surrounded by ice-cold water, and very

with the halogens; its oxides, and also the sulphids CS,, are distinctly acidic in their re-actions; it sxhibits allotropy in a most decided way; the spectrum of C is very complex; yet in some of the physical properties of graphite and dense amorphous carbon, this element approaches the metals (v. supra).) Carbon stands at the beginning of Group IV. in the periodic classification of the elements; the other members of this group, except Si, are more metallic than non-metallic; C shows closer relations to Si, the first odd-series member of the group, than to any other element in the group (v. Cannon or our or elements). Both elements are remarkable for the great number of compounds which they form with II, O, and N. Most of the elements of Group IV. except C, form characteristic compounds with F, or double confpounds with F and other elements.

Reactions.—1. Unchanged by action of acids.
2. Heat, in absence of air, produces no change (comp. Properties of Diamond). - 3. Whom strongly heated in excess of orygen, CO, is formed: the combination is much retarded if tho C and O are carefully dried (Baker, C. J. 47, 349) .- 4. Heated with sulphuric acid and potassium dichromate C is slowly burnt to CO2. 5. Oxidised to CO, by heating with molton nitrate or chlorate of potassium.—6. Reacts with sulphur vapour at high temperatures to form CS .. - 7. Combines with hydrogen to form C2H2; by passing electric sparks between C poles in atmosphero of H.-8. Combines indirectly with nivogen to form cyanogen .- 9. Graphite is oxidised by potassium entorate and nitric acid to graphitio acid (?Ch.H.O. or C.H.O.). Brodio (7. 1859, 249) heated an intimate mixture of 1 part purified and very finely divided graphite and 3 parts KClO3, with enough very cone. HNO3Aq to bring all into solution, at 60° for 3-4 days, until yellow vapour ceased to come off; the contents of the refort were then poured into much water; the insoluble matter was thoroughly washed by decantation, dried on a water-bath, and again oxidised by KClO, and HNO, Aq, as before. These operations were repeated (usually 4 times) until no further change was produced, and the insoluble matter formed a clear-yellow solid. Analysis of this yellow solid, dried at 100°, gave the formula C₁₁H₁O₃. This body-called graphitic acid by Brodic forms small, transparent, lustrous, yellow plates; it is slightly soluble in water; ins luble in water containing acids or salts; turns blue litmus slightly red; shaken with solutions of alkaline bases it appears to form insoluble salts, but the composition of these is very uncertain; when heated it burns explosively, leaving a fine, black residue; it is easily decomposed by reducing agents such as (NH.)HS. SnCl., HIAq. &c. (v. infra). Brodie supposed this body to be a compound of a hypothetical element which he called graphon, and to which he gave the atomic wyight 38; he formulated graphitic acid as Gr. 11,0, and regarded it as the carbon analogue of a silicia acid Si,H.O. obtained by Wöhler from graphitoidal silicon. Gottschalk (J. pr. 95, 321) placed a very intimate mixture of 1 part (50 grms.) purified, very finely divided graphite with 3 parts KOlO, in a large

slowly added enough HNO Aq, S.G. 1:525, to completely moisten the whole; he then digested at 50° to 60°, and then at 60° to 70°, for 25-80 hours; he poured off the greater part of the acid and dissolved KNO, washed with hot water by decantation, dried in vacuo and then at 100°; he repeated this treatment 5 or 6 times; finally he washed the residue with HNO, Aq S.G. 1.28, removed the acid by pressing between paper and then by washing with alcohol, washed with other to remove alcohol, pressed again, and dried on the water-bath in the dark. Gottschalk's analyses lead to the formula $C_{11}H_1O_6$ for graphitic acid; he describes a salt, $C_{23}H_{10}K_2O_{18}$, obtained by treating with conc. KOHAq and

washing with cold water.

The action of KClO3 and HNG3 on graphite has also been investigated by Stingl (B. 6, 391), and by Berthelot (A. Ch. [4] 19, 399). Berthelot calls the compound produced as described graphitic oxide, he says it does not react as an acid; he calls the carbon-like mass left on heating graphitic oxide pyrographitic oxide; the body is completely dissolved by heating with KClO, and HNO₃. The porons, amorphous, insoluble body obtained by heating 1 part graphitic oxide with 20 parts HIAq S.G. 20 to 280°, Berthelot calls hydrographitic oxide; this body is not explosively decomposed by heating, treated with KClO, and HNO, it yields graphitic oxide. There appear to be differences between the graphitic acids obtained from different kinds of graphite. Berthelot distinguishes the three allotropic forms of earbon, by their reactions with K(30, and conc. HNO3; amorphous carbon is oxidised to brown humnslike bodies, which dissolve in water; graphite forms graphitic acid; diamond is unchanged. -10. Both graphite and amorphous carbon are said to yield mellitic acid C, (CO2H), by the action of K2Mn2O8 in KOHAq (Schulze, B. 4, 802). - II. Carbon combines with many metals whon strongly heated with them, e.g. with Fe, Ni. Co, &c.; none of these carbides has been isolated as a pure compound (v. Careides).

Carbon, halogen compounds of. does not combine directly with the halfgens. These compounds are represented by the formulæ CX_{\bullet} , C_2X_{\bullet} , and C_2X_{\bullet} , where X=Br or Cl; when X=I only CX_{\bullet} is known: no fluoride of C has been isolated. The chlorides have been gasified and V.D. of each determined; the formulæ are therefore molecular. The bromides decompose, partially or wholly, when heated: the formulæ are probably molecular. The iodals is easily separated by heat into C and I. The methods of preparation, and reactions, of the chlorides and bromides are very similar. [C,Cl'] = 21,030; $[C^2,Cl^4] = -1,150$ (at const. press. Thomsen). Besides these compounds, several bromochlorides of oarbon exist; CBrCl₃; two isomerio C₂Br₂Cl₄, C₂Br₂Cl₂, C₂Br₂Cl₂, C₂Br₂Cl. [For more details of the halogen compounds of carbon v. the halogen derivatives of ETHANE, ETHYLENE, and METHANE.]

CARBON BROMIDES. Carbon dibromide C.Br. (Tetrabromethylene). White crystals; M.P. 53°; produced by heating C.Br., or better, by reducing C.Br. with Zn and H.SO, Aq; also by reacting with Br on alcohol or ether, adding

treating C.HBr., Br. with alcoholic KOH (Lennox, C. J. 14, 209). Decomposed by hot Zn, Cu, Fe, ZnO, CuO, &c., giving metallic bromide and C or CO, (Lowig, A. 3, 292).

Carbon tribromide C.Br. (Hexabromide Tetrabromethylene dibromide). Hard, rectangular prisms, easily soluble in CS, insoluble in alcohol or ether; decomposed to C,Br, and Br, at 200°. Produced by brominating C.H.Br., and by heating C.H.Br with Br and H.O to 170°-180° (Reboul, A. 124, 271).

Carbon tetrabromide CBr. (Tetrabromo-methane). White lustrous tables; S.G. 3·42; M.P. 91°. B.P. 189·5° (at 760 mm.) with partial decomposition. Insol. in water, very sol. in alcohol, ether, or CHCl. Partially decomposed with liberation of Br, at 200°; with alcohol at 100° gives HBr, CHBr, and CH, CHO; with alcoholic NH, at 100° gives CHBr, and a little guanidine. Formed by the reaction botween (1) Br, in presence of I or SbBr3, and CHBr3 or CS, or CBr₃(NO₂); (2) Br, in presence of I, and CIICl. Best prepared by heating 1 part CS2 with 11 parts I and 7 parts Br to 150°-160° for 48 hours in a closed tube, shaking contents of tube with NaOHAq, distilling in steam, pressing between paper, and crystallising from alcohol (Bolas a. Groves, C. J. [2] 8, 16I; 9, 773).

CARBON CHLORIDES. Carbon dichloride C₂Cl₄ (Tetrachlorethylene). Colourless liquid; cthereal odour; S.G. at 10° 1·62 (R.), 1·612 (R.); S.G. at 0° 1·6595 (R.), B.P. 122° (R.), 116·7° (G.), 121° (B.). V.D. 5·82. Easily combines with Cl in sunlight forming C₂Cl₄. Prepared by reducing C₂Cl₆; C₂Cl₆ is placed in a flask with water and Zn, Il₂SO₄Aq is added from time to time, the flask being kept cold and frequently shaken; the C.Cl. is distilled over in steam, dried, and fractionated (Faraday, T. 1821. 47; Regnault, A. Ch. J., 377; Geuther, A. 107, 212; Bourgoin, Bl. 23, 344).

C.Cl. (Tetra-Carbo trichloride chlorethylene dichloride. Carbon hexachloride). Hard, colourless, rhombio prisms; S.G. 2.0. M.P. 187° and B.P. the same (Städel a. Hahn, B. 9, 1735). V.D. 8.15. Insol. in H.O. sol. in alcohol or ether. Easily reduced, e.g. by Zu and II.SO,Aq, or by alcoholic KHS, to C₂Cl₄; with KOHAq at 200° gives KCl, H₂O, and K₂C₂O₃. Prepared by leading Cl into boiling C2H4Cl2 till saturated, cooling by ice, prossing between paper, dissolving in alcohol, ppg. by H.O. pressing, and eryställising from alcohol (Faraday, T. 1821. 47; Regnault, A. Ch. 69, 165; 71, 371; Liebig, A. I, 219; Geuther, A. 60, 247; Berthelot, A. 109, 118).

Carbon tetrachloride CCl. chloro-methane). Colourless liquid, with ethereal odour; S.G. 30 1.63195; B.P. 76.74° (Thorpe, C. J. 37, 199). V.D. 5.24. Prepared by leading dry CI into boiling, CHCl, centaining a little SbCl, or IOI, in a large flask with inverted condenser, removing excess of Cl by shaking with Hg, and fractionating. Also by passing CS. and Ci through a lot porcelain tube (Kolbe, A. 45, 41; 54, 146). Unchanged by KOHAq; with alcoholic KOH slowly gives KCl, K₂CO, and H₂O; passed through a hot tube gives C₂Cl₄ reacting with Br on alcohol or ether, adding C.Cl., and C; heated with SO, gives COCl, and ROHAQ to remove HBr, and distilling; or by S.O.Cl.; with P.O. gives POCl, and COCl.

CARBON. 680

(Regnault, A. Ch. 71, 877; Dumas, A. Ch. 73,

CARBON IODINS. CI. Dark red octahedra; S.G. 22 4 32; sol. in alcohol, einer, or CS.. Decomposed by heat to C and I; boiled with H₂O or dilute HIAq gives CHI₃. Prepared by mixing equal vols. CCI, and CS, with saturated solution of Al₂I₆ in CS₂, then diluting with H₂O out of contact with air. The solution of Al, I, is prepared by placing the proper quantities of Al (in small pieces) and I in a stoppered flask and adding 3 times the quantity of CS2 (Gustavson B. 14, 1705).

CARBON BROMOCHLORIDES: Trichlorobromo-methane CCl₃Br; two letrachlorodibromo-ethanes C.Cl.Br.; dieblorotetrabromo-ethane C.Cl.Br.; chloropentabromo-ethane C.ClBr, ; dichlorodibromo-ethylen C.Cl. Br.; chlorotribromo-ethylene C.CIBr. (v. these compounds under METHANE, ETHANE, and ETHYLENE).

Carbon, hydrate of (?) By treating pig-iron with (1) CuSO, Aq, (2) Fc, Cl, Aq containing IICl, a brownish-black substance remains, containing, according to Schützenberger a. Bourgeois (C. R. 80, 911) carbon and water in the ratio 11C:3HO. Besides the C and HO, the substance gives about 10 p.c. ash. It loses 311,0 at 250°

Carbon nitride = Cyanogen (q, v).

Carbon, oxides of. Two oxides certainly exist, CO and CO.; these formula are molecular; each bears the relation of anhydride to an acid, CO is formic anhydride (the acid is II, CO2), CO2 is carbonic anhydride (the acid is HCO3) (v. infra). Both oxides can be obtained by direct combination of O with C; either can be produced from the other, by combining with O or with C, respectively. Both we stable gases; CO is an energetic reducer; CO, in a few cases acts as an oxidiser. Brodie (Pr. 21, 245) and Berthelot (Bl. [2] 21, 102) have described bodies, p. sheed by the induced electric discharge on CO, as oxides of C. Brodie goticed a gradual diminution in vol. of the CO and the formation of a red brown film on the glass tube; the solid was soluble in water giving a markedly acid solution; its composition appeared to differ in different experiments; Brodie gives the for-Berthelot got brown, mulae C,O, and C,O, amorphous, humus-like bodies which dissolved in water with acid reactions, gave brown pps. with AgNO₃Aq, BaOAq, and Pb2NO₃Aq; at 300°-400° CO and CO. (equal vols.) were evolved, and another dark body remained, to which B. gave the composition C₂O₃. B. also (A. Ch. [5] 17, 142) states that by the action of electric sparks on pure CO, a gas was produced which reacted violently with Hg and oxidisable bodies.

CARBON MONOXIDE. CO. (Carbonic soxide; CARBON MONOXIDE. CO. (Carbonic soxide; more properly, although rarely, carbonous oxide; formic anhydride.) Mol. w. 27-93. S.G. -9078 (air = 1). V.D. 14. (c. -186°) (Wroblewski, C. R. 98, 982). S.H. p. 2346. S.H. v. 16844 (E. Wiedemann, P. 157, 1). Z.E. -003667 (Regnant). S. (6°) 0287; (9°) 0269; (13-5°) 02315 (Bunsen). S. alcohol (2°) 20356; (13°) 20416; (16°) 20566; (24°) 20452, (17°) 1, 1000301; 4 $\mu_0 = 1.000301$; $\mu_E = 1.000350$; $\mu_0 =$ (Bunsen). 1.000391 (Croullebois, A. Ch. [4] 20, 136). [CO, O] = 67,960 at const. press., and 67,670 at

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tively (Thomsen). Does not exactly obey Boyle's law; $\frac{PV}{P_1V_1} = 1.00293$ (Regnault, Acad. 1862. 26,

229). Liquefied by cooling to -136° at pressure of 200-300 atmos, and then decreasing pressure, not too cnickly, to not lesythan 50 atmos. (Wroblewski a. Olszewski, A. Ch. [6] 1, 112; v. also Natterny, W. A. B. 12, 199; and Cailletet, C. R. 85, 1213 a. 1217, and A. Ch. [5] 15, 132). First obtained in 1776, by Lasonne, by heating C with ZnO; obtained by Priestly, in 1796, by heating charcoal with iron oxide, but supposed by him to be H; proved by Cruickshank not to be a? hydrocarbon; true composition determined by Occurrence. In the gases from burning coal

or charcoal; from the partial combustion or putrefaction of organic matter; or from the reduction of metallic oxides by charcoal, e.g. in the blast-furnace (Bareswil, J. Ph. [3] 25, 172; Bansen, P. 46, 193; 50, 81). During the oxidation of gallic and tannic acids by exposure to air in alkaline solutions (Boussingault, A. Ch. [3] 66, 295; Calvert, C. R. 57, 873). In pig-iron and steel according to Troost a. Hautefeuille, also

Parry (J. 1873, 997; 1874, 1083).

Formation. - 1. By passing steam over excess of red-hot C; the product may contain about 281 p.c. CO, 561 p.c. II, 141 p.c. CO2, and traces of CII, (v. Naumann a. Pistor, B. 18, 164). 2. By passing a slow current of CO, over red-hot C, and washing the gases through KOHAq and soda-lime,-3. By heating CO2 with those metallic oxides which do Ast readily part with O, e.g. ZnO, PbO, Pe.O. oxides which readily give up O yield but little CO, as it is again oxidised to CO₂, -4. By passing CO₂ over red-hot Cu, or over hot Zn-dust (Noack, B. 16, 75), -5. By heating CO₂ to 1300° (Deville, C. R. 59, 873),— 6. By electric sporks through CO2 (Buff a. Hofmann, A. 113, 140).— 7. By heating powdered CaCO₃ or K₂CO₃ with ope-sixth its weight of powdered charcoal; Na SO₄ heated with C also yields CO (with Na S).—S. In very small quantities (with COS) by passing CO, and S vapour through a red-hot tube (Berthelot, A. Ch. [5] 30, 547).—9. By heating dry 11, C,O,, or by reaction between H₂C₂O₄ or an oxulate and hot conc. H₂SO₄.—10. By heating 11.CO₂11, or a formate, with cone. If SOc.

Preparation. - 1. One pt. dry powdered K, Fe(CN), is heated, in a capacious vessel, with 8-10 its. conc. 11.80; as soon as frothing begins the lamp is lowered to a small flame; the gas is passed through milk of lime and KOHAq, to remove CO, and the SO, formed in the process; SO, is evolved only in the oarlier the process, so, is evolved only if the owner stages of the reaction (Grinnia a Raindohr, A. 98, 127). 15 g. K, $Fc(CN)_s$ yield about 4 litros CO: $K_1Fc(CN)_s + 6H_2SO_4 + 6H_2O$ = $6CO + 2K_1SO_4 + 3(NH_1)_2SO_4 + FcSO_4$ (Fownes).

2. Dry CaC₂O₃, or BaC₂O₄, is mixed with about the pt. dry CuO.II., and the mixture is strongly heated in a hard glass Assk; the gas is passed through milk of lime, and is then dried:—CaC₂O₃ gives CaCO₃ + CO; the CaH₂O₂ absorbs any CO2 formed .- 3. According to Chorrier (C. R. 69, 138) pure CO may be prepared by passing the gases produced by heating H2C,O const. vol.; [C,O] = 29,000, and 29,290, respectivith H,SO, through a red kot tube filled with

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charcoal, and then through a mixture of CaOAq oxidised. L. Meyer's experiments (B. 19, 1099), and KOHAq.

Properties .- A colourless, tasteless, slightly odorous gas; liquefied at low temperature and great pressure (v. supra). CO is combustible but a non-supporter of combustion: the temperature of the flamt of CO in air as about 1400° (Valerius, J. 1874. 58). Absorbed by C, and by several metals, e.g. K, Ag, Au; equickly absorbed by Cu,Cl, in a little HClAq (v. infra); decomposed at very high temperature to C and CO,; decomposed when moist by inductionsparks; CO is an energetic reducer; it combines with moist KOH (or NaOH) to form K formate; combines directly with Cl and Br in sunlight. CO is extremely poisonous; it removes O from the blood and combines with the hamoglobin. CO may be detected in the blood by observing the absorption-spectrum; this is almost identical with that of oxygenated blood, and is characterised by two bands between D and E; on adding a little ammonium sulphide these bands disappear in the case of oxygenated blood, and the spectrum shows one band midway between D and E; if the blood contains CO the two bands remain unchanged for several days (Vogel, B. 11, 235; Hoppe-Seyler, Fr. 3, 439).

Reactions. -1. Electric sparks cause a partial decomposition to CO, and C; if the CO, is removed the change proceeds (Berthelot, A. Ch. [5] 30, 547). According to Berthelot (Bl. [2] 21, 102; A. Ch. [5] 17, 142) CO is decomposed by the induction-discharge, with production of CO, and (?) C,O, and C,O, (v. ante; beginning of this art.). According to T aff a. Hofmann (C. J. 12, 282) the induction spalk does not decompose dry pure CO. Dixon (C. J. 49, 103) found that CO was decomposed (only about 1 p.c. of the total gas) by sparks from a Leyden jar .- 2. Heated to about 1300° CO is partially decomposed to C and CO, (Deville, C. R. 59, 873).-3. A mixture of CO with oxygen is brint to CO₂ by application of a flame or electric sparks. Dixon (T. 1884. 617) has proved that if both gases are perfectly dry no chemical change occurs when a spark is passed; that a mere trace of steam renders the mixture explosive; that the exidation of CO by O takes place very slowly if only a very small quantity of steam is present; and that as the quantity of steam is increased the rapidity of the explosion is increased also. The steam acts as a carrier of O to the CO; it is probably reduced, and the II is then again oxidised: the reactions which econf are very probably these (v. Dixon, C. J. $49,^{1}91$):

 $(2CO + 2H_2O = 2CO_2 + 2H_2)$ $(2H_1 + O_2 = 2H_2O)$

Or (Armstrong, C. J. 49, 112) the changes may be represented by the formule, before explosion O.H.O.CO; after explosion O.H.O.CO. Small quantities of gases other than H.O were tried (H.S. C.H., H.CO., N.H., C.H., HCl; SO., CS., CO., N.O. C.N., CCl.); if the gas contained H. explosion becurred; i the gas did not contain H the mixture did not explosion-decurred; in the gas did not contain H the mixture of CO and steam is heated to about 600°, a portion of the CO is oxidised to CO.; the amount of CO oxidised depends on the conditions (v. Dixon, C. J. 49, 94; references to other memoirs are given); if the CO. is romoved as it is formed the whole of the CO can be

however, seem to prove that a mixture of dry CO and O can be exploded it a very strong spark is used, and the temperature is thus made very high. The gases must be under considerable pressure; the more dilnte the gaseous mixture tho more difficult is it to explode it .- 5 When sparks from an induction-coil are passed through a mixture of CO and steam, CO, s little formic acid, and in some cases C, are formed (Dixon, C. J. 49, 94).—6. When to t mixture of dry CO with hydrogen, oxygen in sufficient for complete combustion is added, and the mixture is exploded by the spark, CO, and H2O are formed; the ratio of CO, to H2O depends on the shape of the vessel, and the pressure up to a certain limit; above this pressure - the 'critical pressure'—tho catio CO2:H2O is independent of the shape of the vessel. The larger the quantity of O nsed the lower is the critical pressure. So long as the volume of H is more than twice that of the O the ratio of $CO \times H_2O:CO_2 \times II_2$ remains constant, provided no H2O can condense, and the pressure is above the critical pressure: when the vol. of H is less than twice that of 6 the value of the ratio diminishes. The presence of an inort gas, c.g. N, increases the formation of CO, and diminishes that of H2O, hence it lowers the value of the ratio $CO \times H_2O:CO_2 \times H_2$. This ratio is called by Dixon the co-efficient of affinity of the reaction (v. Dixon, T. 1884, 617; C. J. 49, 94; Horstinann, B. 12, 64; v. also Chemical Chance).-7. CO is oxidised to CO₂ (1) by bichrome and sulphuric acid (Ludvig, J. 1872. 218); (2) by palladium charged with hydrogen, in presence of oxygen and water, H.O. being also produced oxygen and water, 11,02 teing also produced (Traube, B. 15, 2325, 2854; 16, 123; Remsen a. Keiser, B. 17, 83); (3) by mixing with oxygen and passing over latinum-black; (4) by nitrogen dioxide [NO.] (Hasenbach, J. pr. [21 4 1); (5) by he ting with most metallic oxiles; (6) by heating with many oxysalts, e.g. alkaline snlphates (sniphides produced) .- 8. Many experiments have been made to determine whether CO is oxidised by contact with moist oxygen in presence of slowly oxidising phosphorus; the balance of evidence seems to show that CO, is not produced (Remsen (and others) Am. S. [3] 11, 316; B. 17, 83; Am. 6, 153; Leods, B. 12, 1836; C. N. 48, 25; Baumann, B. 16, 2146; 17, 283).—9. CO reacts with moist potash or soda to form alkali formato (Berthelot, A. Ch. [3] 61, 463); the reaction proceeds most quickly at 190° 200°, and is best accomplished by leading moist CO over soda-lime (Fröhlich a. Geuther, A. * 202, 317) .- 10. With ferrous oxide at 300° 400°, CO, and a little C are formed (Griner, C. R. 73, 281) .- 11. CO appears to react with ecrtain metallic peroxides to form carbonates, but, according to Wright a. Luff (C. J. 33, 540), CO, is formed by partial reduction of the peroxide and reacts with the lower oxide to produce carbonate .-- 12. Mr. ny of the preceding reactions exhibit CO as a reducing agent; it also reduces PdCl.Aq to Pd.—13. When sodium or potassium is heated to redness in CO, alkali carbonate and C are formed.

Combinations.—1. With potassium at about 80° to form the explosive compound KCO (Brodie, C. J. [2] 12, 269), v. Potassium.—

2. With chloring or broming in sunlight, to form COCI, or. COBr. (v. CARBON, OXYCHLORIDE, and OXYBROMIDE, out. 8. With sulphur to form COS (v. Carbon, Oxysulphide ork.—4. With platinic chloride to form C₃O₂PtCl, and C₂O₂PtCl, (Schützenberger, A. Ch. [4] 21, 350).—5. CO is absorbed by anhydrous HON (Böttinger, B. 10, 1122); by several metals, e.g. Fe, Ag, Au; by carbon.—6. CO does not combine with eyanogen, nor does it react with Mg(CN)₂.

Estimation.—CO in a gaseous mixture is

absorbed by Cu.Cl₂ solution. Thomas (C. N. 37, 6) prepares the solution by filling a vessel of 120 c.o. capacity; full of Cu turnings, adding 6 g. crystallised CuCl₂ and 20 c.c. cone. HClAq, and shaking until solution of the CuCl, is effected; he then adds 30 c.c. water and shakes briskly for some time, and then adds

30 0.0 water.

OARBON TOXIDE. CO. (Carbonic anhydride, often called carbonic acid.) Mol. w. 43·89. S.G. gas 1·53; S.G. liquid 1·0ε7·at - 34°; 1·016 at - 25°; '966 at - 11·5°; '91 at - 1·6°; '84 at + 11°; '726 at + 22·2° (Cailletet a. Mathias, C. R. 102, 1202). S.G. solid (hammered) slightly under 1·2 (Landolt, B. 17, 309) [-65°] (Mitchell); [-57°] (Faraday); [-78·2°] (Regnault, A. Ch. [3] 26, 257). V.D. 22; 22·42 at 800°; 21·2 at 180° (Movor a. Goldsehmidt, B. 15, 1165). 1180° (Meyor a. Goldschmidt, B. 15, 1165). S.H.v. 33 (equal vol. of air = 1), 2169 (equal weight of air = 1) (Regnault, C. R. 36, 676, &c.;

v. also Wiedemann, P. 157, 21). S.11.v.

to 1.305 (Amagat, Röntgen, C. R. 71, 336; 77, 1325). C.E. 0037 (Regnault, Magnus, Joly). =1.00722 (Reguault, C. R. 20, 975). P.V. 200° CO. obeys Boyle's law (Amagat, C. R. 68, 1170; 73, 183). C.E. liquid CO. very large, 120° C. at -20° become 150 vol:, at +30, (Thiloriei, A. Ch. 60, 427). Critical temperature = 30°9° (Andrews, T. 1869, 575). Vi.pourpressure of liquid CO₂ (Regnault) in atmospheres: -25°, 17·1; -5°, 30°9; 0°, 35·4; +5°, 40·5; 15°, 52·2; 25°, 66; 35°, 82·2; 45°, 100·4. Vapour-pressure of solid CO2 (Fariday) in atmos.: -57° , 5.33; -70.5° , 2.2; -99.4° , 1.14. B.P. of solid CO, -i.e. temp. at which vapour-pressure =760 mm.—is much lower than the M.P.; Regnault (and Pouillet) found -78° to -79° (P. 77, 107); Thilorier, -95° to -98°; and Faraday, as shown by values for vapour pressure, nndor -99°. By evaporation of solid CO2 mixed with ether, temp. is c. - 100°.

S	. CO _z gas (Bu	nsen, A. 93, 1):
At .0°	1 ·7967	At 11° 1-1116
1	1.7207	l 12 1·1018
2	1.6481	13 1-0653
3	1.5787	• 11 1.0321
4	1.5126	15 1.0020
5	1.4497	■ 16 0.9753 ■
6	1.3901	17 0.9519
7	1.3339	€18 0.9318 •
8	1.2809	19 0-9150
9	1.2311	20 0.9014
10	1.1847	

Absorption-coefficient = $1.7967 - .07761t + .0016424t^2$ S. CO, gas in alcohol (Bunsen):

At 3:20' 4:0442 At 14.2° 3.2357 6.8 3.7374 18 3.0391 22.6 2.8277 10.4 3.4875

Absorption-coefficient = $4.32955 - 0.09325t + 0.00121t^2$.

95,960; [CO,O] = 67,960. H.F.v. [C,O²] = 96,960; [CO,O] = 67,670. [C,O',Aq] = 102,840. [CO, O, Aq] = 73,840.[CO',Aq] = 5,880.

[CO)Aq,nNaQ11Aq]; n=1=11,016; n=2=20,184; n=4=20,592 (Thomson).

Carbon digxide has been known for conturies. The identity of the gases produced during fermentation and by the action of acids on chalk was established by Black. Bergmann recognised the same gas in the atmosphere. Cavendish proved that the same gas was produced by burning charcoal. Lavoisior established the composition of the gas. Faraday liquefied, and Thilorier soliditied, carbon dioxide.

Occurrence.-In the atmosphere (v. Atmo-SPHERE); in mineral waters: issues from the earth in different places; sometimes found liquid in eavities in quartz, &c. Produced by the breathing of animals, by the decay of organio matter, by the combustion of coal, charcoal, &c. In combination as carbonate, of calcium, magnesinm, &c., &c.

Formation .- 1. By burning C in air or O .-2. By oxidation of , nost C-compounds.—3. By burning CC.—4. By reducing many metallio oxides by C.—5. By heating together H.O and CO .- 6. By the reaction between red hot C and steam.-7. By the action of steam on CaCO, at red heat.—8. By heating a mixture of K₂Cr₂O, with Na₂CO₃.—9. By heating several carbonates. 10. During fermentation. -11. By reaction between acids and carbon tes.

MgCO_a in lumps is Preparation. -- CaCO₃ treated with dilute HClAq at the ordinary temperature; the gas is passed through NaHCO3Aq (so remove HCl which may have passed over), and is then dried by CaCl₂. Bunsen recom-mends the use of finely powdered chalk and cone. H2SO4, and addition of a very little water.

· Liquid carbon dioxide was obtained by Faraday by decomposing (NII), CO, by II, SO, Aq in one limb of closed glass tube bent at an obuse angle. Thilorier (A. Ch. 60, 247) decom-poses NaIICO, by dilute II, SC, Aq in an iron vessel connected with an iron cylinder in which the CO, is liquefied by its own pressure. Natterer (J. pr. 35, 169) compresses CO, by a specially constructed air-pump (v. also Gore, T. 1861. 63).

Solid car on dioxide is obtained by allowing the liquid to escape into a tin vessel; part of the liquid becomes gas and part is solidified Landolt allows the liquid to evaporate freely into conical woollen balls; he then compresses the solid CO₂ in content moulds of hard wood by wooden pistons (B. 17, 309).

Properties .- A heavy, colourless, gas; incombustible; non-supporter of ordinary combustion, but strongly heated K or Na, or brightly burning Mg, burns in CO2. Absorbed by water, solution colours litmus wine-red and reacts as a weak acid (v. Carbonic acid). Absorbed by moist alkalis and alkaline earths forming carbonates; rapidly absorbed by mixture of powdered KOH and hydratod Na₂SO₄. Poisonous, by

cutting off supply of O.

Liquid carbon diexide is a limpid, colourless, refractive, liquid; nonconductor of electricity; not changed by strong induction-sparks; very expansible by heat; C.E. is greater than that of the gas., Insol. in water which swims on the surface; mixes with alcohol, ether, &c. Does not dissolve S or P; dissolves I; no reaction with Na or K (Caillett, C. R. 75, 1271).

Solid carbon diexide is a white, loose, snowlike, solid; when compressed by frammering in wooden moulds it resembles chalk (Landolt, B. 17, 309). Very bad conductor of heat. Evaporates slowly, a specimen prepared by Landolt 53 mm. by 71 mm. diam. took 5 hours to volatiliso in the air. Burns, if pressed against

the skin.

Reactions.—1. Heated to c. 1300° in porcelain tube is partly changed to CO and O (Deville, C. R. 56, 729; v. also Berthelot, C. R. 68, 1035) .- 2. Partly decomposed by electric sparks; a condition of equilibrium is attained when change of CO, into CO + O equals that of CO and O into CO₂ (Dixon a. Lowe, C. J. 47, 571).— 3. Mixed with hydrogen, and heated to bright redness or submitted to induction-sparks, II_O and CO are formed; if H₂O is removed the whole of the CO2 goes to CO (Dixon, C. J. 49, 94). According to Dubrunfaut (C. R. 74, 125) CO, and II passed over hot purice give Cand H₂O, -4. A michare of two bon dioxide and sulphur vapour passed through a red-hot tube yield a little COS, CO, and SO₂ (Berthelot, BL. [2] 40, 362).-5. With sulphurcited hydrogen, passed through red-hot tube, forms CO, ILO and S (Köhler, B.11, 205).—6. Decomposed by chlorophytl-parts of plants in sunshine .- 7. Reduced to CO by heating with carbon, iron or zinc, or with copped which has occluded hydrogen (Tissandier, C. R. 74, 531; Schrötter, W. A. B. 34, 27) .- 8. Partly reduced to CO by reaction with ferrous sulphate and a little water, in a closed tube (Horsford, B. 6, 1390). - 9. Reduced to C by heating strongly with sodium, potassium, or magnesium; alkali carbonates strongly heated with phosphorus or boron give CO, which is reduced to C (Tennant, Crellis A. [1793] 1, 158; Pragendorff, J. 1861.111; Leods, B. 12, 1834 a. 2131).— 10. With moist alkalis, or alkaline earlhs, forms carbonates.-11. With water probably forms a solution of carbonic acid, H.CO. (v. CARBONIC ACID) .- 12. With sodium- or potassium-amalgam at c. 350° gives Na (or K) oxalate (Drechsel, A. 146, 141).-13. With sodium CO2Aq reacts to give Na formate (Kolbe a. Schmitt, A. 119, 251). 14. Decomposes moist potassium iodide at high temperature giving III (Paparogli, G. 1881.227).

Carben, exybremido of. The existence of a Br compound of CO analogous to COCl₂ is doubtful. A mixture of Br vapour with excess of CO is slowly, but not fully, decolourised in sunlight; it contact with KOHAq thus gas produces KBr and K₂CO₂ (Schiel, A. Suppl. 2, 311). Emmerling a Lengyel could not obtain a trace of any compound of C, Br, and O, by the reaction between COS and Br at a high temperature (B 2, 547). By the reaction between

H₂SO₄ (50 parts), K₂Cr₂O₇ (20-25 parts), and CHBr₂ (5-10 parts), Emmerling (B. 13, 874) obtained a small quantity of a liquid, which he slowly distilled through Sb, to Femove Br; he thus obtained a colourless heavy liquid, smelling like COCl₂. The B.P. rose from 12° to 30°; analyses seemed to show that the liquid was a mixture of COCl₂ and C oxypromide.

Carbon, exychloride of. COCl. (Carbonyl chloride. Phosgenegga. Chloro-carbonic acul) Mol. w. 98·67. (8·2° at 756 mm.). S.G. \(\frac{9}{2} \) (liquid) 1·432; \(\frac{1850}{2} \) 1·392 (Emmerling a. Lengyel, \(A. \) Suppl. 7, 101). V.D. 50·6 (E. a. L.). [C.O.Cl'] = 54.850 at constant volume; 55.140 at constant pressure (Thomsen). First prepared by J. Davy in 1811 (T. 1812. 144) by the action of sunlight on Cl + CO (hence the name phosgene).

Formation.—1. By leading CO into boiling SbCl₃ (Hofmann, A. 70, 139; v. also Butlerow, N. 1863. 484; Kraut, Gm.-K. I. 2, 386), or over hot PbCl₂ or AgCl (Göbcl, J. pr. 6, 388).—2. By leating CCl₄ with ZnO at 200° in a closed tube; or by passing CCl₄ and CO through pumice in a tube heated to about 400°.—3. By heating CHCl₂ (1 part), K.Cr.₂O₇ (2½ parts), and H.SO₄ (10 parts) at 100°, and passing the gas over Sb to absorb Cl (E. a. L.).—4. By passing Cl and CO over Pt black at about 400° (Schützenberger, Bl. [2] 10, 188; 12, 198).—5. By passing Cl and CO. over hot C (Schiel, J. pr. 6, 388). (For other methods v. Schützenberger, B. 2, 218; Dewar a. Cranston, C. N. 22, 174; Armstrong, B. 3, 730.)

Preparation.—Drv Cl and dry CO are slowly passed through a succession of large bottles freely exposed to sunlight, then through a Uttobo loosely filled with pieces of Sb (to remove free Cl), and finally into a tube surrounded by snow and salt. Hind gas should pass through the drying-bottles at as nearly as may 1. the same rate! 100 litres CO give 140-14.5 g. COCl. in diver. sunlight. Paterno (G. 5, 233) passes the mixed gases through a tube 400 mm. long filled with animal charcoal: combination occurs with production of heat; the tube must be cooled by a wet cloth from time to time (v. also Wilm a. Wischin, A. 147, 150).

Properties.—Colourless gas with penetrating odour; at 8° and under it is a colourless limpid liquid; the gas is soluble in acetic acid, benzene, and several liquid hydrocarbous.

Reactions.—1. Water absorbs COCl₂ with formation of COAq and HCIAq. Berthelot (C. R.87, 591) gives the value [COCl², Aq] = 64,600. 2. Alcohol forms chlorocarbonic elher CO.Cl.OEt. (Q. t.).—3. Several metals decompose COCl₂, when heated with it, to CO and metalic chloride; e.g. Sb, Ås, Na, Sn, Zn; potassium forms KCl, K₂CO₃, and C.—4. With slightly moist potassium carbonate, KCl, H₂O, and CO₂ are formed.—5. Zina oxide produces ZnCl₂ and CO₂.—6. Combines with 4 vols. amnonia to form urea and NH,Cl (Natanson, A. 98, 288; Fenton, C. J. 35, 793).

Carbon, exysulphide of. COS. (Carbonyl sulphide.) Mol. w. 59-91. V.D. 30-4. [CO,S] = 8,030; [CO,S] = 37,030; [COS,O] = 181,010 (Thomsen).

Occurrence. - According to Thom (A. Suppl.

5, 236), in several mineral waters, and in volcanic gases.

Formation 1. By passing CO and S vapour through a red-hot porcelary tube (Thom).—

2. By gently hoating SO, with CS; SO, and S also produced (Armstrong, B. 2, 712).—3. By action of CO, on boiling S; or by electric sparks on CO, mixed with S vapour (Cossa, B. 1, 117, on Co, in see with 5 vapout (cossa, B. 1, A.7), Chevrier, C. R. 69, 136).—2. By leading alcohol and CS, over red-hot Cu (C, ruedley, C. J. [2] 13, 523). For other methods b. Ladenburg, B. 1, 273; 2, 30, 53, 271; Dewar a. Cranston, C. N. 20, 174; Salomon, J. pr. [2] 5,476.

Preparation.—By decomposing KCNS by SO, Aq; KCNS + H.O + 211, SO, Aq H₂SO₄Âq; eCOS + KMSO, Aq + NH, IISO, Aq. KCNS 1s added to a cold mixture of 5 vols. H.SO, with 4 vols. II,O as long as the whole remains liquid; if much gas comes off the vessel is cooled, if very little gas is evolved the vessel is warmed gently. The gas is passed through three U tubes, containing (1) cotten wool charged with moist HgO (to remove HCN and formic acid) (2) cuttings of unvulcanired caoutchouc (to remove CS₂), (3) CaCl₂; the gas is then collected over Hg (Thom). Bender (A. 148, 137) recommends passing the gas through a tube surrounded by snow and salt, and Hofmann (B. 2, 73) through wool moistened with PEt3; the object in either

method being to remove CS.

Properties. - Colourless, heavy, gas, with a pleasant somewhat aromatic odour. Colours moist blue litiuus slightly reddish. Absorbed by water; solution sometimes centains GO, and

H.S. Very sol. in alcohol.

Reactions .- 1. Burns in air to CO and SO .. 2. At full red heat gives CS, and CO, (Berthelot, C. R. 87, 71). -3. With water gives CO Aq and H.SAq .- 4. With potash solution gives K.SAq and K₄CO₅Aq; similar reactions with MI Aq. and BaOAq.—5. Ammonia gas, or alcoholic H₄, gives CO.NH., SNH, (r. Berthelet, A. Ch. [5] 30, 539).-6. Solutions of sc'ts of copper, cadmium, lead, or silver give no pps., but on adding NH₃Aq the sulphides of the metals are ppd.—7. The gas is decomposed by hot mercury, copper, silver, and iron, giving sulphides; by hot sodium, giving Na.S, Na.CO., and C.

Carbon selsnide. Carbon and selenion do No definito compounds not combine directly. No definite compounds have been isolated. Rathke obtained a liquid which probably contained about 2 p.c. of a selenide of carbon (along with CCI,), by heating selenide of phosphorus with meist CCl, (v. A.

152, 181).

Carben, sulphides of. Carben disulphide, CS2, is a well-marked compound. A monosulphide, CS, probably exists. According to Low a sesquisulphide, C.S., can be obtained by the action of Na amalgam on CS. (Z. 9, 173; 10, 20). When sodium and CS. react

known. S.G. 166. CS, was exposed to sunlight for 2 months in a U tube of special construction; the solid which had formed on the walls of the tube was removed by water, a water bath) directly into the bottle in which it washed with CS2, and dried in H (Sidot, C. R. is to be preserved. It is kept in perfect dark

69, 1303; 74, 180; 81, 82). CS is a red powder; insol. in water, alcohol, turpentine, and benzene. Somewhat soluble in CS, or ether. Dissolved by HNO,Aq, not by HClAq or H,SO,Aq. At 200° gives C and S. Heated with S gives CS. CS is not produced by leading CS, over hot carbon or pumice, by leating Sb₂S, with C, by reaction between CO and H₂S, by reaction between CO and H₂S, by reaction between CH, and SO₂ or S.Cl., by heating CSD₂S, by beating Fe spiral in CS₂, by cleetric Sparks through CS₂ or by reaction between CSC₂ are by reaction between CSC₂ and the control of the contr through CS2, or by reaction between CSCl2 and hot Cu.

References .- Baudrimont, C. R. 44, 100 Berthelot, J. 1859, 83; Playfair, C. J. 13, 248; Buff a. Hofmon, A. 113, 129; Hermann, J. pr. 79, 448; Husemann, A. 117, 229; Kern, C. N.

33, 253; Rathke, A. 167, 195.

Carron Disulphide. CS₂. (Thiocarbonic anhydride. Sulphecarbonic acid.) Mol. w. 75'93. [e. -12'] (Wartha B. 3, 80). (16'04° at 760 mm.) (Thoppe, C. J. 37, 362; references in this paper to other determinations). S.G. 99 1.29215 (T.). V.D. 38. S.H. (liquid; 14°-29°) 2168 (Schüller, P. Ergzbd. 5, 116; v. also Hirn, A. Ch. [4] 10, 63 and 91). S.H. p. (equal mass of air =1) 1569; (equal vol. of air =1) 412 (Regnault). G.E. v. Thorpe (l.c.). [C, S²] const. press. = -26.310; const. vol. = -25.430; liquid = -19,610; [CS², O'] = 265,130 (Thomsen). For table of vapour-pressures from 0° to 50° v. Ramsay a. Young, C. J. 47, 653. $\mu_{\rm A}$ 1.6059; $\mu_{\rm A}$ 1.6729 (at 13°) (Kundt, W. 4, 34). For relations between volume as gas and pressure v. Herwig, P. 137, 15, 141, 83; 147, 16t.

Occurren. .. - In crus'e benzole; and in mustard oil. First prepared, in 1796, by Lampadius, by heating iron sulphide with charcoal. Composition was long uncertain. Clement a. Desormes (A. Ch. 42, 121) regarded it as a compound of C and S; it was also thought to be a compound of S and He and at other times of C, S, H, and N. Composition established by Vauquelin, Berzelius, and Marcet, in 1812 (v. G. A. 28, 427 a. 153; 48, 177; S. 9, 284; A. Ch. 83,

252). 4 Fermation.—1. By heating S with excess of C in a porcelain tube, condensing product in vessel surrounded by cold water, shaking with NaOHAq, drying by CaCl₂, and distilling from water bath.—2. By heating C with a metallio sutphide which gives off S at a high temperature, A. Cas or Sb.S. — 3 By heating wax, sugar, resin &c., with S.—4. By heating (CN)₂S. 5. By heating CCl₄ with P₂S, to 200° in a closed tube.

Preparation.—Commercial CS, is distilled off quicklime at 60°- 70°, leaving a little undistilled; the distillate is shaken in contact with powdered K_Mn_O₈, about b grains to 1 litre CS₂, for some time (to remove H₂S₁); it is then decanted and shaken thoroughly with Hg until fresh Hg is snaken thorongary with Tig that Hesh Tig has red-brown solid is obtained which according to Raub has the composition C₂S₂ (C. C. 1870. 579).

Carbon Monobuletibe. CS. Mol. w. uning residue when allowed to evaporate on filter paper; the CS, is poured off, allowed to stand in contact with CaCl,, and then distilled (from ness (Obach, J. pr. [2] 26, 281; for other methode v. Sidot, C. R. 69, 1303; Friedburg, B. 8, 1616; 9, 127; Millon, J. 1868. 928; Cloëz, C. R. 69, 1356).

Properties.—Colourless, limpid, highly re-fractive, liquid; ethereal odour when quite pure; vapour even when much diluted is poisonous, it stops fermentation (v. Cloëz, C. R. 63, 185). According to Wartha (B. 3, 80; 4, 180) CS, may be collisied by placing a small quantity in the veceel of a Carré freezing machine, oxhausting the air, and then opening the stopcock while continuing to oxhaust; in large quantities solid CS2 is obtained by mixing with absolute ether and exhausting the air by a Carré machine. Wartha also obtained a snow-like solid by blowing dry air through CS2 at the Erdinary temperature; according to Ballo (B. 4, 118) this body is a hydrate of CS₂; Berthelot (A. Ch. [3] 46, 490) and Duclaux (C. R. 64, 1099) obtained such a hydrate (probably 2CS_4I2O) by evaporating CS, in moist air (v. also Venables, Am. 5, 15). CS, vapour is very easily inflammable, ignition-temp. = 149° (Frankland, C. N. 6, 3), 170° (Braun); with air or O it forms a very explosive mixture; mixed with NO and ignited it burns instantaneously with production of white light rich in actinic rays (v. Berthelot, A. Ch. [3] 49, 486; Berzelius a. Marcet, S. 9, 284; Frankland, C. N. 6, 3; Sell, B. 5, 733; Delachanal a. Mermet, D. P. J. 214, 483). Water dissolves about 1000 of its weight of CS2 (Sestini, G. 1871, 473); it is miscible in all proportions with alcohol, other, ethercal and fatty oils, and liquid CO₂(v. Turaschmidta, Follenius, B. 4, 583). CS₂ is a solvert for fats, resins, gutta pereha, alkaloids, I, S, P, &c. (v. Liebermann, B. 12, 1294; Gore, P. M. [4] 30, 411). According to Sidot, CS2 is slowly changed in eunlight to CS and S (v. ante, CARBON MONOSUL-PHIDE).

Reactions.—1. Heated, strongly CS, gives C and S (Berthelot, Bl. [2] 11, 45%; Buff a. ilof-mann, A. 113, 129).—2. Burns in air or O to CO, and SO, -3. Decomposed by many metals; Fe is said to give CS at ord. temp. (Kern, C. N. 33, 253; v. also Merz a. Weith, Z. 11, 515); Cu at 200°-250° forms Cu.S, S, and C (M. a. W.); K gives a sulphide and C; Na at 110°-150° forms Na.S and Na.CS, the latter body reacts with diluto HClAq to produce 41,CS, (Löw, Z. 9, 173; 10, 120); Na amalgam according to Löw (l.c.) gives C.S., according to Raab (N. R. P. 19, 449) C.S. (v. also Hermanx, J. pr. 79, 448; 'Reichl, C. C. 1880, 420; Gaignett Bl. 1861. 111). -4. Hydrogen, when passed with CS, over heated Pt black, produces H2S and C; nascent H (Zn and 11ClAq) forms HzS and (?) C2111S2 (v. Vernon Harcourt, C. N. 26, 267; Cossa, B. 1, 117; Girard, C. R. 43, 396; Becquerel, C. R. 56, 237) .- 5. Chlorine reacts with CS, differently according to the conditions: dry Cl at ord. temp. gives 8,Cl, and CCl,; moist Cl, or MnO, and HClAq, or other Gl producer, forms IL Go, and CSCl, (Kolbe, A. 45, 41); Cl and CS. passed through a hot tube give S.Cl. and CCl. (K.); Cl passed into boiling CS, containing a little I forms S₂Cl₂, CCl₄, and CSCl₂ (Müller, C. J. 15, 41); the same products result by action of Repplied to liquids, e.g. mustard oil, by distilling ICl, (v. Weber, W. A. B. 1866, 348; Hannay, a little in a current of air into alcoholic potash; C. M 37, 224).—6. Chlorides which readily give coal-gas may be tested by passing through

up chlorine react eimilarly to Cl: MoCl, and SbCl, give CCl, and S₂Cl₂, SbCl, also producing SbCl, S which separatee into SbCl, and S (Aronheim, B. 9, 1788; Hofmann, A. 115, 264; Husemann, A. 114, 229); PCl, forms CSCl₂ (Carius, A. 113, 193), or according to Rathke (Z. 13, 57) Coll, and PSCl, (at 100°) .- 7. Bromine reacts in presence of I or SbBr3; 2 parts CS2, 14 parts Br, and 3 parts I, heated to 150° in a closed tube form CBr₁₄(Bolas a. Groves, B. 3, 508; v also Berthelot, A. Ch. [3] 53, 145). Hell a. Urech describe a compound C.S.Br₄ obtained by slow action of Br and CS₂ and sub sequent distillation (B. 15, 273).—8. Water, in presence of air, oxidiscs CS, slowly and partially to CO, Aq and SO, Aq (Berzelius); heated to 150° in a closed tubo from 3 to 4 hours, CO2Aq and H.SAq are formed (Schlagdenhauffen, J. Ph. [3] 29, 401); ovaporated in moist air a hydrate (?2CS_1I_O) is said to be formed (v. Properties). 9. Warmed with sulphuric anhydride, COS, SO2, and S, are produced (Armstrong, B. 2, 712). 10. Mixed with carbon dioxide and passed through a hot tube, or over hot Pt black, COS is produced (Winkler) .- 11. Passed through a hot tube with sulphuric acid, CO, SO, 112S, and S are formed (W.) .- 12. Many metallic oxides when heated with CS, react to form sulphides, sometimes also carbonates (Schlagdenhauffen, J. Ph. [3] 29, 401; Müller, P. 127, 401; Fremy, C. R. 35, 27) .- 13. Oxidising agents, e.g. KNO, or K.Mu.O., generally produce CO. and H.SO., sometimes CO. and H.S.; As.O., As.O., (and salts of these) produce As.S. (v. Schlagdenhauffen, Le.; Cloez a. Guignet, C. R. 46, 1110). 14. Boric acid and borates react at red heat to form B₂S₃.—15. Sulphuretted hydrogen mixed with CS, and passed over hot Cu produces CH. 16. Alkalis in Aqueous solution form carbonates and thiocarbonates, (v. THIOCARBONATES, under Carbonie Acid, &c.); alcoholic potash forms K xanthara C.H.O.CS.SK. - 17. Ammonia reacts with CS. probably to form CS(NH_), and (NH₃)₂CS₃ (Laurent, A. Ch. [3] 22, 103; Zouteveen, C. C. 1870, 821); CS₂ and NH₄ passed through a hot she, or CS₂ heated in a closed tube with alcoholic NII, produce II,S and A. 73, 26; v. also Millon, J. Ph. [3] 38, 401; Hofmann, J. 1858, 332); NiI3Aq form (NII3)2CS3Aq and NII, CNSAq.

Combinations .- 1. With many metallic sulphides to form thiocarbonates (q. v.) .- 2. With various annuonia derivatives; c.g. NMe, CS, 3. With zinc ethyl and methyl to form ZnEt, CS, and ZnMe, CS, respectively. -4.

With riethylphosphine to form Pitz, CS.

Analysis. Sulphur, by heating in a glass tube with Na, CO, and a little KNC, and estimating sulphates produced, as BaSO. Carbon, by barning with Phero, as in organic analyses.

Detection and Estimation.—Small quantities of CS, may be detected by adding alcoholic solution of potash, whereby K xanthate (C.11,O.CS.SK) is formed; on now adding soxanthate lution of a copper salt a yellow pp. is produced (Vogel, A. 86, 369). This reaction is applied to liquids, e.g. mustard oil, by distilling

alcoholic potash. A very delicate reaction is to bring the CS, into contact with PEt, when a sharacteristic commine-red compound, PEt₄, CS_p, forms; it may be erystallised from ether. This reaction may also be applied for the estimation of CS₂ (Hofmann, B. 13, 1732). CS₂ is sometimes determined, e.g. in xanthates, ly standardised CuSO, Aq (Grete, B. 9, 921); this carbonates may be converted into the Pb salt (by addition of Pb agetate) and this naw be deconversed by of Pb acetate), and this pray be decomposed by boiling with water, and the OS, led into weighed bulls containing alcoholic potash (Delachanal Mermet, B. 8, 1192).

Carbon, sulphochlorides of.

I. THIO CARBONYL CHLORIDE. CSCI, Mol. w.

11469. (703). V.D 57.5.

Formation.—1. By reaction between Cl and CS₂.—2. By leating CCl₄ with S.—3. By passing CCl, and H,S through a hot tube (Kolbe, 1, 45, 41; Carius, A. 113, 193; Müller, C. J. 15, 41; Gustavson, B. 3, 989). Preparation.—To dry CS, about 2 p.c. I is

added and dry Cl is passed in (a reversed condenser being attached) until the volume of liquid has increased by about 3. The whole is boiled with water (to remove S.Cl.); separated S is removed, water is separated, the liquid is distilled from the water-bath whereby CCI, and CS, distil over; the residue is then distilled till a thermometer in the liquid reaches 175°; the distillate is fractionated, digested with water to remove S.Cl., dried, and fractionated. About 320 g. CSCl, (v. infra) are thus obtained from 1 kilo. CS₂; very finely divided silver (by reducing AgCl) is then added little by little (the liquid being kept cold) until the whole of the liquid is sonked into the silver; it is then distilled, the distillate is shaken with water to remove traces of S₂Cl₂, dried, and fractionated (Bathke, B. 3, 858). The reactions are (1) 2CS₁+5Cl₂=2CSCl₄+S₂Cl₂; (2) CSCl₄+2Ag-2AgCl+CSCl₂.

Properties and Reactions .- Golden red, limpid, liquid; penetrating odour, resembling that of COCl.; fumes in air. Exposed to light and then to a low temperature, large colourless crystals of a polymeride, nCSCL, separate; this body is unchanged in air r melts at 112.5°; is volatilised in steam; at 180° in a closed tube

it gives CSCl2.

B. 3, 858). Prepared by action of Cl on CS. (v. Thrown chorner, Preparation). A clear, golden-yellow liquid; vapour causes free flow of tears; B.P. 146°-147°; S.G. 1'712 at 12.8°. Decomposed, slowly by moist air, quickly by heating with water to 160°, to CO₂, HClAq, and S. Hented to 200° is decomposed to CCl, S, Cl; and

a little USCI... M. M. P. M. GARBONATES v. CARBONIC ACID, CARBONATES, and THIO-CARBONATES.

CARBON TETRA-BROMIDE v. TETRA-BROMO-

DI-CARBON-TETRA-CARBOXYLIC ACID v. ETHYLENE-TRIBA-CARNOXYLIQ ACID.

CARBONIC ACID, CARBONATES, AND THIO-CARBONATES.

CARBONIC ACID. A solution of CO, in water probably contains earbonic acid, H.CO. The chief reasons for this statement are as follows. . The mass of CO₂ dissolved by water at ordinary temperatures and small pressures (less than dissolve in water saturated with Co.; at a

760 mm.) varies as the pressure. But at pressures of 2, 3, or more atmospheres, the mass of CO2 dissolved is less than that calculated by the law of Henry a. Dalton. Khanikoff a. Louguinine (A. Ch. [4] 11, 412) give these numbers :-P = pressure in mm., α - vol. of CO, (measured at 0° and 760 mm.) dissolved by 1 vol. water at about 15°.

Ρ.	a.	- 1	P.	a_
694.71	0.9441	1	2488-65	3:1764
• 809·03	1.1619	- 1	2369.02	3.4857
1289.41	1.8617		255 ± 0	37152
1169.95	2.1623	- 1	2738:33	4.0031
$2002 \cdot 06$	2.9076	۰ اږ	3109.51	4.5006

When the pressure is decreased to 760 mm, over water saturated with CO, at pressures greater than 700 mm, most of the CO, escapes, at lirst rapidly, then slowly; the last traces of CO, can be removed by placing the water in racuo, or by long-continued boiling. Magnesium reacts with a solution in water of CO, to form MgCO_a and II; the quantity of H evolved is almost exactly that calculated on the hypothesis that the reaction is $\text{H.CO}_3\text{Aq} + \text{Mg} = \text{MgCO}_4 + \text{H}_4$ (Ballo, B. 15, 3003). Water holding in solution Na₂CO₃ or K2CO3 dissolves considerably more CO2 than pure water; NaIICOa (or KIICOa) is produced (Ballo, t.c.). Mg reacts with aqueous solutions of NaHCO3 or KHCO3 to form MgCO3, K2 (or Na.)CO₃, and H (Ballo, Lc.). Aqueous solution of CO₂ turns blue litmus wine-red, the blue colour returns on exposure to air; blue litmus in contact with CO_2 at pressures of 1, to 2 atmospheres becomes ermilion-red (Malagnti, A. Ch. [3] 37, 200. When CO_2 dissolves in water, heat is produced; [CO',Aq] = 5.880 (Th. 1, 260). This solution reacts thermally with alkalis as a dibasic acid; thus (Thomsen) -

The thermal value of the second formula-weight of NaOII (9,068) is considerably less than that of the first (11,016); in this respect carbonio acid behaves like sulphurous, sclemous, borie, acid, &c. (r. Acids, Basicity of). When moist ives CSCl₂.

CO. reacts with KOH, K₂CO₂ is obtained; II. Thionyl Peachloride, CSCl₄ (Rathke, from this a great many carbonates may be formed, the composition of which is that of metallic derivatives of a dibasic acid H.CO. The aq cons solutions of carbonates are decompose by almost all acids, not by HCNAq or H.B.O.Aq; hence the aginity of carbonic acid " is small (v. Affinity). But soluble silicates are at once decomposed by CO, Aq, and even insoluble silicates e.g. of Ca, Al, &c. are slowly decomposed by moist CO. The sulphur analogue of carbonic acid - ILCS3 - has been isolated. Finally farious derivatives, both of CO(OH), and C (SH), are known; viz. COCl., f2COBr.), CO(NH.)... COS, CSCl., CS.NHL.SH, CS(NIL).

CARBONATES. Norval carbonates have the composition M_cCO₃ or MCO₃. Certain acid carbonates, MHCO₃, have also been isolated; very few of these are known as definite solids, the principal are when M = Na, K, NH, (? Tl). Some insoluble normal carbonates, c.g. Ba, Mg, Fe, pressure of 4 or 5 atmos, acid carbonates are probably formed. Many 'basic' carbonates exist; these are most simply repeated by the general formula xMO(or M₂O₃).yCO₂.sH₂O. A few double carbonates are also known, usually compounds of alkali carbonates with others, e.g. (NH₃)₂CO₃.MgCO₃; K₂CO₂.NiCO₂.4H₂O. Several ethereal carbonates are known, derived from the hypothetical ortho-carbonic acid C(OH)₄; e.g. C(OEt), C(OP₃), (v. CARBONIC ETHERS.

e.g. C(OFt), C(OPt), (v. CARBONIC ETHERS).

Formation.—1. By reaction between metallic oxides or hydroxides and CO₂ in presence of water; the oxides which act as weak bases, Fe₂O₃, Al₂O₃, &c., do not combine with CO₂ when dry.—2. By ppn. from solutions of salts by alkali carbonate solutions; only the salts of BaO, SrO, CaO, Ag₂O, and HgO, yield normal carbonates; other salts give basic carbonates containing less CO₂, relatively to MO, the weaker is the oxide MO, and the warmer and more dilute is the solution; salts of such weak bases as Fe₂O₃, Al₂O₃, and SnO, give pps. of hydrates free from CO₂,—3. By strongly heating the alkali or alkalino earth salts of organic acids.—4. In some cases by reaction between CO₂Aq and a metal; c.g. Mg, Fe, Zn.

Solubility in water.—Carbonates of Na. K, Rb, and Cs, are easily soluble in water; carbonates of Li and Tl are much less soluble; other carbonates are nearly, or quite, insoluble. All carbonates are soluble, to some extent, in water in which CO_2 has been dissolved. All, except those of NII_4 , Rb, and Cs, are insoluble

in alcohol.

Reactions .- 1. All carbonates, except those of the fixed alkalis, are wholly or partially decomposed by heat alone; BaCO3 begins to decompose only at a full white heat, SrCO, at beginning white heat, and CaCO₃ at full red heat; Ag₂CO₃ gives up CO₂ at 200°, and at 250° the Ag2O gives off O and leaves Ag; MnCO3 heatod to 200°-300° in air gives MnO, and CO,.-2. Heated in steam all carbonates are decomposed to hydrates and ${\rm CO}_2$ (v. Rose, P. 85, 99 a. 279) .- 3. Carbonates are decomposed by aqueous solutions of most acids at ordinary temperatures with evolution of CO., 21CNAq and H2B2O4Aq, however, do not decompose earbonates.-4. Solid earbonates are decomposed by heating with solid boric acid, silica, potassium dichromate, and some other salts. Sulphuretted hydrogen reacts with many insoluble earbonates suspended in water, e.g. of Sr, Ba, Ca, Zn, Mg, Li, to form sulphides and CO2; the change proceeds the further the more water is present (Naudin a. Montholon, C. R. 83, 58). -5. The more stable earbonates when heated with carbon give CO .- 6. Alkali carbonates heated in phosphorus vapour give phosphates and C .-7. Aqueous solutions of acid carbonates do not usually affect the colour of tilinus; they give an alkalino reaction with rosolic acid.

Detection and Estimation.—Usually detected by decomposing by any cid and examining action of gas evolved on Co Mq. Usually estimated by decomposing weighed quantity by acid and

determining CO, by loss.

Aluminium carbonate. Existence doubtful. Pp. produced by alkaline carbonates with Alsalts variously formulated as a highly hydrated basis carbonate (Muspratt, C. J. 2, 206; Lang-

lois, A. Ch. [3] 48, 502; Wallace, C. Gazette, 1858, 410); as a compound of Al₂O₂H₅ and a small quantity of the carbonate employed, e.g., Al₂(HO)₈+2NH₂HCO₃ (Rose, P. 41, 462); and as pule₂Al₂O₂H₃ (Barratt, C. J. 18, 190). By Wibain and Renoul the pp. in the cold is said to be 2Al₂O₃, (D₂,8H₂O and to decompose about 30° (C. R. 88, 4133) (v. also H. Rose, P. 91, 460; Parkmann, Am. S. [2] 34, 324).

Ammenium carbonates. Three definite salts

Ammenium carbonates. Three definite salts seem to exist, the normal, the acid, and the seaqui, carbonate (Divers, C. J. [2] 8, 171). The last is sometimes regarded as a compound of the first and second (Deville, C. R. 34, 880). Divers considers the compositions of the three salts to he.

Normal carbonate, 2CO.,4NH,,4H,O. Sesquicarbonate, 3CO.,4NH,4H,O. Acid carbonate, 4CO.,4NH,,4H,O.

Rose's hyperacid carbonate may be the fourth term of this series.

I. Normatearbanate (NH₄)₂CO₃.H₄O. Prepared (1) by adding excess of NH₃Aq to a warm cone. solution of the commercial carbonato; (2) by warming water with the ordinary carbonate, and allowing solution to cool and crystallise; after this has been done repeatedly the cold mother liquor from the last crop of crystals deposits normal carbonate. Large clongated plates, freely soluble in water, insol. alcohol, sparingly sol. NH₃Aq. Decomposes in air to NH₃, CO₃, with evolution of NH₃, and at 85° to NH₃, CO₂, and H₂O. Is converted into carbamate by digestion in closed vessel at 20°-25°.

11. Acid carbonate NH, HCO₃. Occurs in guano-deposits (Ulcx, A. 61, 44). Prepared by saturating NH₃Aq, or solution of NH, sesquicarbonate, with CO₂, and drying over H₂SO, and KOH. S. (0 \cdot 11 \cdot 9); (10 \cdot 0) 15 \cdot 85; (20 \cdot 0) 21; (30 \cdot 2) 27 (Dibbits, J₄-pr. [2] 10, 417). This is the stable salt to which the other NH₄ carbonates are converted. Large, transpagent, trime-trie er, stals. Dimorphous, but never isomorphous with KHCO₃ (Deville). Solution at 36 evolves CO₂; even at ordinary temperature unstable in solution.

III. Sesquicarbonate.

(NH₄)₂CO₂.2NH₄HCO₂. By slowly heating the commercial salt till melted and then cooling; or crystallises from warm solution of the samo salt mixed with NII₄Aq. Large transparent crystals, losing H₂O and NH₃ in air and giving NH₄FCO₂. S. (13°) 25; (17°) 30; (32°) 37; (41°) 40; (49°) 50 (Berzelius). Solution easily decomposes with evolution of CO₂.

Another earbonate-(NII,),CO, - appears to exist in the mother-liquor from the preparation of the sesquicarbonate (Divers). The commercial carrionate is probably 3NII, 200, H,O. It generally contains 1 p.c. H.O in excess of this formula, and a little ammonia. Prepared (1) by dry distillution of animal matter and subsequent purification by redistillation with elfareoal; (2) by heating to redness NH,Cl and CuCO, in retort with receiver. The first products are H.O and NII, earbamate, subsequent distillation produces the commercial carbonate. A white, transparent, fibrous mass, with strong ammoniacal smell, volatile, but not without some decomposition. Solution strongly alkalina

Barium carbonate. BaCO₄. By adding $(NH_1)_2O_2Aq$ to BaCl₂Aq or Ba $(NO_3)_2Aq$; or Na₂CO₂Aq to BaSAq; or (impure) by strongly heating a mixture of BaSO₄, oharcoal, and K₂CO₄, and extracting the K₂Sy formed with water, leaving BaCO₂. S.=O; in water saturated with CO₂ at 4 to 6 atmos. pressure S=75. The satt remains in solution at Irdinary pressure, but is completely ppal, on boiling (Wagner, J. mr. 102, 233; J. 1862, 185). Found native as By adding | J. pr. 102, 233; J. 1862 135). Found native as Withertte. This mineral can be obtained artificially from amorphous BaCO, by erystallising from fused KCl and NaCl (Bourgeois, Rt. [2] 37, 447). A soft white poisonous powder, easily soluble in solution of NH₄Cl, NH₄NO₃, or ammonium succinate. Above red heat in presence of C it yields BaO and CO. Decomposed at red heat by accous vapour especially in presence of chalk. Yields BaSO₄ when shaken with K₂SO₄ or Na₂SO₄Aq, and is completely decomposed by boiling with NII, ClAq.

Barium acid carbonat, 2BaCO, CO, described by Boussingaul. (A. Ch. [2] 29, 280) but Rose thinks if cannot exist except in solution

Beryllium carbonats. A compound of variable composition is obtained by exposing Bc(HO), to air, ppg. beryllinm salts with alkaline earbonates, or boiling solution of double Be-NH, carbonate. The composition of pp. by last method is 5BeO.CO2.5H2O (Schaffgotsch). Decomposed by boiling water, soluble in alkaline carbonates. The salt BcCO, 411,O is obtained by passing CO, through water containing basic

salt in suspension and evaporating over H2SO4 in atmosphere of CO...

Berylliu'n ammonium carbonate. 2(BeCO₂, 'NH₃)₂CO₃).Be(HO) 2H₂O. By dissolving BeO in conc. Am CO, Aq at gentle heat, then boiling till solution becomes cloudy, filtering, and adding cone, alcohol; crystals are drained, washed with alcohol, and dried by pressing between paper (Humpidge, Pr. 39, 14). By similar method Deville (A. Ch. [3] 44, 5) obtained a salt of formula 3(BeCO₃(NII₃)₂CO₃).Be(IIO)₂

soluble in cold, and decomposed by hot, water.

Bismuth carbonate. Bi₂O₃.CO₂ (Berzelius) a white pp. obtained by dropping a solution of Bi3NO₃ into an alkaline carbonate, According to Lefort this pp. contains \(\frac{1}{2} \) H₂O evolved at 100°. Heated strongly yields Di₂O₂.3(BiO)₂CO₂.2BiO₂H₂.3H₂O. Occurs nativo as Bismuthite in South Carolina.

Cadmium carbonate. CdCO3(?) Occurs native with ZnCO₂. By ppg. solution of a cadmium salt with (NH₁)₂CO₂Aq. The white pp. is said to contain water, which is lost at 80°-ai20° and at a higher temp. to loso CO₂ and leave Cito (Lefort, F. F. [3] 12, 78); Rose (P. 85, 304) says pp. is nearly represented by CdCO₃. Moist Cd(HO)₂ absorbs CO₂ and at 300° loses all its material country (2004) Cd(C). water, leaving 2Cd().CdCO,

Cesium carbonate Cs.COs. S. alcohol (19°) = - 11.1. Ill-defined hydrated deliquescent crystals separate from a syrupy solution. On melting these leave Cs.CO. as sandy powder. Acid carbonate, Cs.H.CO., crystallises from aque-

ous solutions in large prisms.

Calcium carbonate. CaCO., S. (cold) = 0094; $(100^{\circ}) = 1.13$ (Fresenius, A. 59, 117); S. (0°) in water saturated with CO_2) = .07, (10°) = .088

(Lassaigne, J. Chem. Med. 1848. 812; Sohlossing, C. J. [2] 10,788).

Occurrence.-Native; in rhombohedra (hexagonal) as calcspar, S.G. 2.69-2.75; and in right rhombio prisms (trimetrio), S.G. 2.92-3.28, as arragonite; also abundantly as chalk, limestone, &c. Formed when hydrated or anhydrous CaO is exposed to moist air; but not by action of. CO2 on dry CaO.

Freparation. — 1. By ppg. CaclAq by NH₃), CO₂Aq. From not too dilute solutions below 30°, it is ppd. entirely as calcspar; above 30° the pp. contains arragonite, in gradually increasing quantity as the temp. rises, until about 90° the pp. is almost entirely arra-gonite. CaCO₃ which separates as calcspar from a cold, not too dilute, solution of the acid carbonate, is deposited partly in arragonitic form on addition of a very little PbCo, CaSO, or SrSO, (Croher, J. f. Mineral, 1871, 288). The arragonite tends to change to calespar form if left under cold solution. Can be obtained in form of calcite from ppd. carbonate by crystallising from fused NaCl and KCl (Bourgeois, Bl. [2] 37, 447).

Properties and Reactions.—Tasteless, white, slightly alkalino; easily soluble, when recontly ppd., and to a certain extent even when it hecomes crystalline, in aqueous solutions of (NH₄)₂CO₃, NH₄Cl, NH₄NO₃, and ammonium succinate. These salts, therefore, provent complete ppn. of calcium as carbonate. At full red heat is decomposed into CO, and CaO. Temp. of decomposition is lowered by passing air or steam over the CaCO. In closed tubo fuses to marble-like substance. Boiling ammonium chloride solution decomposes CaCO, forming CaCl₂ and (NII₂),CO₃. Sulphur decomposes CaCO₃ forming CaSO₄ without previous formation of H₂S (Podacei, G. 1874, 177). The sulphur is oxidised at expense or oxygen of water (languatelli & Pelloggio, G. 1874. 536), and the formation of sulphuric and is preceded by that of penta- and tetra-thionic acids (Bellucci, S. 1874. 179). A salt of the formula CaCl 2CaCO 611.0 was obtained by Pelouze (Bl. [2] 3, 183).

liydrates: above 30° CaCO, is ppd.; below 30° hydrates are formed containing amounts of water (10-27 p.c.) which vary with the temp. and time occupied in ppn. From a solution of lime in sugar-water cooled to 0°-2°, a hydrate CaCO₃,6H₂O, decomposed at 30°, is obtained; while the same solution at a little higher temp. yields CaCO3. 5H2O crystallising in rhombohedra. S.G. 1.783, decomposed above 15 even in water (Pelouze, Bl. [2] 5. 1-3). Same hydrate found by Scheerer (P, i.s., 382) and Rammelsberg (B, 4, 469). A gelutinous hydrate formed by action of CO, on CaO and water is described by Bondonneau (Bl. '2) 23 100).

Acid carbonate CaH2(CO3)2(?) is known only in solution; obtained by passing CO, into cold water containing suspended CaCO₂. Decomposed with separation of CaCO₃ on exposure

to air, or more quickly on boiling.

Basic carbonate 2CaC.Co. Obtained from CaCO, at a moderate red heat. Hardens by action of water, forming, CaCO, CaO, II, which is also produced by exposing CaO to moist air. This hydrate gives 2CaO.CO at low red heat, and at strong red heat gives and H.O (Fuchs, P. 27, 601).

Cerium carbonate (Ce₂(CO₂)₂)H₁O by exposing Ce₂O₃ to air or by ppn. -A white powder, partially converted on heating strongly in air into Ce.O.

Chromons carbonate CrCO₃. An amorphoue greyish-white substance, prepared by heating a chromous salt with Na CO Aq out of contact

with the air (Moissan, A. Ch. [5] 25, 401). Cobalt carbonate CoCO₃. By heating CoCl₄. with CaCO, to 150° in scaled tube for 18 hrs., or by decomposing CoCl, with a solution of NaHCO₂ supersaturated with CO₂ and heated to 140° in a strong vessel allowing slow escape of CO... Light, roso-coloured, sandy powder; microscopic rhomboliedra; not attacket by acids in the cold (Senarmont, A. Ch. [3] 30, 129).

Hydrated carbonates and double salts are formed on adding cobalt solutions to alkalino carbonates. Hot or cold, not too dilute, colutions yield a rose-coloured pp., which dried at 100° is 2CoCO₃.3Co(HO)₂ (Setterberg, P. 19, 55; Winkelblech, A. 13, 148; Rose, *ibid*. 80, 237). Boiling water partially converts it into Co.O. (Field, C. J. 11, 50); digested with NaHCO, or (NH₄)HCO₃Aq yields 3CoCO₃,H₂O (Deville, A. Ch. [3] 33, 75). Hot very dilute solutions yield blue CoCO, 2Co(110), aq. Heated above 150° both pps. give off H2O and CO2, yielding Co2O3 (Rose).

Cobalt-potassium carbonate

(CoKH)(CO₃)₂.411₂O. Rose-coloured crystals, prepared by action of excess of KHCO₃Aq on Co(NO₃)₂ or CoSO₄Aq. Decomposed by water (Rose; Deville, Deville, also obtained (Rose; Deville) $CoK_2(CO_3)_2.411_2O$.

Cobalt-sodium carbonate CoNa (CO.) .. 4H.O. Prismatic, and CoNa₂(CO₃)₂.10H₂O dark red, crystals; obtained together by the action of Co2NO3Aq on solution of Na sesquicarbonate

Copper carbonate. Unknown except in solution Obtained by dissolving the ppd. basic carbonate in CO.Aq at 4-6 atmospheres pressure (Wagner, J. pr. 102, 233).

Hydrated basic carbonates. CuCO, CuO, II, occurs native as malachite. Prepared by ppg. a cupric salt with an alkaline carbonate. The pp. at first is greenish blue, and is said to contain 1 mol. H.O; left in contact with liquid and washed it has above composition and is dark green. Heated to 200° yields malachite. Boiled with water it yields CO₂ and CuO (II. Rose; Field, C. V. 14, 71). Digested with (NII₁)₂CO₃Aq at 48.8° it is converted into CuCO, 5CuO, a dense black powder, which ie also obtained by prolonged boiling CuSO, Aq with Am, H., (CO,),, filtering from CuCO, Cu(HO), and adding more CuSO, (Field). Digested with Na.CO, yields CuCO, 7CuO.51L.O.

The basic salt 2CuCO, Cu(HO), occurs

Boilod with water yields CuO and evolves Co₂. It dry it is fairly permanent; if moist With hot NaIICO₂ yield p blue solution which after protracted boiling, deposits CuCO₂. CuO₂H₂ the change is less rapid.

Ferrous-hydrogen carbonate. Solution of (Field). Prepared artificially by secret process CO₂ dissolves FeCO₃ and Fo; the latter with CO₂ dissolves FeCO₃ and Fo; the latter with native as asurite in blue monoclinic erystals. (Phillips, A. Ch. [2] 7, 44). Asurite can be formed from malachite at ordinary temps. by addition of CO, and abstraction of ILO in pre-

temp. (Weibel, J. für Mineral. 1873. 245). Crystalliecd Cu(NO₂), heated with CuCO₂ under a preseure of 54 acrospheres yielde crystalline

warty mass of a surite (Debray).

Double salts: Potassic-cupric carbonate
5CuO.K.O.CO. 10H.O. A dark-blue eilky mass,
obtained by fading Cu(NO.) to KHCO.

Sodic-cupcic carbonate CuNa, (CO₃), 6H.O. By action of NaIICO₃ on CuCO₃.CuO₂H₂ at

40°-50°. Rhombic prisms.

Cuprammonium carbonate (NH.) Cu.Co.. Oblained as dark-blue crystals by dissolving basic carbonate of copper in Am₂CO₃ and pouring solution into alcohol. Water decomposes it into Am₂CO₃ and GuCO₃CuO.Cu(11O)₂ (Favre, Traite de Chem. par Pelouzo et Frémy). Also formed when CuO or Cu is digested in NII, Aq with accese of air.

Didyminm carbonate. Di₂(CO₃)3. crystalline powder obtained by passing CO, into water containing suspended DiOaHa. A hydrated earbonate ie obtained, as a very slightly rose coloured pp., by adding an alkaline carbonate to solution of a Di salt. Losee 3 H₂O bonate to solution of a Arrignac). and a little CO, at 100° (Marignac).

tinous pp., soluble in (NII,)2CO3Aq, and ppd. on boiling; insoluble in solution of fixed alkali

carbonates (Winkler, J. pr. 94, 1).

Iron carbonates. Ferric carbonate does not appear to exist (Gmelin; Soubeiran, A. Ch. [2] 41, 326). Double salts of ferricum and alkaline carbonates appear to exist in solution, as the pp. of ferric hydrate obtained by cone. alkaline carbonate solutions gradually redissolves, whereas pure well-washed Fe2O, II, doce not dissolve in these solutions.

Ferrous carbonate, FeCO,, occurs abundantly as spathic orc. Prepared by methods similar to those described for CoCO₃ (q. v.). It is a greyish-white crystallino solid, composed of minute rhumbohedra; scarcely attacked by dilute a us, nearly unchanged in dry air. It is darker and less alterable the higher the temperature at which it has been formed, and the longer it has been heated (Senarmont, A. Ch. [3] 30, 129). Spathic ironstone dissolves under pressure in water saturated with CO2 (S. = .72), and is deposited as a black amorphous pp. on boiling (Wagner, J. 1867, 135).

Highrated ferrous carbonate, FeCo, H.O, occurs native. Amorphous, white, earthy, little altered in air, scarcely decomposed by acide at ordinary temps. (Moissan, C. R. 59, 238). The hydrate obtained by ppg. solution of a ferrous salt with an alkaline carbonate rapidly decomposes on exposure to air. It can be obtained fairly pure as a greenish tastcless powder, by ppg. ferric-free ferrous sulphate with normal or acid alkaline carbonates. The sats are dissolved in de-acrated water, the pp. wached by decantation out of contact with air, and dried in CO2. If dry it is fairly permanent; if moist

evolution of H.

La₂(CO₃)₃.8H₂O. Lanthanum carbonates. Found nativo as Lanthanite in four-sided sence of a dehydrating agent and CO2 at high platee or minute tablee of the trimetrio eyetem.

The carbonate obtained by ppn. forms a white gelatinous mass which changes to shining getatinous mass which changes to snining crystalline scales. 2La_(CO₄), 15H.O; ppd. by Na_CO₄Aq from La_3SO₄Aq, and dried at ordinary temperature. Micaccousticales with silky lustre (Hermann, J. pr. 82, 382).

Lead carbonate. PbCO₃. Occurs native in trimetric crystals as white lead or or cerusite. Has also been found on bronze objects from Pompeii (Luca, C. R. 84, 1457). Prepared by ppg. Pb(NO3)2Aq with excess of alkaline carbenates in the cold (Berzelius; Lefort, J. Ph. [3] 15, 26). According to Resc (A. 89, 235) these pps. are all hydrocarhonates, the ratio of hydrate to carbonate increasing with the temp, and dilution of the solutions. Bodies of the same composition are formed by direct action of CO on Pb(IIO), but differ in being amorphous and opaque, instead of consisting of minute transparent crystalline grains.

White lead is a hydrocarbonate present-

ing varieties of composition represented by (1) 2PbCO₃PbO₃II.; (2) 5PbCO₃3PbO₄II.; (3) 3PbCO₃3PbO₄II.; (Muder, A. 33, 212); (4) 5PbCO₃2PbO₄II. (Phillips, C. J. 4, 165). Prepared by (1) Ditch method. Thin lead sheets are placed over gallipots containing weak acetic acid (21 p.c.); the pots are embedded in fermenting tan at a temp. of 60°-65°. The metal disappears in a few weeks. Oxide of load is first formed, and dissolved by the acetic acid, forming a basic acetate, and this is decomposed by the CO, evolved from the tan. (2) PbO is mixed with water and about 1 p.c. of Pb acctate, and CO, is passed over it. Pb(NO₂), has been used instead of acctate. Nearly insol, i 1 sat. CO.Aq even under pressure (Wagner). The ppd. carbona'e has S = 05 in sat. CO.Aq.

4PbCO₃.Na₂CO₃ Lead - sodium carbona.

(Borzelius, P. 47, 199).

Lead-chlero-carbonate. A compound of the chloride and carbonate of lead is obtained as a pp. by the action of CO, under pressure on

PbCl₂ (Miller, C. J. [2] 8, 37).

Lithium carbonate. Li.CO₂. S. $(0^{\circ}) = 1.539$; $(50^{\circ}) = 1.181$; $(100^{\bullet}) = .728$ (Beynde, Bl. [2] 43, 123). Found in many mineral waters. pared by dissolving an excess of (NII,),CO, in conc. LiClAq, and washing resulting pp. with alcohel, or by strongly heating Li acetate. Not decomposed by heat. Melts at low red heat, and solidifies to a vitreous mass on cooling. The solution is alkaline and decomposes NH, salts, but is decomposed by Ba(110), and Ca(IIO)2. By slow evaporation the solution deposits salt in small prisms. Water-saturated with CO₂ dissolves it more readily than pure water.

magnesium carbonate. MgCO₂. Occurs native as magnesite, in rhomboh dral crystals isomorphous with calespar; Infusible; dissolves slowly in acids. Prepared (1) By suspending in water the washed pp. obtained by adding solution of an alkaline carbonate to solution of a Mg salt (which always contains MgO,H2), passing CO, through the liquid till pp. is dissolved, and evaporating by heat (Rose, P. 42, 366); (2) By heating MgSO, Aq with Na, CO, Aq to 160° in a sealed tube; (3) By inclosing a soluble Mg salt with an alkaline-hydrogen carbonate, super-

saurages, resulting the CO, can slowly escape (Senarmont, Eg. R. 28, 663); (4) By carefully heating MgCO, KHCO, 4aq to 200°, and extracting with water (Eugel, Bl. [2] 44, 355). A white crystalline powder, ismorphous with arragenite by method (1); obtained in rhembe-hedra by (3); isodimorphous with CaCO₃. When neist it is alkaline to litums. CaSO, Aq partially decomposes it, especially in presence of NaClAq (Pleischer, J. pr. [2] 6, 273). Soluble in cold solutions of alkaline borates, ppd. on heating, but redissolved on cooling (Wittstein, Ar. Ph. 3) 6, 40). Hydrates (a) MgCO₂,3H₂O₂. Hexagonal prisms obtained by spontaneous evaporation of a solution of MgCO, in excess of CO Aq. They cose water in dry air but retain their form. (β) MgCO₂5H₂O. Transparent oblique prismatic crystals obtained by exposure of above solution at a low temp. Converted into (a) by exposure to air λ c. Boiled, they yield 4MgCO3, MgH2O2, 4aq (Fritzsche; v. also Engel, C.R. 101, 814).

Hydrocarbonates. As in the case of lead, the composition of pp. formed by adding an alkaline carbonate to solution of a Mg salt depends on proportions, strength, and temp., of solutions. A variable mixture of hydrocarbonates is knewn as magnesia alba; (a) 4MgCO₂, MgH₂O₂, 4aq; a white granular powder (for preparation v. Rose, A. 80, 231). (β) 3MgCO₃ Mg(110)... 3aq, occurs native as hydromagnesite in small white monoclinic crystals (Dana). Prepared artificially (Bezzelius, Fritzsche) it is a white, slightly soluble powder forming alkaline solutions. Dried at 1003 irp air it yields a (Rose). (γ) 3MgO.2CO₂, 3aq(?) (Fritzsche, P. 37, 310).

Denied by Rose.

Magnesium hydrogen carbonate. Mg.Hg(CO3)2 (Soubeiran). Obtained by shaking up magnesia alba with CO.Aq. The sol, has a bitter inste and alkalica reaction; becomes turbid at 75° but clears on cooling. Heated to 50°, or evaporated in vacuo, it yields hydrate a (v. suppa) (Berzelius ; Fritzsche).

Magnesium-aumonium carbonate.

Mg.(NA₄) (CO₃)₂.41l₂O. Translucent rhombohedra, from a mixture of cold MgCl,Aq, or MgSO, Aq, and NII, sesquiearbonate solution.

Magnesium - potassium - carbonates: (1) Mg(KH).(CO₄)₂,4aq. In large crystals from cold aqueous mixture of MgCl₂ or Mg(NO₃)₂ with excess of KHCO₃. The crystals at 100° become ope ue, and less water. Decomposed by water, leaving a residue 6MgCO₂Mg1MO₂ 6mf (Berzelius); also obtained in oblique rlambic prisms (Deville, A. Ch. 33, 75); (2) MgK (CO₃), and by digesting magnesic aiba at 60°-70° for 12 to 15 hours with KHCO Aq (Deville, Lc.).

Magnesium sodium carbonate MgNa (CO3)2. Prepared in anhydrous microscopic crystals as

the corresponding K salt.

On solubity of MgCO, in CO.Aq under pressure v. Merkel (J. 1867, 136) and Wagner (J. pr. 102, 233). On all bility of MgCO, and CaCO, in solutions of calcium and magnesium salts, and the reactions of dolomite or gypsum and magnesite in presence of water centaining CO₂ v. Hunt (Am. S. [2] 42, 49). For reactions of basic MgCO₂ and gypsum with CO₂Aq v. Gossmann (Am. S. [2] 42, 217, 368).

Manganese carbonate. MnCO, S. water = 018; aqueous CO₂ = 028 (John). Occurs native as diallogite, in rhombohedral crystals, isomorphous with calcite. Prepared by heating MnCl₂Aq in sealed tubes to 160° with Na₂CO₃Aq, or to 140°-170° with CaCO, for 12 to 48 hours (Sonarmont, C. R. 28, 693). 2MnCO₄H₂O is obtained by drying in vacuo the pp. formed by alkaline carbonates with manganous salts (Gm. 4, 214; Prior, Fr. 4869, 428). Dried in hir pp. contains Mn₃O₄. Equivalent quantities of Na₂CO₃ and MnCl₂ yield 5MnCO₃.2MnH₂O₂ (Rose, A: 80, 235). MnCO₃ is a fine amorphous faintly rose-coloured powder. The hydrate is snow-white and tasteless. Ahlydrous or hydrated the salt is permanent at ordinary temps. Heated to redness in air it yields Mn₃O₄. Strongly heated in H it yields MnO₂. In chlorine it gives 4MnCO₂ + Cl₂ = MnCl₂ + Mn₃O₄ + 4CO₂ (Wöller). Orlorine water, or calcium hypochlorite solution, converts it first into Mn₃O₄, and then into MnO₂. Solutions of ammoniacal salts dissolve it when freshly ppd.

Mercury carbonates. Mereuric carbonate unknown. Neutral or acid carbonates of K or Na yield brown red pps. 2HgO.HgCO, (Setterberg, P. 19, 60). Mercuric ohloride yields an

oxychloride.

Mercurous carbonate. Hg2CO3. A black or yellow powder. Hg2NO, Aq is mixed with slight excess of Na(ork)HCO3Aq; the mixture is set aside for a few days and frequently stirred, and the pp. then washed quickly, and dried over H2SO4 in vaeuo (Setterberg, S.e.). Easily loses CO.; is converted into HgO by exposure to air; blackened by alkalis with separation of Hg

Nickel carbonate. NiCO₂. For preparation, v. Cobalt carbonate. It is a greenish-white powder in minute zhomboliedra scareely attacked by strong acids at ordinary temperatures.

Hydroearbonates: 4 (1) NiCO, 2NiH, O, 4H, O. O Occurs native as emerald nickel (Silliman, Am. S. [2] 6, 248; Shepard, ibid. 250). (2) 2NiCO, 3NiII, O, 4II, O is the pp. obtained from cold NiSO Aq and Na, CO, Aq, when dried at 100°. Boiled with water takes up water and loses CO, Heated above 100° in air gives off water and CO2, and is partly converted into Ni O, (Rose, .4. 80, 237). Not altered by digestion at 60°-70° with NaHCO3Aq (Deville). The pps. produced by alkaline carbonates in solutions of nickel salts vary with ter perature, strength, and proportions, of solutions employed.

Nickel-potassium earbonates:

(1) NiK₂(CO₃)₂·4H₂O; shining apple-green microscopic needles. (2) NiKH₂(CO₃)₂·4H₂O; lightgreen, apparently oblique rhombic prisms. Obtained similarly to corresponding cobalt salts (Deville).

Niekel-sodium carbonate,

NiNa₂(CO₃).10H₂O is obtained like the cobaft salt, in small prisms [Seville].

Palladian carbonate. A light yellow pp. is formed by adding solution of an alkaline carbonato to a solution of a Pd salt. No CO, is evolved at first, but on continuing ppn. effervescence ensues and pp. turns brown. A small quantity of CO2 retained when dry (Berzelius). . . .

Potassium carbonates. Two salts have been obtained.

I. Normal cardonate: K₂CO₃. S. 95·2 at 3°; 111 at 12°; 204 at 70°. The commercial salt is prepared by freating the ash of plants, especially of beetroot, with water, and evaporating. The esidue, containing 60-80 p.o. K₂CO₃. is sold as 'wide potash.' The impurities-KCl, K.SO., and a little E silicate—are partially removed by digesting for several days with its own weight of cold water, decanting, and evaporating quickly with constant stirring. Tho small crystals obtained are washed with pure K₂CO₃Åq, dried, and heated to redness in metal vessels; the product is 'pearl ash,' which usually contains from 2 to 3 p.c. impurities.

, Puro K. CO, is prepared (1) by heating pure K oxalate; (2) by digesting powdered cream of tartar with water containing a little HCl, washing, drying to render silica insoluble, crystallising from water to remove Na salts, heating in a closed silver dish digesting residue with hot water, filtering, Evaporating to dryness, dissolving in cold water, evaporating, and repeating treatment with cold water and evaporation two or three times (Sta\$; Chem. Proport. 340). (v. also Smith, C. N. 30, 234).

Properties and Reactions. - White solid, melting at red heat (83°) (Carnelley, C. J. 33, 280), volatilised without change at white heat: very deliquescent; [K'CO',Aq] = 6490 (2'h, 3, 198). Solution strongly alkaline; hot solution deposits rhombic octahedra K₂CO₃.2H₂O; cone. solution deposits monoclinic crystals 2K2CO3.3H2O, which at 100° give K2CO3.H2O (Städeler, A. 133, 371; Pohl, W. A. B. 41, 630). Heated with aqueous vapour is partly deconiposed, giving KOH; heated with charcoal gives K and CO (v. Potassium); heated with sulphur forms K sulphide as d sulphate, and CO, (Berthelot, Bl. [2] 40, 362); heated in sulphur dioxide ves K.SO, and traces of K.S (Berthelot, A. C. [5] 30, 547). Solutions, about 1 in 10H.O, partly decomposed by CaO₂H₂ giving KOHAq; amount of change much increased by boiling; reverso reaction occurs with more cone. solutions.

II. Potassium-hydrogen earbonate: KHCO2. S. 22.4 at 0°; 32.2 at 20°; 45.2 at 40°; 16.4 at 60° (Dibbits, J. pr. [2] 10, 417). Frepared (1) by passing CO₂ into solution of commercial normal carbonate; (2) by passing CO₂ into solution obtained by lixiviating residue from licating K-II tartrate in closed vessels, and crystallising. Properties and Reactions.-Large transparent monoclinic crystals, KIICO, H2O solution has slightly alkaline reaction, and gives off CO₂ on gently warming (v. Dibbits, J. pr. [?] 10, 417). At 200° give K_2 CO₃ and

Rubidium carbonate Rb.CO₂. By ppn. of Rb.SO₂Aq with BaOAq, adding (NH₂)CO₃ to Rb.SO₂Aq with BaOAq, adding the constant of the consta water, and evaporating this solution, indistinct crystals of Rb₂CO₃, H₂O are obtained. Soluble in alcohol, strongly alkaline. Heated, lose water and leave Rb₂CO₃, as a sandy powder, which melts at a higher temperature. In air it deliquesces, and yiolds RbHCO, in glassy prismatie crystals; pormanent in air; having very faint alkaline reaction; easily converted by heat . nto Rb, CO, (Bunsen a. Kirchoff) (melting-point,

337°; Carnelley a. Williams, C. J. 37, 125).

Samarium carbonate Sm₂(CO₂)₂,3H₂O. Samarium carbonate Sm₂(CO₂), 3H₂O. Needles insofuble in water. The following double salts have also been preperted. Samarium-ammonium carbonate,

SmNH₄(CO₃)₂2H₄O. Samarium-pc ussium carbonate, SmK(CO₃)₂6H₂O; brillint needles. Samarium sodium carbonate.

SmNa(CO₃)₂.8H₂O; a crustalline pp. (Clève, Bl.

[2] 43, 168).

Silver carbonate Ag₂CO₃. Prepared in crystals by adding ammonia by drops to mixed solutions of AgNO₃ and Na₂CO₃ of definite strength (Vögel, *J. pr.* 87, 288). As a white p., becoming yellow on washing, by adding pp., becoming years of the state of exposure to light. Isocadily soluble in strong NH Aq. Solution treated with absolute alcohol yields a pp. containing Ag.CO, and ammonia (Berzelius). At 200° loses CO, and leaves Ag.O. By ppg. AgNO, Aq with large excess of alkaline earbonate and boiling, a substance, possibly a mixture is obtained which dried at 100° hus formula Ag₂CO₂.2Ag₂O (Rose, A. •81, 202). Ammoniosilver carbonate Ag.CO.4NH,. A grey pp. on adding absolute alcohol to Ag₂CO₃ dissolved in NII,Aq (Keen, C. N. 31, 231). Sodium carbonates. Three salts have been

isolated, besides various hydrates, and several

double salts.

I. Normal carbonate Na₂CO₃. Occurs in waters of several lakes and mineral waters; is a con-

stant constituent of ash of sea-plants.

Formation.—1. From Na SO, by heating with C and CaCO, and lixiviating with ILO (Leblanc's process) .- 2. From cryolite, by heating with CaO and decomposing the Na aluminate formed by CO2-3. By reaction between NaCl

and (NH4)2CO3 in solution.

•Preparation .- 1. Soda crystals are repeatedly washed with cold water until all sulphates, oblorides, de. are removed; the last larges of SiO₂ are removed by dissolving the washed salt in water, evaporating nearly to dryness, adding a little (NII,)2CO, heating till quite dry, dissolving in water filtering, evaporating, and heating (Wurtz, J. 1852. 357).—2. Soda erystals are repeatedly recrystallised, the crystals being obtained as small as possible (Gay-Lussac, A. 12, 15); most of the chlorides and sulphates and iron salts are thus removed. The washed salt is dried, heated in a silver tish, and the residue is repeatedly washed with small quantities of cold water; the salt is now free from iron, but may contain traces of siliea (v. supra) (Stas).

Properties and Reactions. — White solid; melts eat 1, 818° (Carnelley, C. J. 33, 280), giving off a tittle CO₂ (Jacquelain, A. Ch. [3] 32, 205; Mallard, A. Ch. [4] 28, 86; Scheerer, A. 116, 134). Heated in steam gives NaOH. Decomposed at high temperature by carbon, to Na and CO; by phosphorus, to C, CO, and Na orthophosphate (Dragendorff, C. C. 1861. 865); by silicon to C, CO, and Na silicate; by sulphur, at 275°, to Na S and Na S.O., at melting-point, to Na,S, and Na,SO,; by ferric oxide, or ferrois oxide, with evolution of CO, (Stromeyer, A. 107, 366); by ferrous sulphide to Na and Na.Fe sulphide (E. Kopp, Bl. [2] 5, 207). Solubility

111. Sesquicarbonate. Na,H₂(CO₂),3H₂O Occurs native; S.G. 2·112. Prepared (1) by heating NaHCO₂ to sulphide (E. Kopp, Bl. [2] 5, 207). Solubility

200° (Hermann, J. pr. 26, 312); (2) by grapo-

in water increases from 0° to 34°; from 34° to 79° S. is constant: S.=46.2 at 34° (Löwel, A. Ch. [3] 33, 353; Poggiale, A. Ch. [3] 8, 468; Tomlinson, C. N. 18, 2; Gerlach, Fr. 8, 279). Solution is accompanied with production of heat; [Na°CO°, Aq] = 5,610 (Th. 3, 198).

Hydrates of Na CO₃.—(1) Na CO₃.10H₂O; separates from moderately cone. solutions at ordinary temperatures, in clear, monoclinic crystals; these melt at c. 34, leaving Na CCAH O (Schindler, May. Pharm. 33, 14); according to Thomsen (B. 11, 2012) the residue is Na CO, 2H O but this gives up another HO in the air. Crystals of Na.CO. 101LO effloresco in' air; at 12.5 "they give the bydrate with 5H2O; and at 38° in vacuo, or over CaCl., the hydrate Na CO 1. (Watson, 1. M. 12, 130). Dissolve in water with disappearance of heat

[18a°CO-1.0HFO,Aq] = -16,160 (Th. 3, 198).

(2) Na_CO_1.1511_O (Jacquelain, A. Ch. [3]

32, 205). Crystallises from cone, solutions of

Na CO at -20°

(3) Na.CO_{2.}7H₂O (Löwel, A. Ch. [3] 33, 353; Rammelsberg; Marignac, Ann. M. [5] 12, 55). Crystallises from hot saturated solutions by cooling in closed vessels; if air has entrance the 10H.O hydrate ferms. Said to crystallise in two modifications, rhombohedra and rhombic tables, with different solubilities.

(4) Na_CO_2_6H_O; crystallises from Na_SAq standing in air, also from NaClAq mixed with

K₂CO₃Λq (Mitscherlich, P. 8, 441).

(5) Na CO, 51LO; crystallises at temperatures over 33° from molten Na₂CO₂.1011₂O (Berzelms, P. 32, 303); also by the efflorescence of Na CO 101LO at 13 50.

(6) Na_CO_211_O; melting Na_CO_1011_O at

34° (Thomsen); [Na "CO".2H"O,Aq] = 20. (7) Na "CO".H"O; from hot saturated solutions of Na CO3, or from hot solutions of Na_CO_3.10II_O; separates from boiling solutions; also projuced by efflorescence of some of the hydrates with more II.O (Mariguae, Ann. M. [5] 12, 55; Haidinger, A. 5, 369). $[N.t^{2}CO^{3}.H^{2}O,Aq] = 2,250 \ (Th.).$

11. Sodium-hydrogen carbonate; NaIICO, (Bicarbonate of soda). Formation. -1. By passing NE; into NaClAq, and then decomposing by

CO, under pressure;

 $NII_1 + NaClAq + CO_2 + II.O =$ NaHCO3 + NIL ClAq (Ammonia soda process).-2. By reaction between soda crystals in solution and commercial NII, carbonate. - 3. By reaction between CO, and afflore ced soda crystals, or a mixture of a part crystallised a 1 3 parts dry Na₂CO₂. -4. By passing CO₂ into Na₂CO₃Aq as long as it is absorbed (1 part Na₂CO₃ in 2 parts H.O) (L. Meyer, A. Supplbd. 2, 170; Berzelius, P. 16, 134; Mohr, A, 19, 15; 29, 268). Properties and Reactions.—White monoclinic tables; alkaline taste; clunges moist red litmus to blue, but has no action on colour of turmeric paper. In moist air readily goes to Na.CO, xaq. When heated gives off CO. and II.D; solution decomposed on boiling (Ross, 17. 34, 158). S. 88 at 10°; 14·64 at 70° (Loggiale, A. Ch. [3] 8, 468; also Dibbits, J. pr. [2] 10, 417).

Na, H2(CO,), 3H20 Occurs native; S.G. 2-112. Prepared (1) by heating NaHCO, to

rating solutions of NaHCO, in vacuo over H2SO1; (3) by melting together the two carbonates, in the ratio Na₂CO₃.10H₂O:2NaHCO₃, and standing in air till mass becomes crystalline, when it contains crystals of the sesquicarbonato; (4) by pouring alcohol on to a mixture of Na₂CO₃Aq and NaHCO₃, the salt separates in fine needles (Winkler, R. P. 48, 215). Monoclinic crystals; non-efflorescent in air, goes to Na₂CO₃ et red heat; aqueous solution in vacuo over H.SO, gives Na CO, and NaHCO, (Rose, P. 34, 160). S. 12-63 at 0°; 41-59 at 100° (Poggiale). IV. Double Salts. Sodium-potassium

carbonates.

(1) NaKCO, 6H, O; nionoclinie erystals, unchanged in dry air, effloresces in moist air by evaporating solution of equal equivalents of the constituent salts, and crystallising from K.Co,Aq. A. 100° loses 6H.O. S. 185 at 15° (Marignae, C. R. 45, 650; Marguerite, A. 56, 220; Stolba, Bl. [2] 4, 192, 7, 241).

(2) 2Na₂CO₃.K₂CO₃.18H₂O; from motherliquor from which K, Fe(CN), has crystallised out.
May be crystallised from K, CO, Aq (Marguerito).

SODIUM-CALCIUM CARBONATE:

Na_CO_aCaCO_a511_O; occurs native as Gay-Lussite; obtained, in microscopic monoclinic crystals, by reaction between freshly ppd. CaCO2 and cone. Na.CO.Aq at ordinary temperature. When dry this compound is decomposed by water (Fritzsche, J. pr. 93, 339; Boussingault, P. 7, 97; H. Rose, P. 93, 606).

Strontium carbonate. SrCO₃, S.G. (pp.) = 3 62. S. (cold or het) = 0003 (Bineau, G. R. 41, 509). S. (cold) = .005 (Fresenius) solubility diminished by NH3Aq or (NH3)2CO3Aq. S. (10° in satd. CO2Aq) = 12 (Lassaigne). Occurs native as strontianite. Crystals of trimetrie system, isomorphous with arragonite and witherite. Prepared by ppn. with an alkaline carbonate as a smooth white substance; in form of strontianite, by erystallising amorphous carbonete from fused KCl and NaCl (Bontgeois, Bl. [2] 37, 447). Heated in clled vessel CO, given off only at about white hear, but in aqueous vapour SrH O2 is formed at a much lower temp. Alkaline sulphates in solution do not decompose it at any temperature (Rose, P. 95, 284). Ammonium chloride solution boiled with it converts it into SrCl.

Thallium carbonate Tl₂(O_3 , S.G. (fused) 7.06 (Lamy). S. (15·5°) = 4·02; (60°) = 11·7; (100°) = 27·21. M.P. c. 272° (Carnelley, C. J. 33, 275). Formed by exposure of Tl in a saturated solution of TLO to air . Prepared by allowing granulated metal to oxidise in warmair, boiling with water containing excess (NH₄)2CO₃ and filtering. TLCO3 is deposited in groups of prisms (Miller, Pr. 14, 555), which are brilliant, highly refractive, very heavy, arhydrous, colourless; melting, undecomposed, much below redness to clear liquid which solidifies to dark-grey mass, and at red heat decomposes evolving CO... Taste mildly caustic and metallie. Solution has alkaline reaction not completely removed by super saturation with CO₂ (Cookes; Werther, J. 1864.

Thorium carbonate. Th(CO3)2.3ThH4O4.2H2O. Alkaline earbonates throw down a basic sult with evolution of CO2. Moist ThII, O4 absorbs CO2 from air. Tho, is not sol. in water containing CO. (Berzelius). Salt of above formula obtained |

as an amorphous pp. by treating hydrate suspended in water with ${\rm CO}_2$, or by ppg. solution of ThCl, with an alkaline carbonate.

Tin oarbonates. SnCO₃.SnO; by adding solid SnCl₂ to cone. Na₂CO₃Aq in absence of air: very unstable. If (NH₄)₂CO₃Aq is used, hexagonal prist's of (NH₄)₂O₂SnO₃CO₂3H₂O are said to be it med (Deville, A. Ch. [3] 35, 448).

Uranium carbonates have not been isolated. Alkaline carbonates pp. uranous hydrate from UCl,, a basic sulphate from U(SO,), and double carbonates from uranic salts.

Urenyl-ammenium earbonate. $(UO_2)CO_3.2(NH_4)_2CO_3$. S. = 5 at 15°; increased by (NH₄)₂CO₃. Prepared, in small yellow transparent erystals, by digesting in (NH₁)₂CO₃Aq at .60°-80° tho pp. produced by NH₃Aq or (NH₁)₂CO₃Aq from uranic salts, filtering, and allowing to cool. Docomposed slowly at ordinary, more quickly at higher, temps. leaving UO₃; solution boiled evolves NH₃ and CO₂, and deposits yellow pp., containing uranium, of doubt-ful composition (Arfredson, Péligot; Ebelmen, A. Ch. [3] 5, 189; Delffs, P. 55, 229). Uranyl-potassium carbonate.

 $(UO_2)CO_3.2K_2CO_3$. S. = 7.4 at 15°; insol. in alco-Prepared, as a bright yellow crystalline crust, by dissolving in KHCO3Aq the pp. formed from uranic salts by K,CO,Aq, and evaporating. At 300° evolves CO2; at red heat leaves mixture of K uranate and carbonate. KOHAq pps. all the U as K uranate, even in presoneo of excess of K2CO3.

Uranyl-sodium carbonate. (UO₂)CO₂2Na_CO₃. Preparation and properties similar to K salt. Two Ca salts,

similar to K salt.

(1) (UO₂)CO₃.CaCO₄.10H₂O; (2) (UO₂)CO₅.CaCO₅.5H₂O; occur nativo (Sliith, A. 66, 253). Yttrium carbonate. Y₂(CO₃)₃. Na₂CO₃Aq pps, it from yttrium salts with 12H₂O in the cold, and H2O at 100°. Not easily decomposed by he sparingly soluble in water containing CO. Solution in (NH,) CO3Aq, if concentrated, deposits a white crystalline double salt which does not redissolve iu (NII,)2CO3Aq; also soluble in K.CO. and Na.CO.Aq (Betzelius).
Zino carbonate. ZnCO. Occurs as calam-

Not obtained by precipitation. Pp. formed by KHCO₃Aq in ZnSO₄Aq is 2ZnCO₃.3ZnO₄H₂ (Berzelius). ZnCO₃ unaltored at 200°; slowly evolves CO₂ at 300° (Rose). Hydrocarbonates. Nativo hydrocarbonates.

bonates are (1) zinc bloom ZnO.ZnCO3.3ZnH2O2 (Berzelius), or ZnCO, 2ZnH, O, 4aq (Smithson a. Borndorff, Gm. 6, 15). (2) Auricalcite or green calamine: 2ZnCO₃, ZnH₂O₂, in which Zn is partly replaced by Cu. (3) Buratity, a hydrocarbonate containing Cu and Ca.

The pps. formed by alkaline carbonates in solutions of zinc salts all appear to contain water, and to vary in composition with strength, temperature, and proportions, of solutions. For results obtained under varying conditions, v. Roso (P. 85, 107), Schindler a. Boussingault (Gin. 1, 15). They all evolve CO₂ and H₂O at 200°, yielding ZnÖ (Rose).

Ammonio-carbonate of zinc (NH,Zn)CO, Deposited in crystals from a solution of ppd. zine carbonato in conc. (NH4)2CO3Aq (Favre, Traitede Chimie, Pelouze et Fremy, 2nded. 8, 47).

z Zinc-potassium carbonate 6ZnCO, 6K,CO, 7H,O (?). Crystallises from a solution of ZnCl, mixed with K sesquicarbonato (Deville, A. Ch. [3] 32, 75).

Zinc and sodium carbonate 8ZnCO, 3Na, CO, (?). Small crystals, obtained as potassium salt (Deville).

Execst of alkaline Zircenium carbenate. carbonate solution produces a pp. in solutions of Zr salts, soluble in Na (orK,)CO, Aq. Composition seems to be variable (Hermann, Klaproth. Vauquolin).

*Emo-camonro acro. H₂CS₃, Mol. w. un-known. A dark yellow very strongly smelling oil; obtained by adding cold dilute HClAq to K.CS, or No.CS,; very easily decomposed, by heating, to CS, and H.S (Zcisc, S. 41, 105; Ber-

zelins, P. 6, 450).

THIO-CARDONATES. These salts have the composition M_2CS_3 , or MCS_4 , when $M_2 = Na_2 &c.$, and M = Ca &c. A few basic safts are also known. The composition of the salts of the alkali and alkaline earth metals has been determined; several other thio-carbonates seem to be produced in the reactions between solutions of •metallic• salts and K.CS.Aq or Na.CS.Aq, but the composition of very few of these thie-carbonates of the heavy metals has been determined. Thio-carbonates are formed by reactions between CS₂ and aqueous solutions of the monosulphides of the alkali and alkaline earth metals, M.S and MS. By using MOHAq and CS2, thio-earbonates and carbonates are formed simultaneously; with MO.H.Aq and CS. (M = Ca,Ba,Sr) basic thio-carbonates are formed, e.g. CaCS₃,2CaO,H₂6H₂O. NH₄Aq renets with CS₂ to form (NH₄)₂CS₃ and (NH₄)CNS (Gelis, J. 1861. 340). The thio-carbonates are yellow, red-yellow, brown, or black, solves; the hydrated salts are yellow. The salts of the alkali and alkaline earth metals are soluble in water; those of the heavy metals are more or less soluble in excess of M₂CS₂Aq (M = Na & The thio-carbonates are not very stable; those of the heavy metals easily decompose to metallic sulphide and CS₂; conc. solutions of the alkali salts change to H₂S and alkali carbonates when boiled, dilute solutions decompose by standing in air to earbonates and S. Heated alone, most of them give metallic sulphide and CS₂; K₂CS₃ gives K₂S₃ and C. The thio-earbonates have been chiefly investigated by Zeise (S. 41,105); Berzelius (P. 6, 450); Walker (C. N. 30, 28); Sestini (G. 1871, 473; B. 5, 327); Gelis (J. Ph. [3] 39, 95; C. R. 81, 282); P. Thénard (C. R. 79, 673); Husemann (A. 123, 67); Mermet (C. R. 81, 344).

Ammenium thiecarbenate (NII,) CS, . Prepared by rixing a saturated alcoholic solution of NH, with 1 its vol. CS, cooling after the liquid has become brown, pouring off liquid, and washing the crystals deveral times with alcohol, then with ether, and pressing between paper (Zeise). Yellow crystals, v. sol. in water, insol. in alcohol or ether; may be sublimed in dry air by gentle warming; dery hygroscopic. Aqueous solution heated to 90°-100° evolves H2S, and NH4CNSAq remains (Gélis).

Barium thiecarbenate BaCS,. By shaking BaSAq with CS2, washing with alcohol, and

drying in vacuo.

Calcium thiocarbonate. CaCS,. By digesting CaS with excess of CS, and evaporating in vacuo. Citron-yellow; sol. in alcohol or water; milk of lime shaken with CS, gives an orange-red pp. of CaCS, 2CaO, II, 6II, 0 and this at 30° gives red liquid from which red crystals of CaCS, 3CaO, H2.7II.O separate (Walker; Sestini).

Potassium thiecarl)nate. K, CS,. K.SAq is digested with CS2 at 80° in a closed vesse, or CS, is dissolved in a cone, alcoholic solution of K.S, yellow deliquescent crystals separate; dried at 60°-80° these give K.CS, a red-brown solid; v. sol. in water, vl. sol. in alcohol.

The other thiocarbonstos which have been fairly well examined and analysed are those of Lithium, Maghesium, Sedium, And Strentium. Thioearbonates of Bi, Cd, Cr, Co, An, Fe, Pb, Mn, Hg, Ni, Pt, Ag, Sn, Zn, seem also to be formed by adding the solution of an alkali thiocarbonate to a solution of a salt of each of these metals. M. M. P. M.

CARBONIC ANHYDRIDE CO, v. CARBON, OXIDES OF.

CARBON TETRA-CHLORIDE v. supra and Tetra-chiono-methane.

CARBONIC ETHERS. There are three classes of carbonic ethers: viz. acid ethers CO(OR)(OH), normal ethers CO(OR)2, and others of ortho-carbonie acid C(OR). In these formulæ R may be any alkyl. They are described as salts of the alkyl: e.g. ETHYL CAR-BONATE, METHYL CARRONATE, PHENYL CARBONATE, &c.

Orthecarhenic ethers are formed by the action of sodium alcoholates on chloropierin (Williamson a. Basset, A. 132, 54). They are converted by ammonia into guanidine.

Nermal carhonic ethers.

Formation. — 1. From alkyl iodides and silver early mate (de Clermont, A. 91, 375).— 2. By the action of Na, K, solid NaOEt (4 mol.) or KOEt (4 mol.), upon alkyl oxalates (1 mol.) (Ettling, A. 19, 17; Löwig a. Weidmann, A. 36, 301; Genther, Z. 1868, 656; Cranston a. Ditt-mar, Z. 1870, 4).—3. By the action of alkyl chloroformates upon sodium alcoholates, e.g.:

Cl.CO.OEt + NaOMe = NaCl + MeO.CO.OEt(Roese, A. 205, 240). The mixed other prepared from ethyl chloroformate and sodium methylate is identical with that from methyl chloroformate and sodium ethylate. -4. From COCl, and sodium alcohomites.

I ro erties. The LdYing points and specific gravities of the fatty carbonic others are as fellows (Hococ, A. 205, 214):-

Ether	Pailing-point	S.G.
Me,CO. Meliticos Et,Co. MePrcOs Pr.COs Me(PrCH.)COs Et(PrCH.)COs Et(PrCH.)cOs Et(C,H ₁₁)COs (C,H ₁₁)cOs	90·6° 109·2° 125° 139·8 169·2° 145·6° 160·1° 190·3° 182·3° 228·7°	1.065 at 17° 1.00 at 27° 97 98 at 27° 95 at 17° 95 at 27° 93 at 27° 92 at 15° 92 at 27° 91 at 15°

Reactions.—1. Ammonia converts the others R₂CO₂ into oarbamio ethers, and finally into urea.—2. PCl₃ forms chloroformic ethers. In mixed ethers RR'CO, the alkyl which is converted into obloride is the smaller of the two: $Et(C_5H_{11})CO_3 + PCl_3 = EtCl + ClCO_2C_5H_{11} + POCl_3$ The amides of the chloreformates, which may be regarded as half chloride half amide of carbonio acid (carbamic chlorides), are obtained by the action of COCl. upon the hydrochlorides of amines, c.g. COCl2 + NEtH2 = COCl(NEtH) + HCl (Gattermann a. Schmidt, B. 20, 118) cf. Chloro-FORMIC ACTD. - 3. When an alkyl carbonate is heated with an alconol containing a heavier alkyl, the heavy alkylelisplaces the light one (R.).

Chloro - imido - carbonic sthers ClN:C(OR),. These are formed by leading colorine into a cooled solution of NaOH and KCN in an alcohol (Sanduncyer, B. 19, 862). They are crystalline, and converted by dilute acids or by aqueous ILS into the corresponding carbonic ethers. Aqueous potassium arsenite reduces them to imido-carbonic ethers (cf. Chloro-imido-

CARRONIO ETHERS).

Imido-carbonio etbers HN:C(OR)... Prepared as above, are alkaline liquids, readily decomposed by aqueous acids into NH, and carbonic ethers (cf. IMIDO-CARBONIC ETHERS).

CARBO-DI-NICOTINIC ACID v. PYRIDINE TRI-CARBOXYLIC ACID

CARBONIC OXIDE. Namo usually given to

CO, v. Carbon, oxides of CARBONIC - OXIDE - POTASSIUM v. po-

tassium salt of Hextoxy-benzene. CARBONOUS OXIDE CO, v. CARBON, oxides of.

CARBON TETRA-IODIDE v. TETRA-10DO-METHANE.

CARBONPIMELIC ACID v. iso-Pentane TRI-CARBOXYLIG ACID.

CARBONYL. The divalent radicle C:O. When attached to two carbon atoms the product is a ketone, when attached to one carbon atom and to hydroxyl the compound is a carboxylic soid; when attached to one carbon atom and to one hydrogen atom the product is an aldehyde. Two or three carbonyls attached to CH render the hydrogen displaceable by metals. Many carbonyl derivatives of amido compounds are

described under the amido-compounds from which they are formed by the action of COCl... CARBONYL - DI - m - AMIDO - DI - BENZOIC ACID v. DI-PHENYL-UREA-DI-PU-CARBOXYLIC ACID.

CARBONYL - AMIDO - PHENOL v. Anhydride of Oxy-Phenyl-C - AMIO A ID. CARBON IL BROMIDE v. CARBON, OXY-BRO-

CARBONYL DI-BIURET v. BIURET.

CARBONYL-CARBAMIC ETHER C.H NO. i.e. CO:N.CO.OEt or (C,II,NO3)3. Carboxethyl cyanate or cyanurate. [119°]. Formed by the action of chloroformic other on potassium cyanate. If dry ether be present a second compound C₁₀H₁₂N₂O₃ [107°] is also formed. Rl'oubic plates, sl. sol. co[1, alcohol, v. sol. CHCi₂. When heated with water to 100° it loses CO. forming cyanurio ether. Its formula should therefore possibly be trebled. The compound C₁, H₁₅N₁O₅ when distilled with water behaves similarly (Whrtz a. Henniger, C. R. 100, 1419; A. Gir. [6] 7, 132).

Compounds with cyanic ether (a) C₁₀H₁₃N₃O₅ or (CO:NCO₂Et)(CO.NEt), [107°]. Formed as above, or together with oarboxy-carbamic (imido-diformic) ether [50°], (226°) when the ther is wet. Neodles, which lose CO2 on heating, yielding cyanurio ether

(W. s. H.).
(b) C₁₁H₂N₂O₇ i.e. (CO.N.CO₂Et)₂(CONEt).
[123°]. Foliand together with imido-diformic ether when KCNO acts on an aqueous othereal solution of chloroformic ether for a long time. On distilling it forms cyanuric other (W. a. H.).

CARBONYL CHLORIDE v. CARBON, OXYCHLO-RIDE OF.

CARBONYL-GUANIDINE v. AMIDO-DIOYANIO

CARBONYL DI-PHENYLENE v. DIPHENYL-ENE KETONE

CARBONYL-DI-PHENYL OXIDE v. DI-PHENYLENE RETONE OXIDE.

CARBONYL-PYRROLE C.II, N.O i.s.

 $OC < NC_1H_1$. Di-tetrol-urca. [63°]. (c. 238°). Formed, together with di-pyrryl-ketone, by the action of carbonyl chloride upon pyrrolpotassium (Ciamician a. Magnaghi, B. 18, 414; 1829). Large monoclinic crystals, =1.1688:1: 7189. V. sol. alcohol and ether, insol, water. By heating to 250° it is transformed into a mixture of pyrroyl-pyrrol C₁H₁N.CO.C₁H₃NH CO(C₁H₃NH)₂ and di-pyrryl - ketone

CARBONYL SULPHIDE v. CARBON, OXYSUL-

PHIDE OF.

CARBONYL-UREA v. UREA.

CARBOPETROCEME v. PRIROCENE.

CARBO - DIPHENYLENE v. DIPHENYLENE

CARBO-DIRHENYLIMIDE v. DI-PHENYL-CYANAMIDE

CARBO-TRI-PHENYL-TRIAMINE v. Dr. PHENYL-MIDO-BENZAMIDINE

CANDO-PHENYL-TOLYL-IMID V. PHENYL TOLYL GYANAMIDE.

CARBO-DI-PROPYL-DI-PHENYL-IMIDE v. DI-PROPYL-DI-PHENYL-CYANAMIDE.

CARBO - PYRIDENIC ACIDS v. PYRIDINE CARBOXYLIC ACIDS

CARBO-PYROTRITARIC ACID v. DI-METHYL-FORFURANE CARBOXYLIO ACID.

CARBO-PYRROLIC ACID v. Pyrrol-CAR-BOXYLIC ACID.

CARBO-PYRROLYL-FORMIC ACID PYRRYL-OLYOXYLIC AOID.

CARBOSTYRIL C.H,NO i.e. C.H.

CH:CII $C_{s}II_{s} < color = color$ Lactam or lactim of o-

(Py-3)- $\bar{O}xy$ -quinoline. amido-cinnamic acid. [199°].

Formation .- 1. By boiling o-amido-oinnamio acid with HClAq (Chiozza, C. R. 34, 598; A. 83, 117; Tiemann a. Oppermann, B. 18, 2070).—2. Obtained by reguoing tri-chloro-oxy-quinoline

with HI (Rotheit, J. pr. [2] 29, 800).

Preparation. — o-Nitro-oinnamic ather is heated with alcoholic (NH₃) to 100 ander pressure, the solution is evaporated to dryaces, taken up with NaOH and the carbostyll mod by CO₂ (Friedländer a. Ostermeyer, B.·14,

1916).

Properties.—Prisms (from alcohol); or long thin threads (Ontaining aq) from dilute aqueous solution. May be sublimed. V. s. sol. cold, v. sol. hot, water. Sol. alcohol and other. Alkaline KMnO, oxidises it to isati's and oxaloxyl-amido-benzoic acid (carbosty filic acid).

Salts.—The K and Na salts form easily soluble plates. The barium salt A'.Ba: spar-

ingly soluble plates.

Methylether: (247° mcor.). Colourless

liquid. Smells of oranges.

Ethya ether: [below 0°]. (256°). Prepared by the action of ethyl iodide on sodium-ear-bostyril, or of alcoholic KOII on chloro-quinoline. Volatile with steam. Colonrless liquid. Sweet smell.—B'HCl: hygroscopic crystals.

Phenylether: [69°]. Sublimable. Glistening plates. Sol. ordinary solvents (Fried-

länder a. Ostermayer, B. 15, 335).

Reference. - AMIDO-CARBOSTERIL, BROMO-CAR-Hydrocarbostyril is described BOSTYBIL, &c. under Amido-Phenyl-Probionic acts. Ethylpseudo-carbostyril is described as (Py. 3, 1)-Oxy-KTHYL-QUINOLINE.

CARBOSTYRIL-CARBOXYLIC ACID v. Oxy-

QUINOLINE-CARBOXYLIC ACID.

CARBOTHIALDINE C.H, N.S. Crystals which separate on adding CS, to an alcoholic solution of aldehyde-ammonia (Redtenbacher a. Liebig, A. 65, 43). Also from aldehyde and animonic thio-carbamate (Mulder, A. 168, 235). Insol. water. Sol. acids.

Reactions .- I. HCl splits it up into aldehyde, NH3 and CS2 .- 2. KMnO4 forms H2SO4, CO2, KCy, and acetic acid. - 3. HCl and Fe Cl, forms NH Cl, aldehyde, and NH_CS.S.Co.N.J. (Gnareschi,

G. 8, 246; B. II, 1383).

CARBO-TRI-THIO-HEXABROMIDE v. 11mxa-BROMO-DI-METRYL TRI-SULPHIDE.

CARBO-POLYLENE-DI-PHENYL-LATRA-MINE v. DI-PHENYL - TOLVLENE - TETEL - AMIDO-

CARBO-DI-TOLYL-IMIDE v. D1-TOLYL-CYAN-

S(CO,Et)2. DICARBOTHIONIC ETHER (180°). From ClCO, Et and alcoholic Na.S. Oil; decomposed by baryta-water or alcoholic KOH into Et₂S and CO₂ (V. Meyer, B. 2, 298).

CARBO-VALERTHIALDINE C. II. N.S.. [109°]. (G.); [117°] (S.). V.D. 60. From isovaleric aldohyde (5g.), CS. (3g.) and aqueons NII. (Schröder, B. 4, 469). From isoval-ric aldebra (Schröder, B. 4, 469). hyde and ammonium thio-carbamate (Mulder, A. 168, 237). Colourless needles (from Alcohol). Re.Cl. on witning gives the sulphocyanide re-action. KMn0 forms HCN, H₂SO, and valeric acid. Fe.Cl. and HCl form in the cold a yellow powder (S.CSML₂): Carbovale adding may thereforc be dithiocarbamate of di-valerylidene ammonium H₂N.CS.SN(CH.CH₂CHMe₂)₂ (Guareg-chi, A. 222, 310; G. 13, 500)

CARBOVINIC ACID is Hydrogen ETHEL

CARBONATE (q. v.).

CARBONATE (q. v.).

CARBONATE (q. v.). CABBOXANIDO - CARBINIDANIDO - BEN-TOIO ACID e. p. 157.

CARBOXAMIDO - CYANAMIDO - BENZOYL v. p. 155.

CARBOXAMIDO-HIPPURIC ACID v. p. 164. CARBOXAMIDO-O-OXY-BENZOIC ACID

C13H12N2O2. A product of the action of urea on amido-salicylic acid at 200° (Griess, J. pr. [2] I, 235). Minute needles, v. sl. sol. most solvents.

CARBOXETHYL CYANATE or CYANURATE v. CARRONYL-CARBAMIC ETHER.

CARBOXY-ACETO-GLUTARIC ACID v.

METHYL PROPYL KETONE TRI-CARBOXYLIC ACID. v-CARBOXY-o-AMIDO-BENZOIC ACID v. ISATOIC ACID.

v-Carboxy-m-amido-benzois other v. p. 157. CARBOXY BENZENE PHOSPHONIC ACID CO.H.C.H.PO(QH), [above 300°]. Prepared by the oxidation of p-tolucne phosphonic acid Calla (CH3). PO(OH)2 with KMnO3. Needles or tables. V. sol. water, m. sol, aqueous HCl or alcohol. On deating to 300° it decomposes, giving metaphosphoric and benzoio acids,

Salts. - A"Aga: slightly soluble pp. -A"H2Kaq: fine needles sol. water, sl. sol. alcohol. - A" HK: long prisms sl. sol. water.

Chlowide CaH, (COCI) (POCI,). [83°]. (315°). Colourless crystals. Mented with PCL it gives p-chlorobenzoyl chloride PCl₂ and POCl₃ Trimethyl other A"Me,; thick liquid (Mi-

chaelis a. Panek, B. 14, 405).

CARBOXY BENZOYL-ACETIC ACID v. ACE-TOPHENONE DI-CARBOXYLIC ACID, p. 37. CARBOXY - BENZOYL - AMIDO - BENZOIC

ACID v. Phthaloxyl-amido, benzoic acid. CARBOXY-BENZOYL-ETHENYL TRI-CAR-

BOXYLIC ACTO v. PHENYL-RETONE TETRA-CARBOXYDIC ACID.

CARBOXY BENZOYL PROPIONIC ACID C₁₁H₁₀O₃ i.e. CO₂H₁CO₂C₄H₄CO₂H. The free acid is unstable, but its Na salt is formed by dissolving phthalyl-propionic acid in NaOHAq $^{\prime\prime}$ (Clabriel a. Michael, \hat{B} : $\hat{\Pi}$, 1680).

o - CARBOXY - BENZYL - ACETO - ACETIC ETHER

C₁₁II₁₆O₅ i.e. CO₁II.C₆II₄ OII₄ OII₄ OII₆ (92°). Formed by reducing phthalyl-acetoneetic eteer with zinc-dust in glacial acetic acid (Bülow, A. 236, 190). Needles; v. sol. hot water, alcohol, ether, and HOAc. The ammomium salt melts at [121]. Boiling barytawater converts it into benzyl-acctono o carboxylic acid.

Though hydrazide C20H22N2O4. Deec nposes slewly forming alcohol and $C_{18} \Pi_{18} N_s O_3 [229]^2$.

o-CARBOXY-BENZYL-MALONIC ACID HO2C.CaH1CH2.CH(CO2H)2. Formed by saponify ing e-carboxy-benz 1-majornic ether (Wisliconus. A. 242, 37). Prisms. V. sol. hot, sl. sol. cold, water. Heated to 190 gives off CO and forms

o-earboxy-phenyl-propionic neid [166].
Salt.—A"A"A": V. sl. sol. hot water.
Di-eth M ether CO.H.C.H.CH.CH.(CO.Et).. [86"]. S. 045 at 17. Formed by reducing phthalyl malonic ether with acetic acid and zinc (Wirlicenus, A. 242, 32). Fino needles. V. e. sol, ether and het alcohol, sl. sol. hot water.

Salts.—A'Ag: white needles.—A'Na: deli-quescent needles. V. sol. alcohol, ansol. ether. Tri-sthyl-ether A"Et .. (250°) at 45 mm. CARBOXY-CARBAMIC ETHER C.H., NO.

i.e. NH(CO.Et)₂. [50°]. (226°).

Formation.—1. By the action of chloroformic ether on potassium cyanate in the presence of wet other. A second compound C10H15N3O,[107°], insoluble in water, is also formed, while a small quantity of yellow oil [170°] is found; in the aqueous extraot. If absolute alcohol be used instead of other, the Geoord compound is not formed.—2. Chloroformic ether (34 gr.) and curbamic ether (24 gr.) are heated together at 120°.

Properties.-Long prisms. It forms biuret and alcohol when made with aqueous NI1,. The salt C_aH₁₀NO Ag crystallises in cubes (Wurtz a. Henniger, A. Ch. [5] 7, 135).

CARBOXY-CINNAMIC ACID v. CINNAMIC 4cm.

 $\begin{array}{ll} \textbf{CARBOXY-CORNICULARIC} & \textbf{ACID} & \textbf{C}_{10}\textbf{H}_{14}\textbf{O}_{5} \\ \textbf{i.e.} & \textbf{C}_{6}\textbf{H}_{3}\textbf{.C}(\textbf{CO}_{2}\textbf{H})\textbf{:CH.CO}.\textbf{CH}(\textbf{CO}_{2}\textbf{H})\textbf{.C}_{6}\textbf{H}_{4} \end{array}.$ Formed as a by-product in the reduction of pulvic acid to dihydrocornicularic acid. It was not isolated, but was converted into the lactone by means of acetic anhydride.

Carboxy-cornicularic-lactone

OC---O CO.II $[215^{\circ}].$ C₁₆I1₁₂O₄ or C.H.C.CH.C: C.C.H.

Long felted needles or short prisms. In cold aqueous NII, or alkaline carbonates it dissolves forming salts of the formula CisHinOiM, but on heating it gives salts of earboxy-cornicularic acid C₁₈H₁₂O₅M₂ (Spiegel, B. 15, 1546).

CARBOXY-CYANAMIDO-BENZOYL v. p. 155. DI-CARBOXY-GLUTACONIC ACID C, II, Og.

Propylene tetracarboxylic acid.

Ethyl ether (CO_Et),..CII.CII:C(CO_Et),.. (270°-280°). S.G. 1131 at 15°. From its sodium salt by 11Cl. Oil, soluble in alcohol or ether. Boiled with HCl it gives off CO, and forms glutaconic acid (q. r.) and iso-aconitic ether. NaOH acts similarly. Sodium derivative (CO,Et)2.CNa.CH:C(CO,Et)2. [260°]. From malonio ether, NaOEt and chloroform (Conrada, Guthzeit, A. 222, 251). 2(CO_Et)CNa, + CllCl,= (CO2Et)2.CNa.CH:C(CO2Et)2 + BNaCl. yellow prisms. Insol. ether, sl. sol. celd water or cold aloohol, v. sol. hot water or hot alcohol. Gives a violet colour with ferric chloride, and pps. with metallic salts. Sodium amalgam reduces it to dicarboxy-gluturic acid, (CO₂H)₂CH.CH₂CH(CO₂H)₂ [167°]. When this acid is heated it gives off CO, becoming glutaric acid.

Mothyl , di carbe ., glutaconic ether. Sodium di-carboxy-glutaconio ether heated with alcoholic Mea at 150° forms methyl-di-carboxyglutaconic ether, (CO,Et), CMe.CII:C(CO,Et), an oil, which on suppnification gives riso to (CO₂H), CMc.CH:C(CO₂H), and this readily splits off CO₂, forming methyl-glutaconic acid, CO₂H.CHMe.CH:CII.CO₂H [‡37°]. Methyl glutaconic acid forms white crystals; v. sol. water, alcohol, or ether.

Bensyl-di-carboxy-glutaconic sther (CO_Et)_c(C,H_).CH. (CO_Et)_c, [78°], is formed in a similar way, using benzyl obloride. Insol. water, v. sol. hot alcohol, ether, or conc. H_SO_. When saponified by caustic soda it gives off CO₂ carbonic acid and forms benzyl-glutaconic acid. CO, H.CH(C.H.).CH:CH.CO,H, [145°].

. ...

DI-CARBOXY-GLUTARIC ACID v. DI-CAR-BOXY-GLUTACONIC AUID.

CARBOXYL. Dxatyl. The monovalent acid radicle CO2H i.e. CO.OH. Its hydrogen is always displaceable by metals, cf. Acros

CARBOXYLIC ACID, so-called, v. DI-OXY-BENZENE-DI-DUINONE.

Di-hydre-carboxylic acid v. Terra-oxy-

QUINONE. Tri-hydro-carboxylic soid v. Hexa-cxy-benz-

Oxy-carboxylic acid v. Benzene TRI-QUINONE. v-CARBOXY-OXAMIC ACID Di-cthyl ether. C,H,, YO, i.e. CO, Et. NH.CO.CO, Et. [45°]. From CICO.CO.Et and oxamic ether (Saloman, J.pr. [2] 9, 292). Needles (from ether); sol. water and alcohol.

· o-CARBOXY-PHENOXY-ACETIC ACID

C.H. (CO.H.) O.CH. CO.H. Carboxy-phenyl-gly-collic acid. [187°]. Formed by oxidation of o-aldehydo-phenoxy-acetic acid with KMnO. White needles. Sol. alcohol, ether, and hot water.—A"Ag_: white, sparingly soluble pp.

Di-ethyl ether A"Et.: liquid.

Di-amide C.H.O(CO.NH.). [158°]. Long yellow needles, sparingly soluble in ether, benz-

ene, and hot water, v. sol. chloroform and hot alcohol (Rossing, B. 17, 2995).

m. Carboxy-phonoxy-acetic acid C₆H₄(CO_.H).O.CH_..CO_.H [1:3], [206°], Prepared by oxidation of m-aldehydo-phenoxy-acetic acid with KMnO, Needles; v. sol. alcohol, other, and acetic acid, sl. sol. cold water.—A'Ag2: crys-

tallino (Elkan, B. 19, 3014). p.Carboxy-phenoxy-acetic acid C₈H₄(CO₂H).O.CH₂.CO₂H [1:4]. [278°]. Prepared by oxidation of p-aldehydo-phonoxy-acetic acid with KMnO. White needles; v. sol. alcohol, other, and acetic acid, more sparingly in benzene, chloroform, and ligroin, sl. sol. cold water.

Salts.—A"Ag: white sparingly soluble pp. The I'L Ru, and Fe salts are sparingly soluble ppsastife Cu and Ba salts are soluble (Elkan, B,

o-CARBOXY-PHENYL-ACETIC ACID C, H, (CO, II) CH, CO, II [1:2]. Homophthalic acid, Phenyl-acctic-carboxylic acid. Isuvitic acid. [174°]. Formed by saponification of benzylcyanide-o-carboxylic acid by boiling with dilute KOH. Colourless crystals. V. sol. alcohol and KOH. Colourless crystals. hot water, insol. benzene.

Salts. - A"Ag2: insoluble amorphous pp .--A"Ca 2aq: sparingly soluble crystalline powder. -A"Ba: casily soluble crystals (Wislicenus, B. 18, 173).

prisms; v. sol. other and chloroform. Formed

by the action of acetyl chloride on the acid.

Ethyl ether A"Et.: (292°); thick are matic cil (Gabriel, B. 20; 2499).

Imide C,H, CH2,CO CO.NH di-exy-isequincline);

[c. 233°]. Formed by dry distillation of the ammonium-salt. Short colourloss needles (from alcohol or acetic acid). Sublimos in long crystals. Sl. sol. alcohol. Dissolves in aquocus caustic alkalis. Heated with POCl, at 1500170° it is converted into di-chloro-iso-quinoline.

Methyl- mide C.H. | : [123°];

(314°-318°); long colourless needles; v. sol. ordinary solvents. Dissolves in aqueous alkalis. Obtained by dry distillation of the pachylamine salt of the acid. By McLand modhyl alcoholic KOH at 100° it is converted into a tri-methyl CMe₂CO

dorivative C₆H, CMe₂CO | [103°], which is

also obtained by methylation of the imide (Gabriel B. 19, 1654, 2364, 2363). The imide and methylimide combino with diazo-benzene in alkaline solution.

Amic acid C₆H₄(CO₂H).CH₂.CONH₂ (homophthalamio acid): [187^o]; colourless needles. Formed by slowly warming benzyl-cyanide-ocarboxylic acid C₆H₄(CO₂H).CH₂CN with conc. H₂SO₄ to 70°, and pourieg into water.

Amic methyl ether C₈H₄(CO₂Me).Cll₂.CONH₂: [112⁵]; erystalline solid (Gabriel, B. 20, 1202).

Benzen 3-azo-carboxy-phenyl-acetic-imide

C.H. CH-N₂-C_aH₅. [260°]. Formed by tho

action of diazo-benzeno chloride upon an alkalina solution of the imide of carboxy-phenylacetic acid. Orange-yellow needles (Gabriel, B. 20, 1205).

CARBOXY-PHENYL-BENZ-GLYCOCYAMI-DINE v. BENZGLYCOCYAMIDINE.

CARBOXY-PHENYL-GLYCOLLIC ACID v.
CARBOXY-PHENOXY-AGETIC ACID.

CARBOXY-PHENYL-MAIONAMIC ACID.

Ethylether CO.Et.Ch. Co.NH.C. II., CO.II. [173°]. A product of the action of malonic etter on mamido-benzoic acid (Schiff, A. 232, 144; B. 17, 403). Silvery needles When heated it freaks up into malonic et r and malonyl-di-amido-di-benzoic acid:

 $\begin{aligned} & \textbf{2CO}_{.}\textbf{Et}.\textbf{CH}_{.}\textbf{CO}.\textbf{NH}.\textbf{C}_{.}\textbf{H}_{.}\textbf{CO}.\textbf{H} = \\ & \textbf{CO}_{.}\textbf{Et}.\textbf{CH}_{.}\textbf{CO}_{.}\textbf{Et} + \textbf{CH}_{.}\textbf{(CONH}.\textbf{C}_{.}\textbf{H}_{.}\textbf{CO}_{.}\textbf{H})_{2}. \end{aligned}$

o-CARBOXY-PHENYL-METHYL-AGETIC
ACID C.H.(CO.II).CHMc.CO.II. a. Methyl-homoo-phthalic acid. Hydratropic-o-carboxylic acid.
[147°]. Formed by heating the inide with fuming
HCl at 200°. Colourless crystalline powder.—
A"Ag₂: crystalline pp.
CHVICO

Imide C,H CHMe.CO CO NH

Di-oxy-methyl-isoquinoline. [145°]. Formed by heating o-cyano-phenyi-methyl-accto-nitrile C₀H₄(CN), CHMc.CN with conc. II_SO, at c. 130° and pouring into water. Glistening prisms. Can be distilled undecomposed. Dissolves in aqueous alkalis. By digestion with alcoholic KOH and MeI it is converted into the methyl-imide of carboxy-phenyl-di-methyl-acctic acid

C.H. CO. NMe By POCI, it is converted

into (Py. 2:4:1)-di-chloro-methyl-isoquinoline, whilst (Py. 4:2:1)-chloro-oxy-methyl-isoquinoline is formed as a by-product (Gabriel, B..20, 2503). o.Carboxy-phenyl-di-methyl-acetic acid

C.H.(CO.H).CMe.CO.H. Di-methyl-homo-o-

phthalic acid. [c. 123°]. Obtained by dissolving the anhydride in boiling aqueous NaOH and ppg. with HCl. On heating it is reconverted into the anhydride. By distillation with soda lime it gives isopropyl-benzone.—A"Ag₂: crystalline pp.—A"K₂aq: plates.

Anhydride C.H. CMo.-CO [85°].

(312°) at 760 mm. Prepared by heating the imide or methylimide with fuming HCI (4 pts.) at 210° for 4 or 5 hrs. Flat crystals, slowly dissolved by aqueous alkalis, forming alts of the acid.

Imide Coll CO (di-mothyl-homo-

phthalimide): [120°]; (318°) at 770 mm. Prepared by the action of methyl iodide on a solution of the imide of carboxy-phenyl-aectic acid in methyl-alcoholic KOII at 100°. Flat needles.

Methyl-imide C₆H CO (tri-

methyl-homo-phthalimide). [103°]. (295°) at 770 mm. Formed by further methylation of the preceding imide or of the methylimide of o-CH₂CO

carboxy-phenyl-acetic acid C_sH_sCO.NMe Long needles. Slowly sublimes at 100°. V. sol.

ordinary solvents, insol. alkalis (Gabriel, B. 19, 2303; 20, 1198).
CARBOXY-PHENYL-METHYL-\n-m-PYRRYL-BENZOIC ACID C10H12NU, i.e.

HO,C.C:CM N.C. II, CO,H [210°]. Obtained

by saponification of its mono-ethyl-ether which is obtained by several days' standing of an acetio acid solution of acetophenone aceto-acetic ether (1 mol.) and m-amino-benzoic acid (1 mol.). Aggregates of colourless needles. Sol. ordinary solvents except water.

EtO C.C:CMe
| | N.C. II. CO II:

[160°]; slender yellow needles (from dilute acetic acid); sol. alcohol, ether, &c. Hs Ca, Ba, Sr, and Mg salts are white granular pps. (Pual a. Schneider, B. 19, 3162).

o-CARBOXY-PHENYL OXAMIC ACID C₀N-NO₅ i.e. C₆II₄(CO₂H^{*}.NH.CO.CO₂H. [210°]. S. 11 at 10°.

S. 11 at 10°.

Formation.—1. Y. ting — lie acid with co-amido-benzoic acid at 120°.

2. From carbo-styril and alkaline F MnO. (Friedlander a Ostermaier, B. 14, 1914; 15, 334).—3. From acetyl-quinoline tetrahydride and cold dilute KMnO. (Hofmann a. Köni., B. 16, 734).—4. From eynurin or cynurenic acid and alkaline KMnO (Kretsely, M. 4, 156; 5, 16).—5. Formed by oxidation of (P3.)-bromo-quinoline with KMnO. (Claus a. Collischonn, B. 19, 2767).

Properties.—Silv ry rectiles (containing aq), (from water), or geodes from ether). Decomposed by dilute acids, or by long boiling, into oxalic and amido-henzoic acids.

Salts.—(NH),A'': minute felted needles.— KHA'',aq.—BaA''aq.—BaH $_2A''$,aq.—CaA''2 $_2aq$.—CaA''2 $_2aq$.—CuA''CuO 4aq.—Ag $_2A''$.

Mono-ethyl ether C,H,(CO,H).NH.CO.CO,Et. Ethyl-oxalyl-anthranilic acid. [181°]. Felted needles. Formed by exidation of indexylic ether or indexanthic ether with CrO, (Baeyer, B. 15, 777).

m-Carboxy phanyl-oxamic soid CO2H.CO.NH.C.H4.CO2H. Oxaloxyl-amido-ben-

, zoic acid. Formation.-1. Bytboiling an aqueous solution of the barium salt of cyano-carbimicoamido-benzoic acid.-2. By heating (equal mols. of) m-amido-benzoio acid and anhydrous oxalio acid for an hour at 180° (Griess, B. 16, 336; 18, 2412).

Properties.—Small white plates, V. sol. hot water, m. sol. alcohol, insol. ether.—BaA" 2aq.

Mono-ethyl ether
CO_Et.CO.NH.U.H..CO_H(ethoxal benzamicacid)
[225°]. Formed by boiling mamido-benzoic acid with oxalic ether (Schill, A. 232, 132; B. 17, 402; G. 15, 534). Silky needle; (from water or alcohol). When heated above 225° it splits up into oxalic ether and carboxy-phenyl-oxamide.

Amide-ether CO.Et.CO.NH.C.II, CONH... Amide-ether [191.5°]. Got by heating m-amido benzamido with oxalic ether. Gives with aniline the annideanilido CO(NHPh)CO.NH.C₆H₄CO.NH₂ [c. 310°]. Anilide ether CO.Et.CO.NH.C.H..CONPhH [180°]. From m-amido benzamilide and oxalio ether. Satiny needles.

Amic acid CO(NH2)CO.NH.C.H4.CO2H v.

PHENYL-OXAMIDE CARROXYLIC ACID.

CARBOXY-PHENYL-OXAMIDE v. PHENYL-OXAMIDE CARBOXYLIC ACID.

CARBOXY-PHENYL-OXY-ACETIC ACID v. CARBOXY-PHENOXY-ACETIC ACID

o-CARBOXY - PHENYL PHENYL - CARBA -MATE C.H.(CO.H).O.CO.NHC.H.

Methyl ether Call, (CO2Mc).O.CO.NIIC, H3. [238°]; long needles; sublimable. Formed by heating methyl salicylate with phenylcyanate (Snapc, B. 18, 2431).

m-CARBOXY PHENYL PHOSPHORIC ACID C,H,(CO,H).O.PO(OH)F [201°]. From its chloride and water. Scales, v. sol. water, alcohol, and other. Water at 160° decomposes it into phosphoric and m-oxy-benzoic acid (Anschütz a. Moore, A. 239, 333).

Chloride C,H,Cl,PO, i.e.

C_aH₄(COCl).O.PO.Cl₂. (170°) at 12mm. S.G. ²⁰ 1.548. From m-oxy-benzoic acid (I mol.) and PCl, (11 mols.). Further treatment with PCl, (1 mol.) produces C.H.Cl.PQ. (178°) at 11 mm.

This is probably C.H.(6000°) t0.PCl, and is converted by water into C.H.(100-H).0.PO(011). further quantity of PCl, converts C,H,Cl,PO, into C.H.Cl.CCl.

o-CARBOXY-8-PHENYL-PROPIONIC ACID HO, C.C., H, CH, CH, CO, H. [166°]. Formed by heating o-carboxy-benzyl-malonic acid to 190° (Wislicenus, A. 242, 39, cf. Gabriel a. Michael, B. 10, 2204). Prisms, v. sol. lot, sl sol. cold, water.

CARBOXY-PHENYL-SEBACAMIC ACID CO2H.C8H18.CO.NH.C8bc4.CO2H. [193°]. From its ether by saponification. Prisms.

Mono-ethyl ether CO₂Et.O₆H₁₀.CO.NH.C₆H₁.CO₂H. [146°]. From sebacio ether and m-amido-benzoic acid (Pellizmari, A. 232, 146; B. 18, 215; G. 15, 550).

m-CARBOXY-PHENYL-SUCCINAMIC ACID C₂H₃(CO₂H)CO.NH₂O.H₄,CO₂H. Bensamsuccinic acid. [223°]. Colourless prisms. Its ethyl-ether is formed, togother with di-phenyl-succinamide distribuyine acid, by boiling amidobenzoio seid with an alcoholic solution of snccinio ether. On heating to its melting-point it loses H.O and is converted into succinyl-amidebenzoio acid U2H2 CO N.C4H4.CO2H [235°].

Ethyl ether C2H2(CO2Et)CO.NH.C4H4.CO2H [1740]; glistoning plates from water.

C2H2(CONH2)CO.NII.O6H4.CO2H Anvide [229°].

4 nilide C2H2(CONHPh)CO.NU.C4H4.CO2H [252°] (Pellizzari, A. 232, 146; B. 18, 214; G. 15, 550; Muretoff, J. R. 4, 298).

p-Carboxy-phenyl-succinamio acid [4:1] CaH, (CO, H). NH. CO. CH, CH, CO, H. [226°]. From p-tolyl-succinimido and dilute aqueous KMnO, (Michael, B. 10, 577). Needles, sl. sol. cold water and cold alcohol. Boiling cone. IIClAq gives succinic and p-amido-benzoio acids.—AgHA".

o-CARBOXY-PHENYL-SULPHURIC CO.H.C.H. O.SO. OH. Salicyl-sulphuric acid. Prepared by the action of K.S.O. on a solution of salicylic acid in strong KOH. By heating the K salt to 190° it gives K,SO, and salicylide. -A"K, Colourless spikes. Readily decomposed by dilute acids into salicylic acid and KHSO. (Baumann, B. 11, 1914).

m-Carboxy-phenyl-sulphnric

[3:1] CO.H.C.H.O.SO.OH. Prepared by the action of K.S.O. on a solution of moxybenzoic acid in strong KOH .- A"K2: [2200-2250 with decomposition]; needles, more stable towards dilute acids than the ortho- compound; is decomposed however at 100°.

p-Carboxy-phonyl-sulphnric acid 6: [4:1] CO₂H.C₄H₂O.SO₂OH. Prepared by the action of K.S.O. on a solution of proxybenzoic acid in strong KOH.—A''K₂: leaflets or tables. Does not decompose till heated to 250°

CARBOXY-PROPYL-ACETIC ACID v. ETHYL-

CARBOXY - PYRRYL - GLYOXYLIC ACID C4NII3(CO2II).CO.CO2H. Formed by the oxidation of pyrrylene - di - methyl - di - ketone (CH, CO), C, H, NH, or of pyrryl-methyl-ketone carboxylic acid, with alkaline KMnO. Crystalline; sol. ether, alcohol, and boiling water,

insol. benzenc.—A"Ag2: yellow pp.
Di-methyl ether A"Mc2: [[145°]; long colourless needles; m. sol. hot alcohol, sl. sol. cher and benzene. Potash-fusion gives pyrol di-carboxyie acid (Ciamician a. Silber, B. 19, 1412, 1957; G. 10, 373, 379).

CARBOXY-TARTRONIC ACID ev. Di-oxy-

TARTARIC ACID.

CARBYLO-DIACYTONAMINE v. p. 27. CARCYLAMINES v. CARBAMINES.

GARDAMOMS, OIL OF. Employed in medicine as a carminative.

2. From Cey' n. The secds of Elettaric major contain 3.5 p.c. of an essential oil which consists of a terpeno (170°-178°), terpinene, a solid substance [61°], and terpineol C10H1, (205°-220°). The latter is converted by into di-pentene hydrochloride, C10H12HCl [520

and by HI into $C_{10}H_{10}$ 2HI [76°]. The terpene also gives a hydrochloride $C_{10}H_{10}$ 2HCI [52°]. A solid tetrabromide could not be got (Weber, A. 238 98).

233, 98).

2. From Malabar. The bil fr. Elettaria Cardamomum slowly deposits crystals of C₁₈H₁₈H₂O (Dumas a. Péligot, A. Ch. [2] 57, 334).

CARDOL C₂₁H₂₀O₂ (2) An fil, occurring with anacardic acid (a, v.), in the pericarp of the cashow nut (Anacardium occidentale). It is sol. alcohol and ether. It is not volatile; it blisters the skin. It gives with basic lead acetate a pp. of C₂₁H₂₀(PbAc)O₂PbO (Slüdeler, A. 63, 137). Anacardio aoid C₂₁H₂₀(OH)CO₂II in alcoholic solution gives with metallic solutions pps. of the salts: AgA'. — CaA' 2aq. — BaA' aq. — MgA' aq. — Its methyl ether is an oil (Ruhemann a. Skinner, C. J. 51, 663; B. 20, 1861).

CARICIN. An oily substance present in the seeds of the Papaw tree (Carica papaya) (Peckelt, Ph. [3] 10, 343).

CARMINAPHE v. N. PHTHOQUINONE.

CARMINIC ACID C₁₁H₁₀O₁₀. The colouring matter of cochineal which is obtained from insects of the genus Coccus, chiefly Coccus cacti. Cochineal contains only 10 p.e. colouring matter (Pelletier a. Caventou, A. Ch. [2] 7, 90; 8, 255; Warren de la Rue, A. 64, 1, 23; Schützenberger, A. Ch. [3] 54, 52; Schaller, Bl. [2] 2, 414; Mène, C. R. 68, 666; Dieterich, C. C. 1867, 287; Liebermann, B. 18, 1969). The lead salt is ppd. on adding lead acetate to an aqueous infusion of cochineal; by this means the quantity of colouring metter is coclinical may be estimated. Carmine, a red pigment prepared from cochineal, app art to be a compound of carminic acid with flumina, lime, and semo organio acid. Cochineal also contains a fat (in which are ethers of myristic acid, of C₁₄H₂₆O₂ sed of C₁₂H₂₇O₄ and a waxy abstance, cocerin Raimann, M. 6, 891; Liebermann, B. 19, 328).

Properties. — Purplo mass, sol. water and alcohol, sl. sol. ether. Its solution forms red pps. with the alkalino earths and with acctates of Pb, Zn, Cu, and Ag. Alum and Na CO, give the aluminum lake.

Reactions.—1. Boiling diluto II SO, forms carnine-red and a sugar $C_0H_1O_4$ (Iflasiwetz a. Grabowski, A. 141, 329). According to Liebermann the formation of sugar is questionable.—2. Potash-fusion gives eoccinin, each acid, and sucinic acid (H. a. G.).—3. Conc. II.SO₁ at 130° forms a compound $C_{ce}H_{20}O_{12}$ and ruficoccin $C_{16}H_{10}O_4$ (Liebermann a. Dorp, A. 183, 105).—4. HNO2 forms nitro-coccic acid $C_{26}H_{20}O_{20}O_{2}$ —5. By distillation with sinc-dust a small quantity of a solid hydrocarbon $C_{16}H_{10}$ is produced, this forms plates melling at [187] (Furth, B. 16, 2169).

Ceccinin C₁₄H₁₂O₂. Prepared as above. Yellow laminæ (from alcohe); insol. water, v. sol. alcohol, sl. sol. ether. Sol. alkalis. The alkaline solutions are yellow and absorb oxygen, becoming green and, finally, purple. The solution in conc. H₂SO₄ turns indigo-blue on warming. On distillation with zinc-dust it yields a

small quantity of a hydrocarbon $C_{i,i}H_{1}$, which forms plates melting at [187°].— $C_{i,i}H_{1}(NH_{i})O_{i}$.

Acetyl derivative. Yellow crystals, sol alcohol and acetic acid, insol. water (Furth, B. 16, 2169).

CaC₁₆H₃O₆.

*Compound C₃₂H₃₆O₁₃. Black insoluble powder. Forms violet solutions in KOHAq and cone. H₂SO₄. Both this compound and ruficocoin give C₁₆H₁₂ [187³] when distilled over zine-dust.

Carmine red C₁₁H₁₂O₂. Formed by boiling carminic acid with dilute 11.8O₁ (v. supra). Dark purple mass with green lustre; searlet when powdered. Alcohol and water form red solutions. Insel, ether. Potash-fusion forms coccinin. Water at 200° forms ruffearmine C₁₆H₁₂O₆.—Salts: K₂C₁₁H₁₆O₇.—CaC₁₁H₁₆O₂.xaq.—BaC₁₁H₁₆O₇.xaq.—Zn(C₁₁H₁₆O₇.xaq.—Zn(C₁₁H₁₆O₇.xaq.—

If carmino-red be dissolved in acetic acid, and treated with bromine two products are obtained, named provisionally (a) and (b) bromocarmine. The (a) bromo-carmine is sparingly soluble in acetic acid and separates in crystals (yield: 10 p.c.), whilst the amorphous (b) bromocarmine remains in solution and is ppd. on adding water (the yield is 20 p.c.).

'(α)-Bromo-carmino' C₁₋₁Π₁Br₁O₃ crystallises in colourless neadles, [278'], v. sol. alkalis. By belling with strong aureons KOII it gives

boiling with strong aqueous KOII it gives

(a) - Bromo - dioxy - carmine, so called,

(a) IBr.O₅, which forms colourless crystals,

(2082). By its behaviour on etherification it is
shown to contain one CO.II and one phenolic

OII group. On exidation with KMnO₄ it yields
two bodies:—(a) An acid C₆II Br.O₄ which forms
calconriess crystals [2112]. By its reactions on
methylation it is proved to contain one OII
and CO.II group, whence it probably has the
constitut on C₆(CII₂)Br₂(OII)(CIIO)CO₂M or

C₆II₁(CI₂)Br₂(OII)CO.CO₂II. (b) A neutral body

C₄II₁Br₂O₄, [1952], which by its reactions is
shown to be a di-bronto-oxy-methyl-phthalio
anhydride C₆MeBr₂(OII)C₂O₂O [1:2:1:3:5:6].

(3). Bromo-carmine,' so called, is the second product of the bromination of carmine-red, and separates in yellow amorphous flocks on adding we er to the acceptancial solution. It is v. sol. alcohor, agente acceptance, out good not be obtained in a pure state. By boiling with strong agreems KOH it is converted into

(B)-Bromo-oxy-carraine' C_1H , Br_2O_4 , which forms glistening yellow needles, [2322]. It is a di-basic acid—and forms red salts. On oxidation with KMnO₁ it yields two bodies:—(a) An soid $C_{1a}H_aBr_2O_4$ which forms colourless prisms (containing aq, and melts at [2302] with evolution of CO_2 . From its reactions it probably has the constitution $C_1MeRr_{21}OH)(CO_1H)CO_2CO_2H_1$. (b) A neutral body $C_2H_2Br_2O_4$ identical with that obtained from the '(a)-brome-bxy-carmine' (Will a. Leymann, B. 18, 3180).

CARMUFELLIC ACID C₁₂H₂₀O₁₆. An acid said to be formed by the action of HNO₂ on the aqueous extract of cloves. Micaccous scales.

ingol. alcohol, ether, and cold water (Muspratt |

a. Danson, P. M. [4] 2, 293).
CARNAUBA WAX. Obtained from the leaves of Copernicia cerifera in Brazil, and largely used there for making candles. It contains myricylic alcohol, C28H38CH2OH [85.5°], a hydrocarbon, [59°] and compound others derived from the following alcohols and acids: myricyl acohol; an alcohol $C_{2a}H_{ab}CH_{a}OH$ [76°]; a di-hydric alcohol $C_{2a}H_{ab}(CH_{a}OH)_{2}$ [104°]; arf isomeride of lignoceric acid, $C_{2a}H_{ab}CO_{a}H$ [72°5°]; an acid isomeric or "identical with corotic acid $C_{2a}H_{ab}CO_{a}H$ [79°]; and an oxy-acid of the formula $C_{a}H_{ab}(CH_{a}OH)(CO_{a}H)$ or its lactone [103°5°]. The alcohol $C_{2a}H_{ab}(CH_{a}OH)_{2}$ gives on oxidation an acid $C_{a}H_{ab}(CH_{a}OH)_{2}$ [102°5°]; and the oxy-acid $C_{a}H_{ab}(CH_{a}OH)(CO_{a}H)$ gives the acid $C_{a}H_{ab}(CO_{a}H)_{2}$ [90] (Stürcke, A. 223, 283; cf. Lewy, A. Ch. [3] 13, 438; Brandes, T. 1811, 261; Maskelyne, C. J. 22, 87; Bérard, Z. [2] 4. an alcohol C26H. CHOH [76°]; a di-hydric 261; Maskelyne, C. J. 22, 87; Bérard, Z. [2] 4, 415). The greater part of the war is myricyl cerotate and myricyl alcohol.

CARNINE C, H, N, O, A substance occurring in extract of meat, and in the product of boiling yeast with water (Weidel, A. 158, 353; Schützenberger, C. R. 78, 493). Obtained by boiling with water the pp. thrown down from meat extract by lcad acetate; the carnine crystallises from the evaporated filtrate (Kruekenberg a. Wagner, C. C. 1884, 107). Crystallises with aq, v. sl. sol. cold water, insol. alcohol and ether. Bromine-water converts it into hypoxanthine

C.H.N.O.—B'HCl: needles.—B'"IL.PlCl., CARPENE C, H14. (156°). Obtained, together with p-crosol, by distilling calcium pohocarpate. Oil, smelling of turpentine; resinities in the air. Forms an oily compound with bromine (Oudenians, B. 6, 1125; A. 170, 252).

CARROTIN C20 II 38 (?) Carrotenc. colouring matter of the carrot (Dancus Carota) (Wackenroder, Geiger's Mag. 33, 144; Zeise, J. pr. 40, 297; Husemann, A. 117, 200). Occurs also as a normal constituent in the leaves of plants, and in the tomato (Arnaud, C. R. 102, 1119; 104, 1113; Bl. [2] 46, 487; 48,64). Inasmuch as no other coloured hydrocarbon is known, carrotin probably egitains oxygen.

Preparation.-The roots are cut up and pressed, dried at 80° and extracted with CS₂. The juice is ppd. with lead acetate and the pp. also extracted with CS. Carrotin, hydroenrrotin, and fat are obtained from the CS2 solutions. The fat is saponified with alequolic potash. Water and Back are a Sec. The Rp. is dried and extracted with acctone. On recrystallising from methyl alCohol hydrocarrotin separates out

first (Reinitzer, M. 7, 597).

Properties.—Small red plates, v. sol. CS., benzono; v. sl. sol. absolute alcohol; and less sol. 90 p.c. alcohol. Insol. Aq. Rapidly absorbs oxygen from the air. Dissolver in conc. ILSO, giving a deep blue colour. Yields a crivative C24H38I2 with iodine; this has a deep green

C₂₂H₁₃B₂ with nonne; this has a deep green colour, and metallic dustre. Chlorine forms a chloro-derivative [120°).

Hydrocarlotin C₂₂H₁₁O (?), [138°]. [a]₀=

-37.4° in CllCl, at -3.4°. Prepared as above. Greatly resembles cholesterin. Colourless, insol. water, v. sol. calcohol, acetone ether, CliCl. and CS, Crystallises from acetone in long

needles, and from methyl alcohol in plates containing water. Resembles Liebermann's oholestol and Hesse's cuprool but differs from phytosterin (Reinitzer, M. 7, 597).

Acety Cericative [128]; coloured green by H2SO4, and rose by addition of chloroform.

Benzoyl derivative [145°] CARTHAMIN C.H.60r. T The colouring matter of safff-wer (Carthamus tinctortus) (Chevreul; Schlieper, A. 58, 352). Washed safflower is treated with aqueous Na₂CO₃, acetic acid is added and pieces of cotton are put in. The carthamine that has been taken up by the cotton is subsequently dissolved off it by aqueous Na,CO3, and ppd. by citric acid. Powder with red lustre (from alcohol); sl. sol. water, insol. ether, v. sol. alcohol. Its alcoholic solution is purple. Decomposed by boiling with water or alkalis. Potash-fusion gives oxalic and p-oxybenzoic acids (Malin, A. 136, 117).

of Origanum hirtum and, together with eymone and a terpene, in oil of Satureja hortensis and S. montana; in oil of mint and of Thymus Scrpyllum (Jahns, Ar. Pn. [3] 16, 277; B. 15, 816; Haller, Bl. [2] 37, 411; C. R. 94, 132;

Beyer, Ar. Ph. [3] 21, 283).

Formation.—1. By boiling carvol (50 pts.) diluted with oil of caraway (50 pts.) with glacial phosphoric acid (12 pts.) for 3 or 4 hours (Lustig, B. 19, 11; cf. Völckel, A. 35, 308; 85, 246; Kekulé a. Fleischer, B. 6, 1088; Kreysler, B. 18, 1704).-2. From camphor (5 pts.) by boiling with iodine (1 pt.) (Kekulć a. Fleischer, B. 6, 934; cf. Claus, J. pr. 25, 264; Schweizer, J. pr. 26, 118; A. 40, 329).—3. From bromo-camplior and ZnCl₂ (Schiff, B. 13, 1408).—4. Pure camplor cymene is converted into its monosulphonic acid and the latter carefully fused with 3 pts. of KOA1 (Jacobsen, B. 11, 1060; cf. Pott, B. 2, 121; H. Müller, B. 2, 130).

Properties .- Oil. Fo2Cl6 colours its alco-

holic solution green.

Reactions. - I. On fusing with KOH isooxycuminic acid CoH3(CO2H)(OH)CH, [1:2:4] is first formed and finally oxy-tercelithalic acid is produced (E. 11, 1060). -2. P₂S₃ gives cymene and thio carvaerol C₁₀H₁₁S₁ - 3. P₂O₃ forms cresol and propylenc. -4. Fe, Cl, gives di-carvaerol. -5. PCl, forms chloro-cymene. -6. Diazebenzene forms C.H.MePr(OH).N.C.H. [800-850] and C, IIMePr(OH)(N2C, II2)2 [126] (Mazzara, G. 15, 214).-7. Chloro-acctic acid in presence of an alkali forms carvacryl-glycollic acid C101113O.CH2.CO211. 8. H2SO4 forms one or two sulplienic feids of the form CaH2MePr(OH).SO3H. According to Julius one only is formed, its salts being: KA'aq. - AgA'2aq. - BuA', 5aq. S. 12.5 at 15°. - MgA', 12aq.

Sodium salt .- C10H13ONa: white crystal-

line powder.

Methyl ether C10H13OMo. (217°). S.G. 2 .954 (Paterno a. Pisati, B. 8, 71; G. 5, 13). Forms with II, SO, two acids C10H12(SO3H)(OMc) whose Bh salts are BaA'₂ 3½aq: v. sl. sol. water, and BiA'₂ 5aq, v. sol. water.

Ethyl ether C,oH,sOEt: (235°); oil having

1884, 787).

Acetyl derivative C16H13OAc: (246°). S.G. 2 10119 colourless liquid heavier than water.

Bensoyl derivative C, II, OBz: (above

200°); thick odourless oil.

Dicarvacrol. $C_{20}H_{20}O_2$ [154°]. Formed by the action of neutral Fe.Cl_a on carvacrol (Dianin, J. R. 14, 141). Thin silky needles (from dilute alcohol); insol. water, v. sol. alcohol and ether.

o-CARVACROTIC ACID C₁₀H₁₂(OII)CO.11.
Oxy-cymene-carboxylic acid. [136°]. Exepared by passing CO₂ over heated codium carvacrol. White silky needles. Sublimable. V. sol. het water, alcohol, and ether, nearly insol. cold water. Alcoholio Fe Cl₈ gives a violet colouration (Luetig, B. 19, 18).

p-Garvacrotic acid. C₁₀III₁₂(OII)CO.H. Oxycymene-carboxylic acid. [80°]. Obtained by oxidation of carvacrotic ridehyde (from carvacrot, CHCl₃, and NaOII) with KMnO₁. Long white silky needles. Can be eublimed and distilled with eteam. V. sol. het water, alcohol, and ether, nearly insol. cold water. Green colouration with alcoholic Fc.Cl. (Lustig, B. 19, 16).
p-CARVACROTIC ALDEHYDE

C₀H₄(CH₃)(C₃H₃)(OH)(CHO) [1:4:2:5] (?) Oxyaldehydo-cymene. (c. 236°). Formed by heating carvaerol with aqueous NaOH and chloroform. Oil. Volatile with eteam (Lustig, B. 19, 14).

An icomeride [96°] has also been described as p-oarvacrotic aldehyde. It is left as a residue after distilling off the volatile aldehydo with etcam. White silley flat plates. Easily soluble in alcohol, ether, and beuzene, sparingly in hot water, insoluble in cold water (Nordmann, B. 17, 2632).

CARVACRYL-AMINE-C. L. I., NH. Methyl-

wopyl-phenyl-amine. (242°). Formed, together with di-carvacryl-amine, by heating carvacrol with amiconiacal Zulbr, or ZuCl, and NII, Br or NII, Cl at 350°-360°; yield, 25° to '.3 p.c. Colourless oil, which solidifies at -16° B'_2H_Cl_PtCl_i, yellow prisms, sl. sol. hot water.

Acetyl derivative Chellar NilAc: [1150]; white glistening tables; el. sol. hot water, v. sol.

warm alcohol.

 $C_{14}\Pi_{13}$.NHBz \bullet derivative Benzeyl[102°]; flat glistening crystals; nearly insol. water, el. sol. cold alcohol, v. sol. hot alcohol,

and benzene (Lloyd, B. 20, 1261)

Di-oarvaeryl-amine (C₁₀H₁₃J₂NII. (344°–348°). Formed as above, the yield is 27 to 40 p.o. Colourless oil. V. sol. alcohol, ether, and bonzene. Its solution in conc. H.SO, is coloured blue by nitrites and nitrates. - B'HCl. -B'H2Cl2PtCl.

Acetyl derivative (C:0H12)2NAc: [78°]; white glietening ecales; v. sol. hot alcohol and ligroin, sl. sol. in the cold (Lloyd, B. 20, 1261). CARVACRYL-GLYCOLLIC ACID C.H.,O, i.e.

C₁₀H₁₃O.CH₂CO₂H. [140°]. From carvacrol, chloro-acetio acid and potash (Spica, G. 10, 345).

Ethyl ether Eth'. [0.100°]. (289°).

Amide C₁₂H₁₀O₂NH₂. [68°].

CARVACRYL-LACTIC ACID C₁₃H₁₀O₃ i.e. C16H13O.CMeH.CO2H. [74°]. From carvacrol, a-chloro-propionic acid, and potash (Scichilone,

an odour of carrots (Lustig, B. 19, 11; C. C. | G. 12, 49). Prisms, v. e. sol. alcohol, ether, and chleroform.

Carvacryl Mercaptan C, H, S 4.2. C, H, MePr(SH) [1:4:2]. (236°). 8.G. 173 998. From camphor or carvacrol and P.S. (Flesch, B. 6,478; Roderburg, B. 6, 669; Kekulé a. Fleischer, 8. 6, 934). Liquid. HNO₂ oxidises it to sulphotoluio acid (Beelder, J. pr. [2] 8, 168).

Salte.—Hg(C₁₀H₁₈S)... [109°] (Fittioa, A. 172, 327).—C₁₀H₁₂S.HgCl.—C₁₀H₁₃SAg.—

C10H1.SAgAgNO

Methyl ether C₁₀II, SMe. (244°). S.G. 99. TRI-CARVACRYL PHOSPHATE

 $PO(OC_{16}\Pi_{13})_3$. [75°]. Colourless prisms or tables. Easily soluble in alcohol, ether, and benzene, more sparingly in petroleum-other. Formed by heating carvacrol with POCl₃; yield, 55-60 p.o. of the theoretical (lefeysler, B. 18, 1704).

CARVACRYL PHOSPHORIC ACID

C₈H₄MePr.O.PO₈H₄ [1:4:2]. Formed by the action of POCl, uppn carvacrol, and treatment with aqueous K. Co,. The potassium calt A'K5aq forms large silvery plates. By alkaline KMnO it is oxidised to oxyisopropyl-salicylio acid C_nH_n(CMc,OH)(OH)CO₂H [4:2:1] (Heymann a. Königs B. 19, 3309).

TETRA-CARVACRYL SILICATE Si(OC, His)4. (380°-390°) at 118 mm. Colourless oil. Formed by heating carvacrol with SiCl,; the yield is 85 p.c. of the theoretical (Hertkorn, B. 18, 1694).

CARVACRYL-SULPHURIC ACID C.H.MePr.O.SO.H[1:4:2]. Cumyl-sulphuric acid. The potassium salt is formed by adding potassium pyrosulphate to a warm colntion of carvaerol in aqueous KOH. Silvery plates. V. sol. water and alcohol. By alkaline permanganate it is oxidised to oxyisopropyl-salicylic acid C, H, (CMe,OH) (O11) CO,H [4:2:1] (Heymann a. Königs, B. 19, 3309).

CARVENE. A terpene present in oil of caraway, v. TERPENES.

Nitroso-carvene r. Chivoxim.

CARVEOL C12H15. OH. (219°). Thick fluid. Formed by reduction of carvol with sodium and alcohol. With phenyl wante it reacts to form carveyl-phenyl-carbamate [84°] (Leuchart, B.

29, 114). CARVEYL PHENYL-CARBAMATE

C₁₆H₁₅O.CO.NPhH. [84°]. Formed by the action of phenyl eyanate upon carveol C10H15.OH. Slender needles. V. sol. hot alcohol, sl. sol. ether and ligroin (Lenchart, B. 20, 114).

CARVOL C₁₆H₁₁O. (228°) (R. Schiff, B. 19, 50.2). S.G. ¹¹ 9667 (Gladstone, C. J. 49, 621); II.F.p. 48,250 ((C,O₁) = 91,000; (H₂,O) = 69,000) (Stohmann, J. p., $\lfloor 2 \rfloor \otimes 4,322$).

Occurrence.—In oil of caraway (oleum carvi) together with carriero (173°) (Völckel, A. 85, 246). In oil of dill (Anethum graveolens) and of mint (Mentua crispa). The carvol in the oils of caraway and of dill is dextro-rotatory, but that from oil of mint is levo-rotatory $[a]^{\mu} = -62.46$ to Plückiger (Ar. 1/h. [3, 21, 238). According to Plückiger (Ar. 1/h. [3] 22, 361) the rotatory power of carvol ie [a]_p = 58.2°.

Properties. Liquid. Carvol from all three

sources forms the same crystallino compound (C₁₀H₁₄O),11.8 [187°] when H₂S is passed into its alcoholic solution. When prepared from oils

of oaraway or of dill this compound is dextrorotatory, $[a]_0 = +5.5^{\circ}$ at 20°, but when obtained from oil of mint it is lævo-rotatory, $[\alpha]_D = -5.5^{\circ}$ at 20°. Dilute alcoholic KOH in the cold liberates carvol from this compound. Protracted treatment with H2S converte carvol in alcoholio solution into the amorphous (C, H14S)2H2

Reactions .- 1. Distillation over solid KOH or P₂O₃ changes carvol into the isomeric carva-crol Kekulé a. Floischor, B. 6, 1088).—2. P₂S₃ forms cymens.—3. P₂S₃ gives thio-carvacrol C10H13SH.-4. Distillation over heated sine-drat gives C₁₆H₁₈ (173°) and cymene (Arndt, Z. [2] 4, 730; B. I, 204).—5. Sodium in alcohol forms ... carveol (q. v.). -6. Dry HCl gas passed into a mixture of carvol (Lorol.) and aceto-acetic ether (I mol.) forms the compound $C_{16}H_{2}$.ClO₄ possibly $C_{9}H_{15}$ Cl:C(OH).CH($CO.CH_{3}$) O_{2} Et. [146°]. Glistening white prisms (Goldschmidt a. Kisser, B. 20, 489). - 7. Hydroxylamine forms the oxim, v. CARVOXIM.

Carvol - phenyl - nydrazide C, 11, N, HC, Hs. [106°]. Formed by the action of phenyl-hydrazino on carvol (Goldschmidt, B. 17, 1578). Slender white needles. Sol. hot water.

Carvol-chiere-hydride C10111.ClO. 111ydrochlorocarvol. Oil. Formed by leading dry HCl into carvol.

 $Oxim C_{10}H_{15}Cl(NOH)$: [132°]; Formed by the action of hydroxylamine upon carvol-chloro-hydride or of HCl upon carvoxim.

Benzoyl-oxim $C_{lo}H_{lo}Cl(NOBz)$: [115°]; needles (from petrolenn-spirit) (Goldschmidt a. Zürrer, B. 18, 2220).

Phony l - hydrazide C10 II 15Cl(N2 IIPla): [I37°]; small white prisms.

Carvel bromo-hydride C,oH,BrO. Oil. De-

composing at about 50°. Oxim C10H15Br(NOH): [1160]; prisms

(from ligroin). Phenyl-hydrazide C10H,,Br(N.HPh):

[119°]; slender yellow needles (Goldschmidt a. Kisser, B. 20, 488, 2071).

Constitution of carvel. HC- Pr = CH Carvol IIC-CMeII -- CO is probably the pseudo-form of IIC - CPr - CH Carvacrol $H\dot{C} - CMe = \dot{C}(OH)$

(Goldschmidt, B. 20, 490). According to Gladstone (C. J. 49, 621) the presence of two pairs of doubly-linked atoms of chebor in the molecule of travol is 'n'ed by its molecular re-

CARVOXIM C10H1. N(OII). Nitroso-hesperidene or nitroso-carvene. [71°]. Large colourless cransparent plates. Sol. ceids and alkalis.

Formation.—I. By the action of hydroxylamine upon carvol.—2. By passing nitrosyl chloride into a methyl-alcoholic solution of carvene, and heating to its melting-point the crystalline by trochloride C10H16ONCl which pre-

Reactions, I. By heating with dilute H.SO. carvol is regenerated.—2. By hassing HCl gas into its methyl-alcoholic solution the oxim of sarvol chloro-hydride (v. supra) is formed.

Hydrochloride B'HCl; white crystalline!

solid; decomposed by water; formed by passing HCl into the ethereal solution.

Methyl ether C10H14:N(OMe): colourless

Benson deris ative C10H14:N(OBz): [950] white glistening needlee, v. sol. alcohol and benzene (Goldsohmidt a. Zürrer, B. 17, 1577; 18, 1729).

Iso-carvogim C₁₀H₁₄(NQH). [1438], possibly CPr < CH.C(NOH) CMC. Obtained, together with a small quantity of oarvoxim, by the action of excess of hydroxylamine on a colution of carvol chlorohydrido or bromo-hydride in alcohol (Goldschmidt a. Kisser, B. 20, 2071). Necdles. sl. sol. alcohol; sol. acids and alkalie. Unlike carvoxim, it does not combine with HCl or HBr. Dilute II SO, forms carvacrol and a compound C₁₀H₁₅NO [94°].

Benzoyl derivative [112°]; scales, v. sol. alcohol. $C_{10}H_{14}(NOBz)$:

CARVYLAMINE C₁₁I₁₅NH₂. Formed by reduction of carvoxim C₁₀H₁₄,NOH in alcoholic solution, by sodium-amalgam and acctic acid. Colourless liquid, of strongly aromatic basic odour. Readily absorbs CO, from the air.— B'HCl: [c. 180°], slender silky needles (from alcohol).

Benzoyl derivativ. C10H15.NHBz: [169°]; white needles (Goldschmidt a. Kisser, R. 20, 486).

CARYOPHYLLIN C, H_{al}O₄ (?) A substance that may be extracted by alcohol from cloves, the dried flower-buds of Caryophyllus aromaticus (Mylius, B. J. 22, 452; Muspratt, Ph. 10, 313). Silky needles in stellate groups; sublimes at about 285°. Sl. sol. cold alcohol, sol. boiling alkalis. PCl₅ forms C₄₀H_{cs}O₅Cl and C, H, O, Cl,

Acetyt derrvative [181°]. Monoclinio crystals (Hjelt, B. 13, 800).

Caryephyllic acid C₁₆H₅O₁₂. From caryephyllin and fuming HNO₂ (Mylius, B. 6, 1053). Amorphota; sl. sol. water, v. sol. alcohol, ether, and HOAc. May be crystallised from fuming . HNO.

Salts. -Na,A"". -Ag,A"". - Ba,A"" 1½aq.

CASCARILLIN C₁₂H₁₈O₄. [205"]. S. 127 at
100°; S. (alcohol) 3:33 at 8°. Extracted from cascarilla bark (from Croton Elemberia and Cascarilla) by boiling water (Mylius, B. 6, 1051; cf. Tuson, C. J. 17, 195; Duval, J. Ph. [3] 8, 91). Minute prisms (from alcohol); tastes bitter. Not alrected by boiling dilute HCl. Cancarilla bark also contains a volatile oil (173"-I80°).

CASEÏN v. PROTEÏDS.

CACEOSE v. PROTEÏDS.

CASSONIC ACID C.H.O. Formed, together with saccharic and oxalic acids, in the oxidation Also from glyconic field and IINO, (Hönig, J. 1879, 667). Syrup. Reducee ammoniacal AgNC, to a mirror.— PaA"xaq.

CASTORIN. Castorenm is a hard black substance (soft when fresh) found in a pair of small sacs situated in the genital organs of the beaver (Castor fiber and americanus). An alcoholic extract deposits first fat, and then castorin. Castoreum also contains a volatile princent oil,

cholesterin, a resin, proteins, CaCO, and in-organic salts (Valenciennes, 7, 1861, 803). CASTOR OIL. A fatty oil obtained by

pressure from the seeds of Ricinus communis. It solidifies at about -18°, has S.G. about 969 at 12°, and is dextro-rotatory, $[\alpha] = 12^{\circ}$ (Popp, Ar. Ph. [2] 145, 233). Castor oil consists chiefly of glycerides of stearic and ricinoleic acids. It is completely dissolved by 5 vols. of 90 p.c. alcohol (Hager, C. C. 1976, 389). Dry distillaition gives aerolein, conanthol (heptoic aldehyde) and an acid (C₁₁H₂₀O₂)_x (Bussy a. Lecanu, J. Ph. 13, 57; Stanek, J. pr. 63, 138; Lecds, B. 16, 290; Krafft a. Brunner, B. 17, 2985). HNO₃ oxidises it to heptoic, oxalic, azelaic, suberic, and (B)-pinfelic acids (Arppe, A. 120, 288). The products obtained by sapenifying castor oil and distilling the resulting alkaline ricinoleato alone or with NaOH are methyl hexyl ketone, sec-octyl alcohol, and sebacic acid (Neison, C. J. 27, 507, 837). Conc. H.SO, converts castor oil into ricinyl sulphune acid C₁₈H₃₃O₂OSO₃II, which by the addition of we ter breaks up into ricinoleio acid and M.SO,. From the fatty acids derived from the Turkey-red oil prepared from castor oil, crystals of a di-oxy-stearic acid separate after some time (Benedikt a. Ulzer, M. 8, 217)

CATALPIC ACID C14H14O6. [206°]. tracted by ether from decoctions of the siliquaccous capsule of the Bignonia Catalpa. It may be isomeric with hydrocardenic acid (Sardo, G. 14, 134). Large white crystals, v. sl. sol. water, sol. alcohol and ether. — BaC, IH 1206 Gaq; white glistening lamina. — Ag. A": a white pp. CATALYSIS v. Chemical change.

CATECHINS $C_{1b}H_{1s}O_s$ 3aq (Illasiwetz; Cross a. Bevan, C. J. 41, 92); $C_1H_{1s}O_s$ (Etti, M. 2, 547); C₂₁H₂₀O₂ (Liebermann & Tauchet, B. 13, 694); C₄₄H₂₀O₄ and C₄₂H₂₀O₄ (Gautier, C. R. 80, 668). This name has been given to various compounds contained in catechu or Terra japonica which is extracted by boiling water from the fruits or twigs of a variety of plants : Boinbay catechu from the fruit of Areca Catechu, Bengal catechu from twigs and unripe pods of Acacia (or Mimesa) Catecha, Gambir catechu from Nauclea (Uncaria) Gambir, and Nubian catechu from some Acacia. Catechu is used in dyeing.

Catechin C., II, 0, (Gautier, C. R. 85, 752); C, II, 0, 3aq (C. a. B.); C., II, 0, 5aq (I. a. T.). [217°]. S. (alcohol) 20 at 15°; S. (ether) 8 at 15° (Wackenroder, A. 37, 311). Obtained from Bombay catechu by washing with water and crystallising from acetic ether (L. a. T., Löwe, Fr. 13, 113; Zwenger, A. 37, 320; Kenbaner, A. 96, 337; Kraut a. Van Delden, A. 128, 285; Hlastvetz a. Malin, A. 131, 118; Etti, A. 186, 337; Schützenberger, Bl. [2] 4, 5; Sace, C. R.

53, 1102).

Properties .- Small needles (from water). V. sl. sol. cold water, v. sol. hot water and acetic ether. The aqueous solution is coloured green by Fe₂Cl_w. The solution in KOHAq absorbs oxygen, turning brown's Lead acetate gives in aqueous solution a pp. of (C₂₁H₂₆O₆).3PbO (?) Catchin solutions are ppd. by albumen, out not by gelatin.

C20H119Cl4O12? (Cross a. Bevan, C. J., 41, 92) which is turned crimson by Na SO. Catschutaunio acid does the same .- 3. Br givss bromocatechuretin $C_{21}H_{2}Br_{2}O_{2}$, a red insoluble powder. 4. Water and PI_{3} give $C_{21}H_{21}O_{8}$, an elastic insoluble mass. — 5. HOAc and BaO₂ give C21H20O10 a colourfess powder which melts below 100° (Schitzenberger a. Rack, Bl. 4, 8). 6. Aqueous K.Cr.O, forms C.H.4010, a brown insoluble powder.—7. Polash-fusion gives Phloroglusin and protocatechnic acid (Hlastwetz, A. 134, 118).—8. Dry distillation gives pyrocatechin.—9. Boiling dilute H.SO, forms insoluble $C_{21}H_{16}O$, (Neubauer, A. 90, 356), or $C_{23}H_{10}O_{12}$ (Etti).—10. Boiling dilute KÖll forms $C_{21}H_{10}O_{12}$? a brown powder, sol. alcohol and alkalis.-11.

III gives iodoform and other products (G.).

Di-acetyl derivative C, 11, O, (OAc),:
[131°]; needles or prisms. Soluble in ordinary solvents except water and ligroin (L. a. T.)

Di-ben pyl derivative Call 180, (OBz),

Flocculent brown substance (S. a. R.). Diacetyl-dichloro-catechin

 $C_{21}H_{16}Cl_2(OAc)_2O_3$: [169°]; needles. Sol. alcohol, al. sol. ether.

Diaetyl-bromo-catechin

 $C_{21}H_{17}Br(OAc)_2O_7$; [120°]. White needles. Sol.

alcohol, sl. sol. ether.

Catechuretin C42H30O176aq(?) or C38H28O12 Formed by passing HCl into a boiling alcoholic solution of catcelin (Kraut a. Delden, A. 128, 291). Formed also by heating catechin with conc. HCl at 170°. Dark reddish-brown in-

soluble powder. Not changed at 190'.
•Di-benzoyl derivative C2,H14Bz2O1(?) Formed, tog ther with di-benzeyl-catechin by heating catechin with BzCl at 190'. Brown

Catechin C₁₀H₂₈O₁₆ 2aq. [205°]. S. 9.9 at 50°. Occurs recording to Gautier (C. R. 86, 668) in Gambir-cateelin together with the two following catechins; *they are extracted by alcohol and crystallise after evaporation with exclusion of air. Monoclinic prisms.

Catechin Carlino, aq . [177°]. Minute needles (v. supra)

Capechin $C_{40}H_{48}O_{16}$ aq. [163°]. S. 5·3 at 50°. Minute needles (v. supra). Catcchiu C, H1,O, aq. According to Etti

(M. 2,547) this is the formula of the catechin in Gambir and Pegn catechins. At 100° it becomes C₁₈H₁₆O₅ at 160° catechutamic acid C₃₆H₃,O₁₅ and "t 180° C₃₆H₃₂O₁₄. The latter is also got by heating catechin for some time with dilute

Reactions. -1. Diazobenzene chloride gives (C,H,N,),C,H16O, n red crystalline pp sol. alcohol and other; it dyes wool golden-brown.— 2. Dilute 1 SO, (1:8) at 140° gives phloroglucin and pyrocate hin. .

Catechin C.H.,O. [165]. In mahogany (Acajon) (Gantie, Bl. [2] 30, 568). Latour a. Cazencuee (Bh [2] 24, 119) give this catechin the formula C₂₀H₁₀O₉.

Catechin $C_{42}H_{36}O_{16}$. [140°]. In brown cate-

CATECHOL v. Pyrocatechin.

CATECHUIC ACID v. CATECHIN and PROTO-CATECHUIC ACID.

Reactions.—1. Boiling dilute H.SO, forms CATECHUTANNIC ACID O21H18O4(?) or catechuretin.—2. With IICl and KClO3 it gives C32H31O13(?) Extracted by water from catechu.

Formed also by heating catechin alone at 130°, with water at 110°, or by boiling it with alkalis, lime, or Pb(OH)₂ (Etti, A. 186, 332; Löwe, J. pr. 105, 32, 75; Z. [2] 5, 538; Fr. 12, 285). Dark reddish brown powder. V. sol. acctic other, v. e. sol. alcohol, insol. ether; m. sol. water. It oxidises in the air. It gives & greyish-green pp. with Fo₂Cl_o. It does not pp. tartar-emette. Its aqueous solution is prod. by gelatin, albumen, and by dilute H₂SO₁. At 162° it changes to C,2H34O,5(?) which resembles catechutaun racid

in all respects – (C., H,,O,) 3PbO.

CATHARTIC ACID. The active principle in senna leaves. It is a glucoside. It contains only O, H, and Q, Its Ba and Pb salts are amorphous (Kubly, 5tt. [2] 7, 356; Stockman, Ph. [3] 15, 749; cf. Lassaigne a. Feneuille, A. Ch. [2] 16, 18; Bourgoin, C. R. 75, 1449).

CAULOSTERIN v. CHOLESTERIN.

CEDAR OIL. Obtained by distilling with water the wood of Juniperus virginiana. Contains cedrene and cedar-camphor. According to Bertagnini (C. R. 35, 800) it contains a com-

be betagnin (C. 7t. 5d, 500) it contains a compound which combines with NaIISO,

Cedar-camphor C₁₅H₂₅O. [74°]. (282°).

V.D. 8·4 (calc. 7·7). Crystalline mass cuclling like cedar-wood. V. sl. soi. water, v. soi. alcohol. Distillation with P₂O₅ splits it up into water and cedrene (Walter, A. Ch. [3] 1, 498).

CEDRENE C₁₅H₂₁. (237°). S.V. 7·6. S.G. 15·984. Obtained as above (Walter, l.c.).

Cedrene. From oil of sage (English). C15H24. (260°). S.G. 15 .915. Yellow or green oil. Inactivo. Resinified by H.SO₄ (4:1) even at 0°. Gaseous HCl turns an ethereal solution purple. The refractive index seems to indicate four C:C groups (M. M. P. Muir, C. J. 57, 686).

The name Cedrene has been used as a generic name for the hydrocarbons C15H21 which occur in the oils of cedar, eloves, patchouli, cubebs, calamus, cascarilla, rosewood, &c. (v. TEIWENES). Cedrenes closely resemble the terpenes in their optical properties, which point to the existence of $1\frac{1}{9}$ pairs of doubly linked carbon atoms (Gladstone, C. J. 49, 617).

CFDRIRET v. CERULIGNON.

CELLULOSE. "{C_uII₁₀O₅}. *S.G. 1·25-1·45. Occurrence.—Cellulose is the basal substance of the skeleton of plants, and indeed may be said to constitute the framework of the vegetable world. The problem of its origin is as much physiological as chemical. It does not appear to be formed as the immediate product of tho synthetical action of the cell upon carbonic anhydride and water, but me liately from starch, sugar, and other carbohydrates, through the intervention of the cell protoplasm. The mechanism of this transformation, as well as tho inverse conversion of cellulose into the simpler carbohydrates, has not been elucidated, but is assumed on physiological grounds to be of the simplest character. There is nothing in this assumption which contravency the evidence afforded by the chemical relationships of the

carbohydrate group, which are likewise simple.

Adapting itself to the infinite variety of structure and function presented by plant tissues, cellulose occurs in multitudinous forms: and in any given structure is subject to differentiation, modification, or variation of elaboration within very wide limits. The scope of this

article, however, precludes such a treatment of the subject as would deal with lesser variations, and we shall therefore confine our attention to those celluloses which constitute fine fully elaborated plant fibres. Plant tissues seldom if ever consist of pure cellulose but contain besides other products of growth, either mechanically bound up with the tissue, and therefore frequently removable by mechanical means and by the action of simple solvents, or chemically united to the cellulose; combinations of this latter kind constitute the compound celluloses, and are only resolved by a chemical process.

Prevaration.—The isolation of pure cellulose

depends upon its relative insusceptibility of chemical change. The general method of preparation from raw fibrous materials consists in exposing the moist fibre or tissue to the action of chlorine gas or bromine-water in the cold and subsequently boiling in a dilute alkaline solution; repeating this treatment until the alkaline solution no louger disselves anything from tho tissue or fibre. The celluloso is then washed with a dilute acid, water, alcohol, and other, and dried.

Properties .- Obtained in this way, or by tho ordinary process of bleaching from cotton or linen (flax), or in the form of Swedish filter paper, the typical cellulose is a white substance more or less transparent, retaining the microscopic features of the raw fibre.

The elementary composition is expressed by the percentage numbers (F. Schulze):

or by the corresponding empirical formula C, II, Os. These numbers represent the composition of the dry and ash-free cellulose. Nearly all celluloses contain a certain propqrtion, however small, of mineral constituents 2 and the union of these with the organic portion of the fibre or tissue is of such a mature that the ash left on ignition preserves the form of the original. It is only in the growing point of certain young shoots that the cellulose tissuo is sometimes found free from mineral constituents (Hofmeister). The proportion of hygroscopic moisture, which is an essential constituent of celluloso under ordinary atmospheric conditions, varies from 7 to 9 p.c.; the mean variation due to variations in the hygrometric state of the air is about 1'p.c.

Cellulose is insoluble in all simple solvents; it is dissolved by certain reagents but only by virtue of a preceding constitutional modification. The most remarkable solvent of cellulose is cuprammonia (Schweitzer's reagent) in which it dissolves without esseutial modification, being recovered by precipitation in a folim which is chemically identical with the original (Erdmann, J. pr. 76, 385), though differing in being amor-

4. Cellulose in its earlier stage of elaboration has no action upon light, but with age it acquires the property of thoulier refraction. This action is independent of the state of aggregation of the cellulose and is therefore an essential control of the cellulose and is therefore an essential control of the cellulose and is therefore an essential control of the cellulose and is therefore an essential control of the cellulose and is therefore an essential control of the cellulose and is the cellulose.

on aggregation of the cellulose and is therefore an essential property of the substance itself (Sachs, Exp. Phys. d. Phys. de. Phys. d. Phys. de. Phys. de.

phous. This reagent has been employed in a variety of forms, a fact which explains the diecrepancies in the etatements as to the solubilities of the various collulosee in ouprammonia. The following methode of applying the reagent are to be recommended.

The substance to be operated upon is intimately mixed with copper turnings in a tube which is narrowed below and provided with a stopcock. Strong ammonia is poured upon the contents of the tube and after standing for some minutes ie drawn oil and returned to the tube; the operation is several times repeated until the colution of the cubstance is directed.

Perhaps the most convonient solution, though not so effective in all cases as the former, is that prepared by dissolving ppd. cupric hydrate in ammonia. In preparing the reagent in this way it is important that the hydrate should be thoroughly washed, preferably out of contact with the air, before diesolving in the strong aqueous ammonia. Cotton is rapidly dissolved by this The soluble compound formed is rosolution. presented by Muldor as $(\hat{C}_0H_{10}O_s)_2Cu(NH_1)...O$. It has been doubted whether this compound exiets actually dissolved in the viscous solution; an investigation of the osmotic properties of the liquid, however, ebows it to be a true solution (Cramer). From an Extended investigation of the optical properties of the solution Béchamp concludes that the solution of the cellulose is not eimple but is accompanied by progressive molecular transformations, the optical activity (dextrorotatory) of the products increasing to a maximum corresponding to a condition of equilibrium ultimately attained (C. R. 100, 117, 279,

The solubie bases (NaOH, KOH) added to the solution give blue gelatinous pps having the composition $(C_oH_{1o}O_s)_2CuM''O$. Digested with finely divided lead oxide the solution yields the compound C₆H₁₀O₅.PbO. Celluloso is reppd., as a gelatinous hydrate, on the addition of acids, as well as of many neutral bodies such as alcohol, suga , and common salt, or even on largely diluting with water and allowing to stand. The pp. dried in vacuo is obtained as a transparent mass resembling gum arabic. On digesting the annonia-cuprio solution upon metallic zinc, this metal pps. she copper, replacing it in the solution and producing the corresponding ammoniazincio colution of cellulose, which is colourless. The property of cellulose of being dissolved by cuprammonia receives an important technical application. A sheet of paper left for a short time in contact with the cuprammonia, so that the constituent fibres are superficially attacked, and then passed between rollers and dried, becomes impervioue to water and its collesion is not affected at the boiling heat. Two sheets thus treated adhere firmly together, and with a cufficient number, artificial boards are produced. A variety of materials are now produced in this way, on the manufacturing scale, useful for roofing and other purposes (C. R. A. Wright, Journ. Soc. Chem. Ind. 1884, p. 121).

Reactions.—Cellulose has already been spoken

of as a comparatively inert substance, and its characteristic reactions are consequently few. One of these ie available for ite identification and is chiefly used in the microscopical exami-

nations of tiesues: thie is ite reaction with iodine. The reaction, although similar to that of etarch, differe in requiring for ite determination the presence of an auxiliary (dehydrating) reagent euch as sulphurioor phosphorie acid or zine ohloride. The most effective solution is prepared in the following way: zine is dissolved to eaturation in hydrochloric acid and the colution evaporated to the sp. gr. 20; to 90 pts. of this solution are added 6 pts. potassium iodide dissolved in 10 pts. water. and in this solution iodine is dissolved to saluration. By this reagent cellulose ie coloured instantly a deep blue or violet.

Compounds of Cellulose .-- Celluloso is generally inactive towards compounds contained in diluto aqueous solution; hence its extensive employment in the filtration of solids from solutions. Nevertheless it exhibits a tendency to incipient combination even with acids and alkalis (Mills, C. J. 43, 153); with metallic salts it forms compounds of sufficient stability to cause their removal from solution, but the combination is of an indefinite and unstable order (Erdmann, J. pr. 76, 385). (Cf. Gladstone, J. pr. 56, 217; Müller, Fr. 1, 84; O'Shea, C. J. Proc. 1, 206.) Certain sarbon compounds, such as the organic astringents, and many of the colouring matters natural and artificial, unite with cellulose to form compounds of various orders of stability; of these we would more particularly instance amongst others many of the derivatives of diphenyl which possess a specific power of direct combination with cellulose. Although such combinations are of great technical importance, being the foundation of the arts of dyeing and printing they are not sufficiently systematised to descree more than this passing notice. On the other hand some of the substitution-compounds of cellulose with acid radicles are both definite and stable.

Acetyl-collulese.—The tri-substituted compound $C_1\Pi_1('\cdot,\Pi_3O)_2')_3$ is formed by heating cellulese with 6-3 times its weight of acetic anhydride at 130°, and separates as a white flocculent pp. ou diluling the syrupy product. Tri-acetyl-cellulose is insoluble in alcohol and in other, and is soluble in glacial acctio acid. It is saponified by boiling with alkaline solutions, the cellulose being regenerated. No derivative containing more than three acetyl groups has been obtained; but a mixture of the mone and di-acetylcellulose is formed by treating cellulose with only twice its weight of acetic anhydride, the formation of these bodies being unattended by their solution.

lose.) -- Whenever cellulose in any form is brought into contact with strong nitric acid at a low temperature, a nitro-product or nitrate is formed The extent of the intration depends upon the concentration of the acid, upon the duration of its contact with the cellulose, and on the state of the physical division of the cellulose itself The first investigation of these substances dates from 1838, when Pel uzo showed the identity of several of these products obtained from paper, linen &c. and starch. Knop and also Kamarsch and Heeren found that a mixture of sulphuric and nitric acids also formed nitrate of cellulose; and still later (1817) Millon and Gaudin employed a mixture of sulphuric acid and nitrate of potash and sods, which they ound to have the same effect. Although gunottons or pyroxylines are generally spoken of s nitro-colluloses, they are more correctly decribed as cellulose nitrates, since they have not een found to yield amido-bodies on reduction vith nascent hydrogen. The following are the eneral properties of these compounds (Eder): 1) When warmed with alkaline solutions, nitrio icid's removed in varying quantities, dependent upon tife strength of the solution emfloyed. 2) Treatment with cold concentrated sulphurie acid expels almost the whole of the nitric acid. (3) On boiling with ferrous sulphate and hydrochloric acid, the ritrogen is expelled as nitrie oxide; this reaction is used as a method of nitrogen estimation in these bodies. (4) Potassium sulpliydrate, ferrous acetate, aud many other substances, reconvert the nitrates into cellulose.

Several well-characterised nitrates have been obtained, but it is a matter of difficulty to prepare any one in a state of purity and without admixture of a higher or lower nitrated body. The following have been described under a nomenclature having reference to a C12 formula:

Hexa-nitrate.—C₁₂H₁₁(NO₃),O₄ (gun-cotton). Prepared by treating cotton with a mixture of HNO, (S.G. 1.5) 3 parts, and H.SO, (S.G. 1.84) 1 part, for 2I hours, at a temperature not exceeding 10°; 100 parts of cellulose yield about 175 of the compound (calc. 183). Insoluble in alcohol, ether, or ruixtures of both, and glacial acetic acid; with acetone it forms a jelly and is slowly dissolved. It is the most explosive of the ceries igniting at 160°-170°. Mixtures of sulphuric acid and nitre do not give this nitrate (Eder). Ordinary gun-cotton may contain as much as 12 p.c. of nitrates soluble in etheralcohol; the hexa-nitrate seems to be the only one quite insoluble in this menstruum.

Penta-nitrate.—C₁₂H₁₅(NO₃₎₅O. This composition has been very commonly ascribed to gun-cotton. It is impossible to prepare it in a state of purity by the direct action of the acid on cellulose. It is prepared by dissolving the hexa nitrate in nitric acid at 80°-90°, croling to 0°, and adding concentrated sulphuric acid, by which the penta-nitrate is precipitated; after mixing with a large volume of water and washing the precipitate with water and alcohol, it is dissolved in ether-alcohol and finally re-precipitated by water. This nitrate is slightly soluble in acctic acid, nearly insoluble in alcohol cortaining only against proportion of ether. Strong potash solution converts it into the dinitrate.

Tetra- and tri- nitrates (collodion pyroxyline) , are generally formed together when cellulose is treated with a more dilute said at a higher temperature and for a shorter time than in the case of the hexa-nitrate, e.g. 4 vols. HNO, (1.38), 5 vols. H₂SO₄ (1.84) at 65°-70° for 5 19 minutes. They are freely and equally soluble in otheralcohol, acetic ether, and mixtures of acetic acid and wood spirit, or al bhol, and are therefore inseparable. They are insoluble in pure alcohol, ether, or acetic acid. On treatment with concentrated nitric and sulphuric acids they are converted into the higher nitrates. Potash and ammonia convert them into the dinitrate.

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Di-nitrate C12H1a(NO2)2O4 is formed as already indicated, and also by the action of hot dilute nitric acid on cellulose. Freely soluble in etheralcohol, acetic ether, acetic acid wood spirit, acetone, and absciute alcohol. The further action of alkalis on the dinitrats results in a complete resolution of the molecule.

The colluloso nitrates have generally much stronger absorption affinities for coleuring matters than the colluloses. They are much less susceptible of attack by acid 'oxidants than cellulose itself, and are therefore used in the filtration of solutions containing e.g. ohromic acid, permanganates, and, of course, nitric acid of any degree of concentration. Natric acid of S.G. 1.42 has a remarkable touthening action upon filter paper: the modification is effected by simple immersion, and the paper so treated is increased in strength tenfold, undergoing at the same time a contraction amounting, in circles, to about 10 diameter. The cellulose so treated contains non-trogen (Francis, C. J.

Celluless and sulphuria acid .- Cellulose is disintegrated and dissolved by the concentrated acid to a colourless solution. The products are sulphates of a series of compounds of which the celluloses and the dextrins may be regarded as the extreme terms. They are easily isolated in the form of Ba salts. The composition of the sulphates may be expressed by the general formula $C_{\nu}\Pi_{\nu m}O_{\nu m-\nu}(SO_4)_{x}$. The variations in mula $C_{c_n}H_{1e_n}O_{5n-2}(SO_4)$. The variations in composition and in physical properties are functions of the temperature (5° 334) and duration of the action. The limits of specific rotation of these sulphates are $[\alpha]_j = -3.65$ and +72.99. These ethereal salts are entirely decomposed by boiling with alcohol: the resulting carbohydrates may be regarded as the corresponding alcohols. In composition and properties (e.g. their reactions with iodine) they constitute an extended series, beginning with soluble celluloses and terminating in achrodexton (Hönig a. Schwoert, M. 7, 455). While it is impossible to determine the mechanism of these successive resolutions of the cellulose molecule with the precision attainable where such changes may be reversed, and therefore completely studied, thoy certainly establish the typical connection of the celluloses with the simpler earbohydrates, and in a much more complete way than the coincidence of empirical formulæ.

Prior to the researches above detailed the initial terms of the transition series had been similarly obtained and described under the term Amyloid, a term selected to indicate their resemblance to starch.

A practical application of the reaction of cellulose with sulphuric acid is found in the manufacture of parchment paper. The process consists in the rapid passage of the paper through the strong, acid (S. f. 1-5-1-6). followed by copious washing. The result may be described as a superficial conversion of the cellulose into amyloid, whereby it acquires the properties which have obtained for it the designation in question (cf. Hofmann, A. 112, 243).

" Callulose and chlorins .- Dry ohlorine has no action upon cellulose; the presence of water determines an indirect oxidising action, but there is no direct combination of celluloss with

chlorins. By heating osllulose nitrates with phosphorio psutachloride and oxychloride at 200° and syaperating the sxcess of the reagents at 170°, a viscous liquid is left, miscible with alcohol and other, which appears to la composed of, or to contain, a chloride of cellulose or a cellulose derivative (Baeyer, B. 2, 54). Broming is without action upon cellulose: specimens left for several months in contact with strong bromins-water were not sensibly attacked (H. Müller, Pflanzenfaser, p. 27; cf. Frauchimont, R. T. C.

Celluloss and Oxygon. Oxycelluloses. - Two of these compounds or series of compounds have

been described

(a)-Oxycellulose (Witz, Bull. Rouen, 10, 416; 11, 189) is formed by the action of solutions of the hypochlorites upon cellulose. Exposed to the action of a solution of bleaching powder (5 p.c.) for 24 hours, the fibre is converted into a friable modification having the composition C 43.0, H 6·2, O 50·8. Other oxidising agents produce similar results; even by exposure to air and light cellulose is slowly converted into these oxidised derivatives. Their formation is accompanied by molecular resolution, as is shown by their reducing action upon alkaline copper solutions: the product giving this reaction is dissolved by alkalis to ryellow solution. These oxycelluloses have a strong attraction for basic colouring matters: also for vanadium compounds, attracting these from a solution containing so minute a quantity as 1 pt. in 1,000,000,000. Upon this property a method has been founded for the quantitative estimation of minute traces of variadium in aqueous solution (Witz a. Osmond, Bull. Rouca, 14, 30). The study of these exycelluloses is of great importsnce to the manufacturer of lettile materials.

 (β)-Oxycellulose C₁₈ II₁O₁₆ (Cross a. Beyan,
 C₄ J. 43, 22) is the residual product (insoluble) of the prelonged digestion of cellulose with nitric acid (20-30 p.c.) at 90° C. On washing with water to remove the acid the substituce gelatinises. It dries to a horny colourless mass. is characterised by its reaction with sulphuric acid: on gently warming it dissolves with develop-ment of a bright pink colour, the reaction resembling that of mucic acid, to which, on other grounds, it is probably related. A fresh preparation, treated with a mixture of nitrie and sulpharic acids, dissolves, and on pouring into water the nitrate C₁₈II₂₃(NO₃)₃O₁₃ separates as

a white flocculent pp.

Chromic acid.—Cellulose treated with potassium dichromate in presence of acetic acid is converted into glucose, dextrin, and formic ucid. Pormanganates under the same condition effect

a similar accomposition.

Chromic anhydrido in presence of sulphuric acid decomposes cellulose rapidly and completely, the carbon being entirely converted into the gaseous compounds CO and CO. It has been proposed to apply this to the quantitative esti-mation of carbon in celluloses and cellulosio mixtures (Cross a. Bevan, C. N. 52, 207).

Alkaline oxidations.— The permanganates

and hypochlorites in presence of alkalis oxidise cellulose to a syrupy mixture of acids of the pectio series (H. Müller, Pflanzenfaser, v. also S. C. I. 3, 206, 291). Fused with potassium

hydrato the cellulose is oxidised to oxalio acid, malic acid being obtained as an intermediats

Electrolytic oxidants.—The nascent oxygen and other electronegative ions liberated in tho electrolysis of various saline solutions have a very powerful action upon cellulose. These actions have been made the subject of interesting rescarches by F. Goppelsroeder (D. P. J. 254, 42).
• Calluless and Hydrolytic Agents.

(1) Dilute acids. A large number of acids, organic as well as mineral, attack celluloso, producing hydration changes, attended by disintegration of the fibre. The action is gradual at ordinary temperatures, and is of course accelerated by applying heat. The study of these actions is of the first importance to the cellulose technologist (Girard, C. R. 81, 1105; Cross a. Bevan, S. C. I. 1885; Crookes, Handbook of Dycing and Calico-printing, p. 19).

It is worthy of note here that the celluloso isolated from grass and hay, and many others less highly claborated than the celluloses which we are at present considering (cotton and linen), are decomposed on boiling with dilute mineral

acids with formation of furfural.

(2) Alkalis.—Dilute solutions of the alkalis are without sensible action upon cellulose, even at temperatures considerably above the boiling point: when, however, oxidising conditions are superadded, molecular resolution sets in. Tho joint action of calcium hydrate and air at the boiling temperature is especially powerful, oxycellulose being produced (Witz, loc. cit.). Concentrated solutions of the alkalis (NaOii, KOII) at ordinary temperatures act in a very remarkable way upon cellulose. There appears to be a 'molecular' combination of the reagents in the proportion $C_{12}\Pi_{20}O_{10}$. Na.O (Mercer) which however is easily resolved by washing with water. But the characteristics of the fibre and the fibro substance are found to have undergone a permaneut modification. There is a considerable shrinkage in linear dimensions: in cotton fabrics treated with caustic soda solution of S.G. 1.225 this amounts to 25 p.c. The corresponding modifications in microscopie features been investigated by Crum (C. J. 1863), the changes being found similar to those which take place in the ripening of the fibre in the plant, viz. from a flattened tube with large central cavity, to a thick-walled cylinder with small lumen. The chemical change produced is, to far as has been ascertnized, entirely one of hydration, and it is remarkable that the only evidence of the change is the increased capacity for hygroscopic moisture. This amounts to 5 p.c. of the weight of the cotton, the proportion ealculated for the formula C12H20O10.H2O being 5.5 p.c. From this fact we may also infer that the normal attraction of cellulose for atmospheric moisture is a residual manifestation of the molecular combinations which are seen in the multitudinous hydrates of cellulose found in, or constituting, growing tissues. Morcer, who appears to have first investigated these phenomena, further found that the addition of hydrated oxide of zine very much increased the action of the caustic solution: thus a solution of sodium hydrate of S.G. 1 100, which is without marked action, is rendered very active by

the addition of the oxide in the proportion ZnO:2Na,O. He also found that the actions were favoured by low temperatures (v. Life of John Mercer by E. A. Parnell, London, 1886).

By these characteristics the hydration phenomena in question are seen to be closely related to those attending ine action of the Schweitzer reagent (supra). The more powerful action of the latter we must refer either to the specific action of the "ammonia upon't the condensed addehydio molecules of which cellulose appears to be constituted, or to the particular relationship of the molecular weight of the cupranumonia in solution to that of the cellulose or cellulose hydrate which it forms.

The action of concentrated solutions of zino chloride is similar to that of the alkaline hydrates above kydrates. It is remarkable on the other hand that a saturated solution of zine

nitrate is without action (Mercer).

(3) Water.—Heated in contact with water, cellulose is attacked at 160°, but not below 150° (Scheurer a. Crosseteste, Bull. Mulhouse, 1833, 62-85). Heated at 200° in contact with water in scaled glass tubes it is fundamentally resolved, being converted into highly-coloured products, insoluble for the most part, with a small proportion of soluble derivatives amongst which are furfural and pyrocatechol (Hoppe-Seyler, B. 4, 15).

(4) Ferments.—There are, it can scarcely be doubted, endless transformations of cellulose determined by the so-called soluble ferments, though but few have been investigated. The soluble ferment of the toxglove is stated to convert cellulose into glucose and dextrin (Kosmann, Bl. [2] 27, 246). The duid from the vermiform appendix of the rabbit has also been found to digest cellulose with liberation of marsh gas and formation of a soluble compound which reduces cupric oxide in alkaline solution.

Preximate Syntheses of Cellulose .- Transformations of the soluble earbohydrates into cellulose, which we may regard as a proximate synthesis of this body are, as already stated, an important feature in the life of plants. The mechanism of these changes has been thus far but slightly studied, and they are of a kind to elnde chemical investigation. Of those which have been studied we may notice (a) There is a change set up 'spontaneously' in beet juice which results in the formation of a hard white substance, having the properties of cellulose. On transferring these lumps to a solution of pure cane sugar, a further transformation of the saccharose into the same shostance is brought about. At the same time there is produced a gummy substance which is ppd. by alcohol as a white caoutchouc-like substance of the same composition as cellulose but swelling up with water and otherwise differing in its physical properties from cellulose. This latter derivative is also formed by the action of diastase upon a solution of saccharose. A similar transformation takes place nnder the induence of certain fatty seeds, e.g. those of rape and colza and it is probable that the formation of collubose from succharose in the plant takes place under the influence of ferments similar to those above described (Durin, C. R. 82, 1108).

(b) More recently A. Brown has investigated

the formation of cellulose by the 'vinegar-plant,' growing in solutions of the carbohydrates, e.g. dextrose in yeast-witer. The cells elaborate an extra-cellular film, which acts as a coll-collecting medium, and they possess, therefore, a two sided activity, i.e. the property above mentioned, in addition to their strictly fermentative activity. The cellulosic film in question was found to contain 50-60, p.e. pure cellulose. It is noteworthy that in a solution of levulose the growth of the 'plant' is unattended by fermentative action, 33 p.c. of the substance being on the other hand transformed into cellulose (C. J. 48, 432).

OTHER FORMS OF CELLULOGE.

We cannot attempt to enumerate the multitudinous varieties of cellulose which the plant world presents. Some of these, when isolated in the pure state, resemble the typical cellulose above described, e.g. the cellulose of hemp and rhea. Others, especially such as require a drastio process of resolution, e.g. the cellulose isolated from jute and wood by the chlorination method (infra), resemble rather the (a) expeciluloses. Tims jute cellulose (3C₈H₁₀O₃H₂O) reduces cupric oxide in alkaline solution, and is much more susceptible of degradation by hydrolytic reagents than those of the cotton type. Cellulose from pinewood is similar in composition and properties.

Cellulose from esparto and straw, isolated by treatment of the plant substance with alkaline solutions boiling under pressury—which are alongst the most important of the staple materials of the paper-neaker—are distinguished by their reaction with aniline salts, being coloured a deep pink on boiling with solutions of these compounds. Many of the celluloses are decomposed on boiling with fillule acids with formation of furfural and formic acid: hay cellulose yields under certain conditions a volatile crystallino body, which appears to be a furfural derivative, but is still under investigation.

It may be mentioned here that the term cellulose is applied by plant physiologists and agricultural chemists to substances which would not come under the definition, upon which this article is based, of cellulese as the (white) insoluble residue which survives the exhaustive treatment of plant substances alternately with chlorine, bromine, or oxidising agents, and boiling alkaline solutions.

Animal ceftulese.—The mantles of the Pyrosomidæ, Salpidæ, and Phallusia mammilaris, freed from chondrigen by boiling in a Papin's digester and further purified by prolonged boiling with potast solution, yields a residual substance which not only has the ultimate composition of cellulose, but has identical properties, e.g. dissolves in cuprammonia, is converted by nitrio acid into an explosive nitrate soluble in ether (Schäfer, A. 160, 312). According to Virchow cellulose is found in degenerated human spleen and in certain parts of the brain (C. R. 37, 492, 860).

COMPOUND CELLULOSES.

Plant tissues, always containing a proportion of cellulose more or less large, frequently contain other constituents so intimately united to

the cellulose as to mask its reactions. From the circumstances of their occurrence and formation it is not to be expected that the line can be sharply drawn between mixtures and combinations of cellulose with non-cellulose constituents of either fibree or tissuee. Frémy (Ann. Agronomiques, 9, 529) recognises the exietence of the following compounde dietinguiehed from cellulose chiefly by their different behaviour to hydrolytic reagents and cuprammonia; (1) Fibrose, constituting the membranes of wood celle; (2) Paracellulose, constituting the membrane of the pith and medullary rays, and (3) Vasculose constituting the mombranes of the vassels. The value of this somewhat arbitrary classification is aucstionable (Sachsse, Farbstoffe, Kohlehydrate, &c., p. 150), and the distinctions which it eeeks to establish have not been generally recognized. On the other hand, there are certain groups of substancee widely distributed throughout the plant world, which, while they have certain featuree in common with the celluloses, are sufficiently distinct to admit of chemical classification apart from them. Generally speaking, these substances are made up of a collulose and a non-cellulose portion, the latter conferring the epocial features of distinction. The compound cellulose thus constituted is resolved, by treatment 47th reagente according to the methods to be described, into cellulosc or a cellulose residue on the one hand, and soluble derivatives of the non-cellulose on the other. Of these groups we shall consider typical members.

The following are the compound callulosee sufficiently characterised to warrant special description under class names: the nomenclature of these compounds is explained by their cha-

racteristic resolutions.

Pectocelluloses

Resolved by hydrolysic (alkalic)
into

Pectio acid and cellulose.

Type: Raw flax
(Kolb, Bull. Mull. 1868, A. Ch. [4] 14, 348).

Lagnocelluloses

Resolved by chlorination into

Chlorinated derivs.: Aromatic and collulose.

Soluble in alkalis J Type: Jute (Cross a. Bevan, C. J. 41, 90).

Adipocelluloses

Resolved by exidation
(Nitric acid)
iuto

Similar derive.:
as by the oxidation and cellulose.
of the fats.

Type: Cuticular tiesue of leaves and fruits (Fremy, C. R. 48, 667; Sachsse, Farbstoffe, &c.).

Pectocellulosee.—The purified bast of a Russian flax investigated by Kolb was found to have the aggregate composition:

C 43.7 H 5.9 O 50.4

The non-cellulose constituent is therefore a substance of lower carbon percentage than cellulose. From its yielding pectic acid on boiling with alkalis, it is obviously a substance allied to pectin. Since the fibre yields about 20 pto of its weight of the acid derivative we infer independently that it is a substance containing approximately 41 p.c. carbon, which confirms the view of its constitution above expressed. Many other plant fibres are made up of or contain pectocelluloses (Welster, C. J. 43, 23); pectic derivatives were identified by Schunck in the products of the action of boiling alkaline solutions upon raw cotton (Proc. Manchester Lit. Phil. [3] vol. iv).

The collular tissue of a large number of fruits, c.g., the apple and pear, and roots, e.g., turnips and collustrates, ie composed of pecto-collustrates.

loses

Lignocellulosee. The course of lignification in plants is marked by profound alteratione in the physical properties of the tissues undergoing this modification; the tissues lose their elasticity, become coloured from grey to brown, and the substance of the tissues unanifosts the chemical properties about to be described.

It may be regarded as composed of

Non-cellulose (Lignin) C₁₁H₂O₁₀ (55 5 p.c. C) 25 p.c. 3 and Cellulose C₁₁H₂O₁₅ (44 4 ,) 75 , though the cellulose isolated from the fibre by chemical resolution differs in composition as already stated from normal cellulose, appearing rather as an oxycellulose. The mineral constituents of the raw fibre (purified) vary from 0.5 to 2 p.c. of its weight; the hygroscopic moisture from 10 12. Attacked by concentrated solutions of the alkalis similarly to cotton; freely soluble in cuprammonia, but is incompletely precipitated on acidifying; the body remaining in solution gives the reactions of the original substance, and may, therefore, be regarded as a hydrated modification. Jute combines freely with the organic astringents (tanmins) and the majority of aromatic colouring matters. It is coloured a bright yellow by solutions of salts of aniline and other aromatic bases. This reaction is probably due to a product of oxidation, sinc. it does not take place with jute which has been boiled for some time in solutions of sulphilities. It is coloured brown by iodire solutions. Moistened with a solution of pilor glue, and treated afterwards with hydrochleric acid it gives a deep red colouration; with pyr.ol also in presence of hydrochloric ao'a it gives a deep carmine colour. A mixture of sulphuric and nitric acids nitrates the fibre, the g in in weight being approximately equal to that of cellulose under the same conditions. The products are orangecoloured and are soluble in acctone. Like the cellulose nitrates, they give no amide derivatives on reduction. Iodine is absorbed by the fibre, the quantity taken up being constant under constant conditions; the resulting compound is not more stable than the iodide of starch. This

tive estimation of the lignocelluloses in cellulosic mixturss. Bromine attacks the fibrs in presence of water, forming ill-defined compounds which are dissolved by alkaline colutions. The operation once or twice repeated eliminates the whole of the non-cellulose; the resulting cellulose amounts to 72-75 p.c. of the weight of the fibre. Chlorine does not act upon the dry fibrs, even when the temperature is raised to 160°, but in presence of water combines rapidly at ordinary temperatures with evolution of heat. chloringted derivative is yellow coloured; it is soluble in alcohol, and from the solution water precipitates fire compound $C_{i,i}H_{i,i}Cl_iQ_{i,j}$ as a yellow floculent mass. This compound gives a characteristic magenta colouration with sodium sulphite solation similar to that of mairogallol 1 (Webster, C. J. 45, 205). It dissolves in solutions of the caustic alkalis with a brown colour, evolving the characteristic odour of the chlorequinones. It yields chloropicrin on distillation with nitric acid. Fused with potassium hydrate it yields protocatechnic aoid. It is therefore an aromatic derivative, and appears to be allied to the trihydric phenols. The chlorinated fibre when boiled with sodium sulphite solution is entirely resolved into cellulose, and soluble derivatives of the non-celluloso or lignin constituent. This constitutes the simplest and most rapid method of cellulose estimation in the fibre. The proportion of cellulose obtained is somewhat higher than by the bromine method (75-78 p.c.) and is further increased by preventing rise of temperature in the chlorination.

Digested with dilute nitric acid at 80° jute is resolved directly into cellulose (oxycellulose) oxalio and carbonic acids and a peculiar acid derivative of the lignin constituent, body has the formula C_2 , $H_{10}(NO_2)O_{23}$; it is of an intense yellow colour, and dyes the animal fibres to a similar shade. It forms salts with the sarthy bases (C25H22(NO2)O22.M"4) which are precipitated by alcohol from aqueous solutions in

the form of bright yellow flocks."

Hydrolytic agents.—Jute and the lignocelluloses generally are much more susceptible of hydrolysis than the simple celluloses: diluts acids effect a simple hydrolysis at 80°, i.e. the portion dissolved has the same composition and properties as the original; on raising the temperature to boiling, furfural is obtained in considerable quantity. Boiling dilute alkalis effect a simple hydrolysis. When the hydrolysis is complicated by the introduction of either reducing or oxidis. Tonditions, resolution into collulose and non-collulose (soluble derivatives) is determined. Thus sulphurous acid, the hisulphitee, and the normal sulphites (alkaline) attack and resolve the lignocelluloses when heated with their solutions under pressure. The

According to Hantzsch a. Schnitter (#: 20, 2033), According to Hantson a Schulter (H. 20, 2033), malrogallol is a species of condensed quinoue chloride—the characteristic molegule being derived from quinoue by replacement of one of its typical O atoms by Cl_n the aromatic linking being at the same time partially resolved. Such a view accords equally well with the properties of the derivative in question.

It is worthy of note that the addition of urea to the dilute acid considerably modifies its action, which becomes one of simple hydrolyds as in the case of sulphuric and by droch oric acids (in ra).

temperatures necessary for efficient resolution,

reaction may be made use of for the quantita- lie. for the isolation of cellulose, are with aniphurous acid (7.5 p.c. SO₂) 90°-100°, with hisulphitss (4 p.c. SO₂) 150°, with normal eulphites (4 p.c. SO₂) 170°-180°, the increase of temperature corresponding with the diminution of hydrolytic power by progressive neutralisation of the acid. The hydrolysis is aided by combination of the reagents with the soluble derivatives, which prevente the reversal of the action at the high temperature, which would otherwise occur. In heating with colutions of the caustic alkalisunder pressure, a high temperature is necessary for complete resolution; a considerable proportion of the reagent is necessary for combining with the products, which under the oxidising conditions are of an acid nature. These facts are of importance in the preparation of cellulose frem lignocelluloses, which is now a widely extended industry (Forestry Exhibition Reports.

Edinburgh, 1885).

Animal Digestion.—It has long been known that the urine of the herbivora contains hippuric acid as a normal constituent, and it has been shown that the benzoyl radicle necessary toform this body is a product of the digestion of lignocelluloes (Meissner). Since the lignocellulose moleculo contains no aromatic compounds in the strict sense of the term (Stutzer, B. 8, 575). the process of digestion must effect the conversion.

Decomposition by Heat.—Cumulative Reso-The celluloses burn in the air with a. lution. quietluminous flame. When heated out of contact with the air they are completely resolved into gaseous and volatile products of the ons hand, and a residual black, mass, containing a high percentage of carbon. Regarding these carbonaceous substances as the products of condensation of cellulose molecules with elimination of water, the process may be viewed as, in the main, one of cumulative resolution by dehydration (Mills, P. M., June, 1877); the cumulate containing hydrogen and oxygen in chemical union with carbon is still in svery sense a compound; taking into consideration, at the same time, its approximation in appearance and properties to the element itself it may be termed a pseudo-carbon. Dehydrathig reagents effect similar resolutions, the lignocelluloses undergoing condensation more readily than the celluloses. The tendency to carbon accumulation which is the main feature of these resolutions is well marked in the vast series of natural products of the decay of cellulosic tissues, from humns to the coals. Though not of pyrogenio origin they neverthelese deserve mention in this connection from considerations of general resemblance.

It is noteworthy that the product; of chlorinating ulmic substances (Sestini, Gaz. It., 1882, 292) are closely similar to those officined from the lignocelluloses. Similar products are also yielded by the cannéi coals.

Other forms of Lignocellulose. Two other varieties of lignocellulose deserve mention, viz. glycolignose, C₃₀H₄₀O₂₁, the substance of fir woods 'glycodrupose,' C₃₀H₄₀O₂₀, the substance of the stony concretion of pears (Erdmann, A. 138, 1; Suppl. 5, 223; Bente, B. 8, 476; Sachsse, Farbstoffe, 151). On boiling with dilute 61 Phil. Mag., 1882, 325.

hydrochloric acid these compounds are said to addition to an interior epidermal layer, which be resolved into a sugar and an insoluble residue, lignose = $\mathbf{C}_{1a}\mathbf{H}_{2a}\mathbf{O}_{11}$, and drupose = $\mathbf{C}_{1a}\mathbf{H}_{2a}\mathbf{O}_{12}$, respectively. On comparing these formula with those of the original substances, the differences are so slight as to be negligible. The action of the acid is probably therefore one of simple hydrolysis. The reactions of these substances are also in other respects identical with those of the jute substance. Although therefore dif-fering from the ligno-cellulose, above described, in ultimate composition, they are essentially of the same order of compounds.

Crude fibre. This term is applied by agricultural chemists to the residual product of the treatment of fodder plants with boiling solutions of certain acids and alkalis, applied successively. The process is a crude imitation of the process of digestion in the animal, and the results which it yields are of purely empirical value. Crude fibre will be found on examination, in most cases, to be a ligno-cellatose and to be further resolved by chlorino in the marner indicated.

ADIPOCELLULOSES. Cork and Cuticular Tissue. From the mode of formation of these tissues it has been concluded, on physiological grounds, as in the case of the lignocelluloses, that they are modified celluloses (Sachsse, Farbstoffe, &c. p. 153). The ultimate composition of cork is represented by the following percentage numbers.: C 65.7, H 8.3, N 1.5, O 24.5. Unlike the lignecelluloses, owever, it is by no means a simple or homogeneous substance, but is resolved by the sition of mere solvents into a number of proximate constituents, some of which are crystalline. The residue from the action of these solvents may be regarded as It is resolved by the true cork substance. the action of boiling nitric acid (20-30 p.c.) into cellulose on the one hand—only a small proportion (3-5 p.c.), however, surviving so severe a treatment- and a series of fatty acids (or products of their decomposition) such as euberic and adipic acids, on the other; the latter amount to about 40 p.c. of the weight of substance treated. If cork be resolved by treatment with sodium sulphite solution, at 166° a soft mass is obtained preserving the structural foatures of the original, but which on slight pressure breaks down into a cellular mass. Cn slight purification this yields a pure cellulose; the proportion obtained being from 9 to 12 pe. of the original cork. From these resolutions into cellulose, and products of decomposition eimilar to those obtained from the fats andwaxes under similar treatment, the substance of cork has come to be regarded as a compound of such molecules; and this view of its constitution, as woll as that of the entire group of substances similar in composition and function, is summed up in the group term Adipocellulose, by which it is proposed to designate them. The cuticular tissues, such as constitute the covering of deshy fruits and leaves, while similar in many respects. are simpler in composition. When purified they are non-nitrogenous and appeal to be home-

Frémy and Urbain have developed (C. R. 100, 19) a somewhat different view of the constitution of these tissues. Taking the protective Ver. I.

is cellulosic, and is soluble in cuprammonia after treatment with boiling hydrochlorio acid, they distinguish the external or true epidermis. which they term Cutose. The substance composing this tissue has the following properties: it is resolved by alkalino saponification into two fatty acids, oleocutic CasH400, a liquid oily body, and stearogutic C_mH₁,O₁, a solid body melting at 76°, soluble in benzene and glacial actic actic, and slightly soluble in boiling alcohol, the solution gelatinising on cooling. The following numbers give the percentage composition of the original entose and of the twe-derivatives.

Cutose . Olcocutic acid . Stearocutic acid 68.4 66.6 \mathbf{II} 8.7 8.2 10.3 $25 \cdot 2$ 22.914.2Whence it is concluded that the molecule of

cuteso is composed proximately of olco- and stearo-cutic adds in the molecular proportion . The two derived neids above described are

marked by a carious instability or tendency to reversion passing under certain conditions, notably by exposure of their solutions to light, into modifications closely resembling the original

With regard to the distribution of entose in the plant world, while it is the main constitueut of the external protective tissues of leaves, tleshy fruits and the stems of annuals, it is found also in the interior tissues, e.g. the bast and \$bro-vascular bundless

These researches are an important contribution to the subject; and while there is no reason to doubt the authors' conclusions in the main, it is probable that they have overlooked the presence of celtulose as a constituent of cutose which occurs as an organised i.e. cellular tissue. We need scarcely observe that the chemistry of these adipocelluloses has been but little investigated.

Constitution of Sellulose.

The physical properties of cellulese throw but little light on the problem of its molecular constitution. We have seen, moreover, that cellulose is chemically inert; its derivative comfounds are few, and of these indeed the nitrates alone appear to merit such a description, their formation being unattended by molecular resoution. From their composition and properties we inter the pressure of alcoholic OH groups in the cellulose molecule. The medual resolution by the action of sulphuric acid, through a series of compounds terminating with achreodextrin, indicates a certain constitutional relationship to the simpler earl ohydrates of which the latter is a typical representative. The exact meelianism of the resolution not having been clucidated, it is impossible to draw any but the general inference from the products to the original cellulose molecule, viz., that the latter is a comple of molecules resembling these simpler and better known carbohydrace. We also infer that the bond which unites them is one or dehydration, but the exact nature of this bond is for the present entirely conjectural. (For a discussion of the probabilities involved in this problem the tissues of the leaf of the agave as the type, in reader is referred to a paper by Baeyer entitled S A

*Wasserentziehung u. ihre Bedeutung für das Pflanzenleben und die Gährung, B. 3, 63.) That such molecules are to be regarded physiclogically as well as chemically, as the proximate constituents of the ocllulose moleculo, is a conclusion which has perhaps been somewhat prematurely drawn. Its ultimate origin is of course to be traced to the carbonic anhydride and water of the air and soil, the synthesising agencies being the protoplasm and chlorophyll of the plant, aided by the supplies of energy from without. It has been supposed that the inediate source of the vegetable carbohydrates is formic aldehyde, and the recent researches of Locw (B. 20, 141) upon the condensations of this aldehyde have at least confirmed the probability of this supposition. The producte of resolution of cellulose, moreover, clearly indicate the presence of aldehydic groups in the nuclecules. Apart from conjecture, we may, to sum up this brief review, regard the cellulosconoleculo as a complex of eimpler carbohydrate groups, containing alcoholic and aldehydic oxygen (v. also Bowman, B. A. 1887).

The most striking features of the empirical formula of cellulose C, H, O, are those characteristic of the entire group of carbohydrates, the molecule containing some multiple of C, and the hydrogon and oxygen having the ratio 211. At present these relationships are merely suggestive of conditions of melecular equilibrium to

be elucidated by future investigation.

The prevalence of the Cg group is remarkable and suggests a relationship to the aromatic group, which is confirmed by the undoubtedly cellulosic origin of the benzenc compounds. On the other hand we have no evidence of the existence of closed chains of C atoms in the cellulose molecule, nor have any simple transitions from the one group to the other been as yet discovered. If we turn, however, from the typical cotton cellulose to other forms of celmlose, such as those isolated from growing tissues, we find in the ease with which many of them yield furfural as a product of acia hydrolysis, some evidence of & more complex union of the C atoms, than the normal type usually assumes. Passing on further to the lignocelluloses we find a molecule in the constitution of which a furfural group undoubtedly plays a part, and in which the linking of the C atoms is such as permits a simple transition, in part at least, to products of undoubtedly 'aromatic' composition.

We have, in this brief statement of the evidence which well as upon which ato found our viewe of the constitution of cellulose, endeavoured rather to indicate the main lines upon which the investigation of the problems involved is proceeding, than to draw premature conclusions. The subject will be enormously developed in the future, and will constitute an important foundation of the natural history of the carbon compounds; a department of fiew of the science which can scarcely as yet be eaid to be within

reach.

NOMENCLATURE.

Considerable confusion has been imported into this subject by the indiscriminate employment of the terms, celluloee, lignin, woody with nearly boiling water, dried and extracted fibre, lignose, 'crude fibre,' in describing the with alcohol. The first extract deposits very

various products, natural and artificial, of which this article treats. The want of uniformity arises from the division of the subject amongst physiologists, agriculturists and ohemists. That which we have adopted appears to bring about

a certain simplification.

The application of the term cellulose we have already defined and limited. To denominate the compound childces, which correspond with the chief modifications of cellulose recognised by physiologists, we use a compound term consisting of cellulose as the substantive portion with a qualifying prefix. To specially distinguish the characteristic constituent of these compounds, that to which we have applied the neutral term non-cellulose, we employ the root of the prefix with the termination in, thus pectin, lignin.

In conformity with this plan it may be ex-pedient to introduce such terms as suberocellulose, cuto-cellulose, euberin, cutin, and many others: but this chould be left to be determined by the progress of investigation. The special classification and nomenulature proposed by Frémy has been already discussed, C. F. C.

CERASIN, the meta acid (v. Arabio acid) existing in cherry-tree and plum tree gums, and in the gums of other trees of the came family. Gum arabic yields cerasin when heated to 150° (Gélis, C. R. 41, 144). Cherry-tree gum is said to contain 52.1 p.c. arabin, 34.9 cerasin, 12 water, and 1 ash. Treated with nitric acid ceraein yield 15.5 p.c. mucic acid.

CERATIN v. PROTEIDS.

CERATOPHYLLIN. ° [147°]. Extracted by lime water from the lichen Parmelia ceratophytia (Hesse, A. 119, 365). Thin prisms, sl. sol. cold, v. sol. hot, water; v. sol. alcohol, ether, and aquious alkalis. Fc2Cl8 coloure its alcoholic solution purple. Bleaching-powder gives a red colour.

CERPERIN. A poisonous, cry;talline glucosids occurring in Carbera Odollam (Oudemans, J. pr. 99, 407; de Vrij, R. T. C. 3, 167).

CEREBRIN C.69·08, H.11·47, N.2·13, O.17·32 From these numbers the formula (Parcus. $C_{10}H_{140}N_2O_{12}$; $C_{10}H_{140}N_2O_{13}$; $C_{10}H_{140}N_2O_{15}$ may be calculated). The body called *Phrenosin* by Thudichum (*J. pr.* [2] 25, 19) is eprobably the same body in an impure state. The name cerebrin was used first by Kühn (1828) and applied to a mixture of cholesterin and lecithin. was used by Lassaigno (1830) to denote the entire substance that can be extracted from brain by alcohol. It was employed by Gobley (1850) to denote so-called 'protagon;' and by Müller to denote the nitrogenous substance free from phosphorue that can be extracted from the brain by alcohol, to which he gave the formula C, H, NOs.

References .- Feurcroy, A. Ch. 16, 232; Fremy, J. Ph. 27, 453; Liebreich, A. 134, 29; Gobley, J.Ph. [4] 19, 846; W. Müller, A. 103, 131;

105, 361.

• Preparation Ox-brain is freed from membrance, waehed with water, squeezed and heated with baryta to beiling. The clear liquid is poured off and the coagulated residue washed

little on cooling, but cerebrin mixed with cholesterin separates from the subsequent extracts. The cholesterin is removed by solution in other. Sc prepared, ccrebrin is free from Phosphorus but contains inorganic matter. 250 grms. were got from 90 brains. It is redissolved in alcohol at 60° which leaves a barium salt undissolved. The barium that goes into colution is removed by a current of CO. The corebin is parified by frequent recrystallisations from alcohol. earlier mother-liquors deposit on standing a gelatinous pp. By recrystallisation from alcohol this pp. may be separated into cerebrin (spherical crystals), homocerebrin (needles), and encephalin (P. Parcue, J. pr. 132, 310).

Properties.—Separates as a white crystalline powder composed of transparent globules from a boiling alcoholic solution. Soluble in acctoue, ohloroform, benzene, and glacial acetic acid. Insoluble in other. Separates from acctone partly as globules, partly as matted threads. Cerebrin dissolves in conc. H.SO, On standing • Cerebrin dees not combine with acids, bases, or salts. It is not affected by boiling for a short time with baryta. It is but slightly decomposed by boiling alcoholic prash. Cerebrin swells up but slightly in boiling water. It is very slightly hygroscopic, absorbing 2 p.c. of water from the air. It is decomposed by boiling for a long time with baryta. Boiled for some hours with dilute HCl it forms a solution that can reduce Fehling's solution.

Homocerebrin C. 70-1-p.c.: II. 11-6; N. 2-2; O. 16-1. The yield is 1 that of the cerebrin. Soluble in the same liquids as cerebrin; soluble also in boiling ether. Swells up in hot water, but does not form a paste. Is not decomposed by boiling water. Treated with hot IICl, it forms a solution that reduces Felding's solution. Boiled for a long time with baryta, it is decomposed. It is not hygroscopic. Its solutions are neutral. It does not combine with acids, bases, or salte. It eeparatee from alcoholic solutions in very fine needles. After separation from alcohol and drying, it is not a white powder (like oerebrin) but a waxy mass. It is more soluble

in absolute alcohol than cerebrin. Encephalin. C. 68.4; H. 11.6; N. 3.1; O. 16.9. Resembles homocerebrin rather than cerebrin. It separatee from solutions in flexible plates. From acotone it separates in granular masses. It swolls up with hot water forming a complete paste. Boiled with HCl, it forms a solution

that can reduce Fehling's solution.

CERIC ACID, C. 64-2 p.c.; II. 8-8 pm.; O. 27-0 p.c. Obtained by the action of HNO, on cerin, or according to Kügler on phellonic and $C_{22}H_{12}O_3$ [96°]. Phellonic acid is obtained, together with glycerin and stearic acid, by the action of alco-holic KHO on cork that has previously been exhausted by ohloroform and alcohol (Döpping,

GERIN C₂₀H₂₀O(?). [250° A substance that may be extracted by chloroform from cork, the bark of Quercus Suber, in which it occurs to the extent of 2 p.c. (Kügler, Ar. Ph. [3] 22, 217; ef. Chevreul, A. Ch. 96, 170; Döpping; A. 45, 289). Needles, insol. water, solf other solvents. According to Siewert (7. 1978, 383) an alcoholic extract of cork contains phellyl alcohol $C_{11}H_{26}O$ [100°], a neutral, crystalline substance, sl. sol. alcohol.

CERIUM. Ce. At. w. 139.9. Mol. w. unknown as olemont hae not been gasified. S.G. (abt. 18°) 6.63 to 6.73 (Hillebrand a. Norton, P. (150, 472). M.P. considerably above that of Sb (150°) but below that of Ag (950°) (H. a. N.). SH. (0°-100°) '01479 (E., P. 158, 7). SVS. Chief lines in emission spectrum 5352.2, 27.1. Oner mes in comson special for the first of the fir Berzelius and Hisinger, separated a new earth from a Swedish mineral and called it Ceria (from the then recently discovered planet Ceres); in 1839-41 Mosander (P. 11, 406) proved that ceria was a pixture of at least three metallic oxides, oxides viz. of Ce, La, and Di. Tho metal cerium was first obtained by Mosander in 1826, it was more fully examined by Wöhler, Bunsen, and other clemists.

Occurrence.- In many Swedish minerals, exposed to the air, the liquid becomes covered more especially in cerite a silicate of Ce (abt. with a purple skin, which afterwards turns green. 56 p.c. Ce₂O₃); occurs as silicate, phosphate. carbonate, fluoride, &c., also in many apatites; generally accompanied by La, Di, Fe, Al, &c. It has been recently found that clay used for brickmaking at Hanistadt (near Sceligenstadt in the neighbourhood of Frunkfurt) contains from 8 to 12 p.c. of Ce₂O₃ (Strohecker, J. pr. [2]

33, 133 a. 260).

Preparation. - Cerite is heated to redness, powdered, neixed with considerable excess of core. H.SO., and the mixture is gradually heated to incipient redness in a Hessian crucible; after cooling, the mass is powdered and then added. little by little, to ice-cold water, whereby sulphates of Ce, La, and Di are dissolved; the treatment with H2SO, &c. is repeated with the portion insoluble in cold water; the aqueous solution is heated to boiling, whereby basic oulphates are ppd.; solution in ice-cold water and ppn. by heating to boiling are repeated several times (Marignac; Bunsen; Wöhler). The solution in cold water is ppd. by addition of exalic acid solution; the oxalates are washed with . water, Tried, and heated in a Pt dish until completely converted into oxides. The mixed oxides (of Ce, La, and Di) may now be treated in many different ways. The following method was used by I ranner (C. J. 47, 834) in order to prepare pure Ce 380, for atomic weight determinations; it is based upon the formation of basic Co nitrate insoluble in water. The mixed oxides were dissolved in moderately conft: HNO, Aq. excess of acid was removed by evaporation, and the syrup-like liqual was poured into much pure boiling water; 'he ppd. basic ceric nitrate was washed (on a fungel connected with a pump) with boiling water containing a little nitric acid. The nitrato was dissolved in HNO, Aq, excess of acid was removed by evaporation, the liquid was poured into boiling water, &c., as already described. This treatment was repeated 10 or 11 times. From the pure basic ocric nitrate thus obtained other Ce compounds may be prepared. Brauner prepared Ce. 380, by dissolving the basic nitrate in dilute H. SO, Aq, and H. SO, Aq, evaporating to dryness in a Pt dish, and heating with due precaution, dissolving In a little icecold water, ppg. by absolute alcohol, washing

ppg. by aloohol, dissolving again, heating to 100° and stirring with a glass rod; Ce₂3SO₄6H₂O was thus obtained; it was dehydrated by heating for some weeks at 440° in molten sulphur; at full redness the sulphatengives off SO, and

SO, and leaves pure CeO...

Robinson (Pr. 37, 150) prepared puro Cc. Cl. The mixed oxides from cerito were dissolved, in HNO, Aq (v. supra), and the liquid, after concentration to a syrup, was poured into boiling diluto H.SO.Aq; the basic sulphates were dissolved in HNO.Aq, the Ce salt oxidised by boiling with PbO₂ (Gibbs, And S. [2] 37, 352), the liquid evaporated, and & a and Di salts removed by treatment with HNO, Aq; the pure basic ceric nitrato remaining was converted into chloride by treatment with ECIAA, from this Ce oxalate was obtained by ppn. with oxalic acid; the oxalato was heated in dry HCl, and pure Ce2Cl6 was thus obtained (for details v. original). Robinson (l.c.) also recommends evaporating the nitrie acid solution of the mixed nitrates to complete dryness, heating the brown mass to full redness until the residue is pale-yellow, and treating this with boiling dilute HNO, Aq; te basic nitrate remains while nitrates of Di and La dissolve.

Other methods for preparing more or loss pure Ce salts are described by Bunsen, P. 155, 375; Candnowicz, J. pr. 80, 16; Watts, C. J. 2, 147; Holtzmann, J. 1862, 136; Jolin, Bl. [2] 21, 533; Erk. Z. [2] 7, 100; Popp, A. 131, 361; Pattison a. Clark, G. N. 16, 259. Wöhler (A. 144, 251) prepared Ce by adding pieces of Na to a molten mixture of Ce.Cl., KCl, and NH,Cl; the product, however, was ifapure. Hillehrand a. Norton (P. 155, 633; 156, 166) by electrolysing Cc.Cl., covered with NH Cl, using a thick iron wire as negative electrode (4 Bunsen's cells) obtained approximately pure ecrium. (For details

v. original papers.)

Properties and Reactions.—Steel-grey, very lustrons, very ductile, metal; malleable, unchanged in dry air, but in moist air is superficially oxidised; heated in air burns to oxide with production of heat and much light; burns in Cl. Br. I vapour, S vapour, P vapour, forming compounds with these elements; easily dissolved by IICIAq, dilute HNO, or H2SO4Aq, no reaction with cold cone. HNO, or H2SO, Aq. Ce slowly decomposes cold water, quickly decomposes hot As the V.D. of no Co compound water. has yot been determined, the At. w. of the metal must be fixed by chemical considerations and by S.H. The value given to the At. w. for many years was abt. 92, and the formulæ of the oxides, and chloride, &c., were CcO, Cc2O3, · CeCl, &c.; Cc was thus placed, along with La and Di, in the same class as Al and the earth metals. Considering the relations between the properties of compounds of Ce, La, Di, and of other elements, Mendelejeff (A. Suppled. 8,186) thought that Cc ought to be classed with those elements which form well-marked oxides MC, i.e. with Ti, Zr, Sn, & If this were dono tho At. w. of C. would become $92 \times \frac{3}{2}$. Further investigation has completely justified Mendeleieff's proposal; Ce is now classed with C, Si, Ti, Zr, Sn, Pb, and Th; La is placed with Al and the earth metals, and Di is classed with the elements

with abs. alcohol, dissolving in ice-cold water, of Group V. whose characteristic highest ofide is M2O5.

The At. w. of terium has been determined (1) by estimating SO, in cerous sulphate (Beringer, L. 42, 134; Rammelsberg, P. 55, 65; Hermann, J. pg. 30, 184; Marignac, A. Ch. [3] 27, 200; 38, 148). (2) by estimating Cl in cerous chloride (Boringer, Le.; Robinson, Pr. 37, 150). (3) from combustions of cerous oxalato (Jogel, A. 105, 45; Rammelsberg, P. 108, 44). (4) by conversion of Co₂SO₄ into CeO₂ by hoat (Brauner, C. J. 47, 879). (5) by determining S.H. of approximately pure Ce (Hillebrand a. Norton, P. 156, 471). The most accurate determinutions are those made by Robinson and Brauner, respectively: both lead to the number 139.90 (0 = 15.96).

Cerium is distinctly metallicein its physical and chemical properties; it forms the oxides Cc.O. and CcO. (and probably also CcO.), both of these are salt-forming in their reactions with acids, but the woll-marked salts (cerous salts) are those corresponding to Ce₂O₃, e.g. Ce₂3SO₄ &c.; in this respect Ce differs from Te, Zr, Th, Sn, and Pb, the salts of which metals belong to the forms MX and MX2(X = SO4, CO3, 2NO3, &c.); and shows analogies with the elements of Group III. (Al &c.). Ce does, however, form a ceric sulphate Ce2SO,; besides the oxido CeO, a fluoride CeF, analogous to TiF, ZrF, &c., is known (v. Carbon oroup of ELEMENTS; also

TITANIUM GROUP OF ELEMENTS.)

Detection and Estimation .- Ce salts are generally soluble in water: all alis pp. white hydrated Cc.O.; NaClO. 1 pps. yellow hydrated Cc.O. which dissolves in HClAq in presence of alcohol to form a colourless liquid; solutions of Ce salts are ppd. by oxalic acid, the ppd. oxalate is white and ins huble in NH, salts; saturated K.SO Aq pps. a white crystalline double sulpliate, scarcely soluble in cold water, decomposed and dissolved by hot water containing a little HCl; Co salts toiled with PbO, and a lattle HNO, Aq give a yellow solution. When a slightly acid solution of a Ce salt is mixed with ammonium acetate, a little H.O.Aq is added, and the whole is warmed, a yellowish pp. is formed (Hartley, C. J. 41, 202); if this pp. is moistened with NH₃Aq, again treated with H₂O₂Aq, and warmed, au orange solid (? CeO,) is produced (de Boisbaudran, C. R. 100, 605). Co can be separated from La and Di only by very prolonged treatment, v. Pravaration. Co is usually estimated as CeO, but it is very difficult to obtain this oxide pure. For methods of separation and estimation a manual of analysis must be consulted.

Cerium, carbide of. Said to be obtained as a dense black powder, insoluble in hot cone. acids, by heating cerous formate or exalate in a stream of H, and treating with HClAs to dissolve out any oxido formed. Analysos agree approximately with composition CeC. (Delafontaine, J. 1865. 177).

Cerium, halois and oxyhaloid compounds of. Cerium forms cerous haloid compounds, Co.X. or CeX2, corresponding to the oxide Ce2O; and also a perio fluoride CeF: the molecular weight of none of these compounds has been determined.

CEROUS BROMIDE. Ce.Br. or CeBr. Known have been described, but their existence is very as the hydrate CeBr. xH.O.: Obtained by dissolving cerous oxide, Ce. a. in HBrAq and evaporating. Deliquescent needles; anchanged by heating out of contact with air; heated in air, partially decomposed with less of Br. Forms double salt CeBr₃.AuBr₃.8H₂O (Jolin, Bl. [2] 21,

CEROUS CHLORIDA. Ce.Cl. or CeCl. Obtained by heating Ce in Cl; or by dissolving Ce₂O₃ in HClAq, adding NH,Cl, evaporating to dryness, and driving off NILCI by heating; or by passing a mixture of dry 40 and Cl over hot Ce₂O₃ (Didier, C. R. 101, 882). Here CeCl₃ was prepared by Robinson (Pr. 37, 150) by heating pure Ce₂(C₂O₄) in pure dry HCl gas to 120 -1303 for some time, then to 200°, and then to low redness; the small quantity of C separated was removed by heating at low redness in mixed CO₂ and HCl; finally the temperature was raised to a full red heat and the CO₂ stream was stopped. The chiefide was allowed to cool in HCl gas, transferred to a small tlask, and kept in acuo, over H.SO, and surrounded by CaO, until all HCl was removed. S.G. $\frac{15747}{11697}$ 3:88 (Robinson, l.c.). A mite, deliquescent solid; easily soluble in water with production of heat; decomposed by O, or by steam, to Ce.O. (Didier). A hydrate, CeCl₃.7IL₂O, is said to be formed by digesting Ce.O3 in HClAq and evaporating. salts are described; Various double CeCl₃·4HgCl...10H₂O, CcCl₃·PtCl₄·13H₂O, CeCl₃·AuCl₃·13H₂O (Jolin, l.c.).

CEROUS CYANIDE (and double cyanides) v.

CYANIDES.

CEROUS FLUORIDE. CeF. ⊖btained as a gelatinous pp. (2CeFaH_O) by adding NaFAq to

CeCl, in IIClAq (Jolin).

CERTE PLUOMDE. CeF, II. . An amorphous brown powder, insoluble in water, obtained by treating CeO2.6H.O with HFAq, washing, and drying at 100°. Decomposed by heat with loss of H₂O and HF and formation of CeF₃; heated strongly in contact with moist air CcO, and HF are formed. Combines with KF toy treatment with KF.HF) to form 2CeF, 3KF.2H.O (Brauner, C. J. 41, 69).

CEROUS IODIDE. Ce.I. or CeI. Obtained as the hydrate Cel₃,911.0, in colourless crystals, by dissolving Ce₄O₃ in HIAq, evaporating in a current of H.S, and placing in vacuo over H.SO.: soluble in water and alcohol, very quickly decomposes in air (Lange, J. pr. 82,

134).

 \mathbf{T} lie CERIUM OXYCHLORIDES. eompound Ce₄O₃Cl₆ (= Cc₂O₃·2CeCl₃) is said to be formed when CeCl₃ is heated with Na (in preparation of Ce) and the mass is treated with water. Dark purple, lustrous, powder; insoluble, in water (Wöhler, A. 144, 254). The same oxychloride is said to be obtained, as iridescent scales, by passing a mixture of steam and N over a fused mixture of CeCl, and NaCl; easily soluble in dilute acids; heated in air gives Ce.O. and 11Cl (Didier, C. R. 101, 882).

Cerium, hydrexides of, v. CERIUM, OXIDES OF. Cerium, oxides and hydrated oxides of. The best studied oxides are cerous oxide Ce.O. and cerio oxide CeO,; a peroxide CeO, also probably exists. Other oxides, e.g. Ce,O, and Ce,O,

CEROUS OXIDE. Ce.O., (Cerium sesquioxide.) Mol. w. unknown. The white, bulky pp. obtained by adding KOHAq to a solution of a cerous salt is hydrated cerons oxide; as the pp. at once begins to take up O and CO, from the air the hydrate has not yet been obtained pure. The oxide $Ce_{\cdot}O_{3}$ is prepared by heating cerous exalate in a stream of pure H. The explate is prepared from the basic nitrate (v. Chrium, Preparation) by dissolving in a little HNO, Aq ana ppg. by oxalic acid (Popp, A. 131, 361; Rammelsberg, B, 6, 85).

Properties and Reacti no. A grey solid; nuchanged by heating in II. Dissolves in many acids to form cerous salts of the form Ce_nX_{so} X SO₀, CO₀, 2NO₀, 2ClO₀, \(\delta c \).

CERIC OXIDE. CeO, (Cerium dioxide). The pale yellow pp. obtained by suspending Co₂O_{3,2} H₂O i • KOHAq and passing in CI is hydrated certe oxide (2CoO_3H_O; Rammelsberg, ₽. 103, 40).

Formation.-I. By washing the hydrate with water containing a little acetic acid until KOM is removed, drying and heating .-- 2. By heating cerous sulphate Ce. 3SO, to full redness in air. 3. By heating CeF, in air (Branner).

• Preparation.—Corous oxalate is prepared from the basic nitrate from cerite (v. Centum, Preparation); it is heated to reduces in a Pt dish with free access of air. Nordenskield (P. 111, 616) obtained colourless, transparent, 'esseral crystais of CeO, by heating CeO, for 24 hours with a little borax in a porcelain oven, and treating the mass with 11ClAq: S.G. at $15^{\circ} = 6.91.$

Properties and Reactions. - Very pale yellow Properties and Rederious.—Very P.ue yellow Solid (Wolf. Framer, Robinson [C. N. 54], Grookes, P. 38, 414). S.G. 674 (Nilson a. Pettersson, /. 15, 1159). S.V.S. 2545. S.H. 0877 (N. a. P., Pr. 31, 46). Dissolves in conc. H.SO, Aq with production of much heat, and evolution of some O; on crystallising, the salt Co.3SO Cc(SO4)...21112O separates (Mendelejeff, A. 168, 45); from the mother-liquor of this salt cenic sulphate Ce2SO, 4H,O is obtained. This reaction shows that part of the CeO, is reduced by the H2SO, to Ce O3, and part reacts with the acid to form Cc2SO, Dissolves in HNO, A1; on adding NII, NO, and orystallising in vacan the double salt 2Ce(NO₃), 4NH,NO₃,3H₂O is obtained. Scarcely soluble in HGlAq; but dissolves in this acid, and also in other dilute acids, in presence of ceducing bodies, ey. filter paper, alcohol, SO Aq, &c.

CERIUM TLEDRIDE. CeO. (Cerium peroxide.) Said to be obta aed as a reddish pp. by adding a slight excess of NII3Aq to Ce.3SO,Aq, and digesting with hydrogen peroxide (de Boisbaudran, C. R. 100, 605; Cleve, Bl. [2] 43, 53).

OTHER OXIDES OF CERTUM are described by Popp (A. 131, 36); Lermann (J. pr. 30, 184; 82, 385; 92, 113); the formulæ Ce O, and Ce O, are assigned to these exides, respectively; but experiments made by Rammelsberg (P. 108, 40) and others tend to show that the only exides which have been isolated are Ce,O,, CcO, and

Cerium, exychlerides of, v. under CERIUM. HALOID AND OXYHALOID COMPOUNDS OF.

II of acids by Cs. Two series of salts are known, cerous salts represented by Cs. 3SO4, and ceric salts represented by Ce2SO... The cerous salts salts represented by Ce2SO4. The cerous salts correspond to the oxido Ce2O4, the general form of these salts is Cs.X, where X SO, CO, 2NO, 2ClO, &c.; the eeric salts correspond to the oxids CeO_2 , their general form is CeX_2 where $X = SO_4$ &c. The cerons salts are considerably more stable than the ceric salts; the latter are readily reduced to the former; but few ceric salts have been obtained, the principal salt is Ce2SO4; several double salts are known of the form Ce2SO, xM SC, and Ce1NJ, xMNO, when M is an alkali metal. Many double cerous salts are known. A few basic salts are also known. Some salts have been isolated which appear to belong to the mixed form rCe₂X₂yCeX₂; and one of the so-called basic nitrates is probably g Ce26NO gy CeO2. For descriptions of the individual salts v. the articles on the various groups of salts, carnonates, sulphates, &c.; the chief salts are the bromate, carbonates, iodate, nitrates, oxalate, perchlorate, phosphates, selenate, sulphates, tungstate.

Cerium silicids. Described by Ullik (Z. [2] 2, 60) as a brown powder; obtained by passing an electric current from 8 Bunsen-cells through a fused mixture of KF and Co.F. in a porcelain crucible, and treating with water the brownish mass formed at the negative pole. The Si was, dsrived from the crucible which was strongly attacked. Analysis gave numbers nearly agree-

ing with the formula Ce.Si₃.

Csrium, sulphides of. Only one sulphide is known, cerous sulphide, Ce.S. It is best obtained by passing dry 11.S over CeO, heated to full redness in a porcelain tube. S.G. 5.1. Vermilion to black according to the temperature at which prepared. May also be obtained in over a fused mixture of dry NaCl and dry Ce Cl, and then washing with water. Unchanged in ordinary air, but burnt to SO, and CeO, below a red heat in air. Dissolves easily in dilute acids with production of H.S; very slowly decomposed by warm water (Didier, C. R. 100, 1461; v. also Lange, J. pr. 82, 129; and Mosander, P. 11, 406).

Cerium, sulphecyanids of; v. sulpho-OYANIDES, under CYANIDES. M. M. P. M.

CEROPIC ACID CaslI ... O. (?) In the leaves of the Scotch fir (Pinus sylvestris), from which it may be extracted by dilute alcohol. Minute

needlen—Bah" aq (Kawaliera d. 88 a 360).

CEROSIN C₂₄II aO. [82]. A waxy substance found on the stem of the sugar-cane. Nacreous lamine (from alcohol). Heating with potash-lime oxidises it to cerosic acid C48H98O2 (?) [93c], which may be crystallised from petrolsum (Avequin, A. Ch. 75, 218; Dumas, A. Ch. 75, 222; Lewy, A. Ch. [3] 13, 438).

CEROTENE C. H., [58] I. A. F. oduct of

the distillation of Chiness wax (Brodie, P. M. [3] 88, 878; A. 67, 210). Resembles paraffin. Chlorine forms $C_{x:}H_{x_0}^*Cl_{1_0}$, $C_{x:}H_{x_0}Cl_{2_1}$, and $C_{x:}H_{x_0}Cl_{2_2}$. A similar body [66°] occurs in hay; it is perhaps $C_{x_0}\Pi_{x_0}$ (König a. Kiesoff, B. 6, 500). 500).

CEROTIC ACID C., H., O. or C., H., O. [78°]. Occurrence. -1. As ceryl esrotats in Chinese

Cerium, salts of .- Salts obtained by replacing | wax; whence it is obtained by distillation, er, better, by treatment with alcoholic KOH .- 2. In the free stats in bte's wax (John, Ottenische Schriften, 4, 88; Boudet a. Boisstnot, J. Ph. 13, 88; Httling, A. 2, 267; Hsss, A. 27, 3; Gerhardt, Rev. scient. 19, 5; Lewy, A. Ch. [3] 13, 438; Brodie, A. 67, 180; Zatzek, M. 3, 677).

Formation .- By oxidation of paraflin with dilute HNO₃ og with chromio mixtero (Gill a. Meusel, C. J. 21, 466). Also by heating ceryl alcohol with soda-lime (Schwälb, A. 235, 141).

Preparation .- The alcoholio extract from bee's wax is recrystallised till it melts at 70°. This is dissolved in alcohol and the lead salt thrown down by alcoholic lead or cubric aco-

Salts. — NaA'. — CuA', at 100°. — PbA', [1f3°].—KA'. —MgA',? [140°-145°].—AgA'. Methyl ether MeA'. [60°] (Nafzger, A. 224

Ethyl other EtA'. [60°]. Fatty plater (from alcohol). Soluble in ether. May be distilled in vacuo. On distillation it gives of C.H. and CO. and the distillate contains cerotic acid and a paratin [44°] (C₂₆H₃₄ or C₂₇H₃₆) while in the retort there remains a ketone [920] which

is (C., H.,), CO or (C., H.,), CO.

Cory l ether C., H., O.C., H., O.C., H., O.

Occurs almost pure as Chinese wax (Brodis) Occurs also in opium-wax (Hesse, B. 3, 638) and in yolk, the sweat of sheep (Buisine, Bt. [2

42, 201).

An acid [79°] isomeric or identical with ecrotic acid is the chief acid present in the product of saponification of carnauba wax. From alcohol it separates as a jelly, but from other solvents (ether, benzene, light petroleum) as a crystalline powder (Stürcke, A. 223, 283; cf.

Bérard, Z. [2] 6, 4[5].

Salt.—PbA'₂ [112²]. Sol. boiling glacial acctic acid and toluene. Insol. boiling alcohol

or ether

CEROTONE C 3H 100 i.e. (C H 110) 2CO (?) [62°] Formed by distilling lead cerotate (Brückner, J. pr. 57, 17). Plates (from ether).

Ceretene (C₂₀H₂₁).CO (?) [92°]. Formed by distilling ceretic acid or its ettyl other (Nafzger,

A. 221, 237). Plates (from acetone).

CERYL ALCOHOL'C₂₇II₅₆O or C₂₆H₅₂O₂. [79°]. Occurrence.—Chinese-wax consists almost entirely of ceryl cerotate (Brodie, A. 67, 180; Schwalb, A. 235, 141). Ceryl csrotate also occurs in the sweat on the wool of sheep. The wax outside ripe heads of the opium poppy contains ceryl cerotate and ceryl palmitats, [79°] (Hesse, B. 3, 637).

Preparation.—Chinese-wax is saponified with alcoholic potash, the product mixed with baric chloride solution and the ceryl alcohol separated from baric corotate by solution in alcohol.

Reactions. -- Is oxidised to corotic acid

C₂,H₃,O₂ by heating with soda-lims.

Iso-veryl alcohol C₂,H₃₆O. [62°]. The portions of the wax of Ficus gummiflua which is

insoluble in cold ether (Kessel, B. 11, 2113).

**dcctyl delivative C.H.,OAc. [57°].

CESPITINE C.H.,N(?). (95°). Occurs in coal ter, and in the product of the distillation of psat (Church a. Owen, P. M. [4] 20, 110; Fritzsche, J. 1868, 402), Liquid, misoible with water. Combines with Eti. Its platinochloride is decomposed by boiling water (De Coninck, Bl. [2] 45, 131). Goldschmidt a. Constam (B.16, 2978)

45, 181). Goldschmidt a. Constam (B. 16, 2978) suggest that it is wet pyridine.

CETAND v. Hexadecand.

CETENE O₁₈H₃₂ i.e. CH₂.(CH₂). CH:CH₂.

Cetylene. Hexadecylene. [4°]. (275°); (155°) at 15 mm. S.G. ½ 7917; ½ 7842. V.D. 8°0.

Formed by distilling cetyl alcohol with P₂O₃.

(Dumas a. Péligot, A. Ch. [2] 62, 4; Smith, A. Ch. [3] 6, 40). Also by distilling cetyl pulmitate or commercial. Dermaceti (cf. Krafit, h. tate or commercial permaeeti (cf. Kralit, B. 16, 3018). Oil; sol. alcohol and ether. According to Berthelot (A. Ch. [3] 51, 81) it forms unstable compounds with HBr and I.Cl. It forms a dibromide G18H22Br2 [14°] whence alcoholic KOH forms oily bromo-cetene (Chydenius, .1. 143, 267). HClO forms chloro-cetyl alecticol C₁₄H₃₂Cl(OH) (c. 300°) (Carius, A. 126, 155). SO₃ forms C₁H₃₁SO₃H [18°], insol. water.—KA': [106°]; S. 1 (Lasarenko, B. 7, 125).

An isomeride of cetene, [42°], 1281°) is got by distilling azelaic acid with baryta (Schorlem-

mer, A. 136, 265).
CETENE GLYCOL. The di-acetyl derivative C1, H12 (OAc), is formed by the action of . AgOAc on cetene dibromide. It cannot be distilled (Chydenius, A. 143, 270).

CETENE OXIDE C16H12O. [below 30]. (below 300°). From chlora-cetyl alcohol iv. CETENE) and aqueous KOH (Carius, A. 126, 202).

Minute needles.

CETIC ACID $C_{15}H_{20}O_{2}$. [55°]. The glyceryl derivative occurs in the oil expressed from the seeds of Jatropha Curcas (Bouis, J. 1851, 462). According to Heintz (P. 90, 137) a very small quantity of this acid is found in the products of saponification of spermaceti

Ethyl ether $\hat{ ext{EtA'}}$, $[21^{\circ}]$ (P.).

CETINENE C₁₀H₃₀, Cetylene, Hexadecinene, Hexadecylidene, [20°], (34°), (160°) at 15 mm, S.G. 2° = 804; 3° = 797, H.F. 118,000 (Beethelot). Large colourless tables. Formed by heating ceteno (hexadecylene) bromide (natural or synthetical) with alcoholic KOII (Krafit, B. 17, 1373; cf. Chydenius, C. R. 61, 180).

CETRARIC ACID $C_{18}H_{16}O_{8}$. Contained, together with licherosteario acid, in Iceland moss (Cetraria islandica) (Berzelius, Schw. J. 7, 317; A. Ch. 90, 277; Herherger, A. 21, 137; Knop a. Schnedermann, A. 55, 144). Hair-like needle. (from alcohol), v. sl. sol. water, sl. sol. ether, v. sol. boiling alcohol. Tastes bitter. Decomposed before melting. Its solutions turn brown on boiling, especially in presence of alkali. Fe₂Cl_d gives a red pp. in neutral solutions.—(NH₄) $_{2}\Lambda''$.-PbA".

picetyl C₃₂H₆₆ Dotracontane. [70°]. (above 360°). V.D. 15.8. Formed by treating an ethereal solution of octyl iodide with sodium (Scrabji, C. J. 47, 37; cf. Lebedeff, J. R. 1884, [2] 299). Scales, v. sl. sol. ether, v. sol. boiling glacial HOAc.

CETYL ACETATE C16H33OAc. n.Prim.hexadecyl acetate. [19°] (Becker, A. 102, 220) (82°) (Krafft, B. 16, 1721). (276°) at 190 mm.; (200°) at 15 mm. (K.). S.G. 15 8640; 25 8612. MM. 18·772 at 20·7° (Perkin, C. J. 45, 421). Needles; sl. sol. cold alcohol.

CETYL - ACETIO ACID is identical with Brearic scid (q. v.).

Dl-cetyl-acetic acid C₁,H₂,O₂ s.e. (C₁₀H₂₃)₂CH.CO₂H. [70°]. Formed by heating di-cetyl-malonic acid at 150° (Guthzeit, A. 206, 365). Crystalline scales, sl. sol. alcohol.-AgA': amorphous pp

CETYL ALCOHOL C14H34O. Asthal. n-Primheradecyl alcohols Mol. w. 242. [509]. (3449); (190°) at 15 mm. S.G. (liquid) \$2 -8176; \$4 -8105; \$4 -7837. H.F. 112,000 (Bertholot).

Occurrence.—Spermeeti is cotyl palmitate (C₁₀H₂₁)C₁₀H₂₁O₂ (Chevreul, Rechercher sur les & cps gras, p. 171; Dumas a. Péligot, A. Ch. [2] 52, 4; Dumas a. Stas, A. Ch. [2] 73, 124; Smith, A. Ch. [3] 6, 40; A. 42, 217; Heintz, P. 84, 232; 87, 553). In the sebaceous glands of geese and ducks (De Conge, H. 3, 225).

Formation. By the distillation of baric sebacate (Schorlemmer, Fr. 19, 32).

Preparation. -- 1. Spermaceti (10 pts.) is boiled with alcohol (5 pts.) and potash (2 pts.) until saponified. The product is poured into water and the cetyl alcohol crystallised from ether. The crude commercial cetyl alcohol contains in addition to hexadecyl alcohol also n-poctadecyl alcohol and probably small quantities of other homologues (Krafft, B. 17, 1627).-The acctate is formed by reducing palmitic aldehyde (obtained by distilling barium palmitate with barium formate) with zine-dust and Acetic acid (Krafft, B. 16, 1721).

Properties, -Small crystalline plates (from alcohol). Gives palmitic acid on oxidation.

Sodium Cetylate Calla NaO. [110°] (Fridau, A. 83, 1).

LEthyl ether C1611. OEt. [20°] (Becker,

A. 102, 220).
TRI-CETYL-AMINE C₁₅II₁₆N i.e. (C₁₆II₂₀)₅N. [397]. From cetyl iodida and gaseous NH, at

150° (Fridau, A. 83, 25). Needles. Its salts aro insol. water, tol. alcohol and ether. -B'HCl. -(B':ICl) P(Cl.: yellow pp. CETYL-ANILINE C., I, N i.e.

 $N(C_nH_1)(C_{16}H_{33})H$ Placetyl-cetyl-aminc. From colvi lodide and aniline at 100° (Fridau, 11.83, 29). Silvery scales (from alcohol). Has no action on litmus, and does not pp. metallic salts. B', H, PtCl,

Di etyl-aniline N(C,Il,)(C,IH,3)2.

ectyl-aniliue and cetyl iodido at 110° (F.). Crystalline.—B'.H.PtCl., CETYL-BENZENE C.H.,(Cl., H.3). Hexadeeylben snc. [27°]. (230° at 15 mm.) Formed by the action of sodium upon a mixturo of iodebenzene and cetyliodide (Krafft, B. 19, 2983).

CETYL-BENZENE-SULPHONIC ACID C. II. (C. II.). SO. I. H. radccyt-benzene-sulphenic acid. Formed by sulphonating cetyt-benzene. The sodium saft A'Na is sparingly soluble (Krafit, B. 19, 2983)

CETYL BENZOATE CieHzOBz. [30°] (Becker. A. 102, 219). Crystalline scales. CETYL BORATE C1611 280.

[58°] (Schiff,

A. Suppl. 5, 198).

CETYL BR MIDE C. H. Br. [15°]. From cetyl alcohol, Br, and P (Fridau, A. 83, 15). Insel. water, v. sol. alcohol and ether.

CETYL CHLORIDE C14 Handl. S.G. 12 .841. From cetyl alcohol and PCl₃. OR, insol. alcohol, sol. ether. Boils above 280° with decompo sition (Dumas a. Péligot, A. Ch. 69, 4; Tüttscheff, Rép. Chim. pure, 2, 463).

CHEMICAL AND PHYSICAL PROPERTIES BODIES, CONNEXIONS BETWEEN. lertain properties are common to all kinds of natter, others are characteristic of this or that ind only. Thus, every material substance is cted on by the fore of gravity in exactly the ame manner, but only a few liquids rotate he plane of polarisation of a ray of light. roperties belonging to the second of these lasses are subdivided into two groups, physic al and chemical properties. Chemistry deals with those changes in the properties of material odies which are accompanied by changes in ho composition of the bodies. Physics deals ith changes in the properties of bodies the emposition of which remains the same. When he totality of properties by which a body is nown remains unaltered throughout any pro-ess, that process is called physical; when he result of any precess is a body or bodies vitle properties so different from the totality d those of the original that the diginal can to longer be said to exist, that process is alled chemical. Physical and chemical proesses are always closely connected in their ccurrence; no chemical change takes place vithout some accompanying physical change, and it is probable that every physical change is o some extent accompanied by chemical hange. Many physical properties are quantiies which may be accurately measured; e.g. nelting- and boiling-points, specific gravity, &c., c. Change of composition of a body or sysem of bodies is very frequently accompanied by shango in the value of one or more of these neasurable quantities; in other words, the physical constants of a hody are conditioned, mong other circumstances, by changes in the variable, chemical composition. By the chemieal composition of any homogeneous kind of natter is meant, in the first place, a statement of ho elements, and of the mass of each element, n a given mass of that body; in this meaning of the term the chemical composition of a body, or system can be accurately stated, and lefinite relations can be determined between changes in the composition and changes in the values of such physical properties as meltingand boiling-points, specific rotatory power, spesific refractive energy, and so forth. When the relations between the two groups of changes; have been studied and generalised, it may become possible to infer the amount and chaacter of a change of composition from measurenents of the changes in the values of a few hysical properties. It would be impessible to study the relations between every chemical shange and the accompanying variations in the physical properties of the bodies forming the lianging system; it is necessary to select ypical eases, and to study there as accurately ind minutely as possible. As a rule, one physial property is chosen for measurement; the composition of the system is defined to begin with; the system is allowed, or caused, to pass into another definite state; and the variation in the value of the chosen property is determined.

But when at is found that several distinct kinds of matter exist, each homogeneous, each listinguished by definite properties, and each sontaining in a given mass the same masses of

the same elements, it becomes necessary 'to widen the meaning of the expression chemical composition. It becomes necessary to frame an hypothesis to account for the observed facts The hypothesis generally adopted asserts that matter has a grained structure, that a mass of any kind of homogeneous matter is composed of a vast but not indefinite number of minute parts; and that the properties of the mass are conditioned by the properties of these parts. These minute portions of matter are called molecules. But the molecule is not necessarily itself without parts. The chemist asserts that overy molecule is built up of a definite number of smaller parts either of one or of several kinds of matter. These parts of molecules are called atoms. The atoms of elements are the ultimate forms of matter with which chemistry at present concerns itself. The hypothesis goes on to assert that the properties of a molecule, and hence the properties of any portion of homogeneous matter composed of molecules of this kind, are conditioned by the nature, the number, and the relative arrangement, of the atoms which together form the In other words, the hypothesis molecule. declares that the molecule is itself a structure. Ou this hypothesis, by the chemical composition of a body is meant a statement of the nature, number, and relative arrangement, of the atoms which form a molecule of the body. know as yet almost nothing about the configuration of atoms in molecules; but chemistry has formed certain more or less clear hypotheses, and attempts are constantly being made to councet changes in the values of various physical properties with variations in the relative arrangement of atoms in molecules, as this arrangement is conceived by the .hy. potheses in question?

But the physical conception of the molecule is derived from the study of various gaseous phenomena: the physicist deals with the molecule as a whole; ho pictures the melecules as performing certain vibratious, on the form, amplitude, and rate, of which the physical properties of bodies depend. The two conceptions, the chemical and the physical conception, of the molecule are therefore to a great extent mutually independent. In how far then, one may ask, can a devolopment of the chemical conception bo looked for hy using physical methods of inquiry? Leoking at recent researches, it seems probable that the ohemical conception of the molecule must be very considerably modified, and must be brought more into harmony with the physical conception. The latter is itself to some extent being changed by the development of the theory of evortex atoms. But it must not be forgotten that the physical conception, in so far as it is a clear conception, has been developed almost wholly from the study of gaseous laws, more especially of the laws which express the relations of the volumes of gases to temperature and pressnre; these relations are dependent on the states of combination of the parts of molecules, and are in no way affected by the nature or number of. these parts. The chemical conception, on the other hand, must be made sufficiently elastic to cover the phenomena presented by gaseous,

therid, and solid, compounds; and most of the as AB+CD=AD+BC, which groups may be of shemical processes which occur among com-pounds belonging to these classes are conditioned both by the nature and number, and by the states of combination of the alms which form the chemical molecules of the reacting bodies. The chemical conception of the molecule will probably be modified when we know more of those properties which like the relation between the volumes of gases and the temperature and pressure of diese gases, are to a great extent, if not altogether, independent of the nature and numbers of the constituent parts of molecules. The physical conception will pro-bably be modified as we learn more of those properties which, like specific heat, are for the most part dependent on the nature and numbers of the constituent parts of molecules. (v. Molecular structure of matter, theories

Chemistry regards not only changes in the composition, but also blanges in the properties, of bodies; sho attempts to generalise not only the laws of composition, but also those of the mutual actions, of bodies. The study of the connexions between changes of composition and variations in physical properties of chemically reacting bodies will throw light on the nature of chemical change. When accurate measurements have been made of the quantities of heat which disappear or are produced in a series of typical chemical processes we shall be able to apply to these processes the knowledge of heat-energy which is generalised in the thermodynamical laws. Chemical change may then perhaps be shown to be a special instance of the working of these laws. The conditions of chemical change on the one hand, and of physical change on the other, must be studied, in order that the laws which express these conditions may be gained; the relations between these two groups of laws must then be ascertained; thus it may become possible to attain to clear mental images of netural phenomena as wholes which now present one aspect to the physicist and another to the chemist.

For accounts of the various physical methods employed in chemistry, and résumés of the more important results, v. Physical methods.

M. M. P. M. CHEMICAL CHANGE. Chemical science is based upon the hypothesis that matter is constituted of extremely small particles or atoms, and that these atoms are capable of aggregating together by virtue of certain inherent properties or forces, their affinities, to form complex atomic structures or groupings.

The recognition of this distinctive force by the older chemists led them to propound various theories to account for its existence and explain the phenomena of chemical action (v. AFPINITY).

Wish the naturo of this force we are not concerned here, but only with the phenomena that accompany its exhibition and the circumstances that modify its action.

By a chemical change, therefore, is meant any alteration either (1) of the character of a permutation in an atomic group, such for instance as is exhibited in the change of ammonium cyanate, NH, CNO, into usea (NH2), CO; or (2) a permutation between two or more such groups,

various degrees of complexity; or (3) the change may arise from a combination such as AB + CD = ABCD, or the converse of this, as is

seen in the phenomena of dissociation.

The majority of chemical changes may be formulated as permutations between two sots of atomic groups; such as the action of bases on acids, the decomposition of one salt by another, of the combination between gascous olements as H₂ + Cl₂ = 2HCl. Examples of chemical change according to case (3) are of less frequent occurrence than the last; such are the fermation of double salts like the alums the combination of certain gaseous molecules with oxides, &c., as CaO + CO₂ CaCO₃ and CO Cl₂ = COCl₂. The number of strikingly marked instances that could come under the head of permutations in an atomic group is very small, but such changes may be of frequent occurrence, producing alterations in physical and chemical properties too slight to be Reognisable.

There are several bodies which are known to undergo remarkable and highly interesting physical, and consequently no doubt chemical, changes, when heated, but whether such changes come under class (1) or class (3) is undecided. Among such substances are phosphorus, paraevanogen, and evannric acid. In the case of phosphorus, the change from the yellow to the red modification, caused by heat or light, is probably due to an alteration in the state of aggregation of the atoms; that is to say, if the molecule of vellow phosphorns he P, that of the res modification is probably Pm; for solid paracyanogen, which is converted by heat into gaseous cyanogen, and for cyanurio acid, the same may be true, with or without a re-arrangement among the constituent atoms of the molecules of Almotropy and Isomenism).

The study of the phenomena attending a chemical change shows that in many instances there is an accompanying evolution of energy, from the changing system, in one form or other, either as heat, or light, or as electrical currents. In other cases to produce a chemical change expenditure of energy is necessary.

Wast may be the nature of the chemical force or affinity that acts between atoms is not known, but it is characterised from gravitative force by this difference, that whereas gravitation act upon all kinds of matter alike, depending merely on the masses of the bodies, chemical attraction or affinity depends upon the kinds of matter that are precented to each other, as well as upon the conditions under which the bodies are brought together; in other words, it is an. elective attraction modifiable by circumstances. For instance, at a red heat metallioiron is oxidised by water vapour and hydrogen is set at liberty, but at a lower temperature oxide of iron is reduced by hydrogen with the formation of metallic iron and water vapour; a mixture of hydrogen and chlorine will femain unchanged for any length of time in dakness, but exposure to sunlight will cause almost instantaneous combination, and the resulting compound (hydric chloride) may be again converted into its original constituents by heat. If to a solution of silver nitrate a pieceof metallic copper be added, metallic silver is, ppd., and coppernitrate formed; and if now to the copper nitrate a piece of zine or iron be added, of an extrancous body which itself undergoes metallic copper is ppd., and zine or iron nitrate no apparent change. For instance, oxygen and is formed.

These illustrations are sufficient to show the relativity of chemical affinity as depending both upon the conditions o which the system is subjected as well as upon the qualities of the materials.

Under whatever conditions a elemical system may exist in which a change is happening, the atomic forces at work will centime to act until a state of more or less stable equilibrium is reached, after which no further action will take pace; and the ultimate limit reached will depend upon (i) the relative quantities of the reacting bodies; and (2) the conditions to which the system is subjected. As the system passes from the initial to the final configuration there will be a los or gain of entry equal in amount to the difference between the total energy of the system in the two states. The rate at which the change takes place will depend also upon the same two circumstances. These two statements amount to this; that, representing a chemical change by the equation A + B: A' + B', all the atomic forces at work producing the transformation have not invariably the same zatios, but that the ratios vary with variations in the conditions as regards heat, light, &c.; and, consequently, any determinations of the relative affinities of the members of the system can only be looked upon as expressing certain ratios that hold good under special conditions. The final stato reached by the system, and the rate at which the change progresses towards that state, vary with the relative masses of the reacting bodies, other things being equal, although the atomic forces or the affinities wmain the same. In other words the final configuration, and tho speed of attaining it, are each a function of the reacting masses and of the atomic forces, the latter being a function of the physical conditions to which the system is subjected.

The phenomena of dissociation furnish many examples of othese facts; as do also those systems which are limited by inverse actions and do not properly come under the term dissociation, wherein both the masses of the constituents as well as the conditions, especially as regards heat, influence the change in its amount and rate. The great field offered for investigation by fractional pptn. will, no doubt, afford many striking instances of the variations of the affinity values under diverse circumstances when the

subject is worked out.

From the foregoing considerations it is clear that a chemical system may or may not undergo ohango by virtue of any intrinsic forces acting among the constituents, but that such will happen only according to the conditions to which it is subjected. Reactions which at moderate temperatures take place with evolution of much energy may be completely suspended by lowering or increasing the temperature, tachuding hight, or altering the prossure; in other words, the forces or affinities resisting change, if greater than those tending to thoduce an alteration under some circumstances, may be reversed when these circumstances are altered.

There are, however, a number of interesting examples in which the stability of a system seems to be overturned by the mere presence of an extraneous body which itself undergoes no apparent change. For instance, exygen and SO₂ do not combine when moderately heated, but if passed over spangy platinum combination readily occurs. Sometimes again the intermediate body does undergo change, as when chlorine is passed over a strongly heated mixture of carbon and silica, whereas without the addition of carbon the silica is not acted upon by the chlorine; or, platinum, which itself is insoluble in nitric acid, may be rendered soluble in the same acid by alloying it with silver (v. post, Catalytic changes, p. 750).

Some equally remarkable instauces of the apparent necessity of the presence of a third body in order to bring about chemical action between two others have been noticed. Wanklyn (C. N. 20, 271) found that perfectly dry chlorine gas has no action upon metallic sodium. Couper (C. J. 43, 153), starting from Wanklyn's observation, has examined the action of dry chlorine on several metals that are acted upon vigorously by the moist gas. He found that dry chlorino has no perceptible action on Dutch metal, whereas with the moist gas combination takes place, with production of heat and light; or on touching . the metallic surface when in an atmosphere of dry chlorine with a drop of water, instant combination occurs. Couper examined a number of metals in the same way with the following results: the chlorino gas used was allowed to stand over CaCl, for several days to thoroughly dry it. Zine, and magnesium, showed no action; silver, slight action; bismuth, combination slow; arsenic, antimony, and tin, rapidly acted upon. It is worthy of note that these last three metals form volatile chlorides liquid at ordinary temperatures. With mercury, combination equally rapid, with dry or moist chlorine. Potassium showed slight action, probably due to adhoring KHO; with proper precautions against moisture, action was slow. Dixon (T. 1884, 617) has observed a somewhat analogony fact relating to the combination of gases under the influence of the electric spark. He has shown that if a mixture of CO and O be perfectly dried by P2Os, and be then subjected to the spark from a largo Leyden jar or a Ruhukorif's coil, union does not take place; if, However, the slightest trace of maisture be admitted to the mixture, and the spark again made to pass, combination occurs. The hypothesis Dixon advances to account for these facts is that the intervention of water molecules is necessary to bring about combination, a molecule of water being decomposed under the influence of the spark by one of carbonic exide to form earbonic acid and free hydrogen, the latter in its turn combining with the oxygen to form water: this cycle of operations being represented by the equations $H_2O + CO = H_2 + CO_2$; $H_2 + O = H_2O$; consequently a comparatively small number of water molecules are necessary to effect complete combustion. (See also C. J. 49, 94.) Phosphorust and carbon have been shows by Baker (C. J. 47, 349) to combine with oxygen less energetically in the absence of moisture than when moisture is present; and Ramsay and Young (C. J. 45, 93) observed that if a mixture of dry If and N is passed through a tube containing iron filings at a red heat no

Several of the elementary bodies are known to exist in two or more different modifications, such for instance as sulphur, scionion, carbon, phosphorus, and oxygena the several forms of each element exhibit more or less strongly marked differences inchemical as well as physical properties. It is probable that such different modifications of one elementary body consist, as in the case of oxygen and ozone, of different atomic groupings or aggregates of atoms. means by which the change from one modification of an element to another is brought about are various. Oxygen is converted into ozone by the electric spark or 'silent discharge,' and ozone is changed again into oxygen by heat; yellow phosphorus is converted into the red modification eithor by light or by heat, and the red modification is again reconverted into yellow phosphorus at a higher temperature; sulphur and sclenion undergo several changes under the influence of heat; in the case of carbon, the conditions ne-cessary to bring about metamorphoses are not fully known.

The study of certain isomeric compound bodies (v. Isomer sm) has shown that the transformation of one isomeride into another is, in some cases, somewhat analogous to the phenomena of dissociation. If solid paracyanogen (CN), is heated in a closel vessel to 860° it is entirely converted into cyanogen gas (CN)2; the pressure increases until the gas condenses and is liquefied on the cooler parts of the apparatus. At temperatures below 500° little or no decomposition occurs. As the paracyanogen is heated above this temperature a slow transformation takes place into gasecus yanogen, and the transformation continues until the pressure of the cyanogen gas attains a certsin definite limit beyond which it does not rise, and there is no further evolution of gas. Exhausting the apparatus and maintaining the temperaturo, the pressure again rises to its previous limit and remains stationary however long the heating is continued. For every such temperature there is a maximum pressure reached which limits the further decomposition of the paracyanoger into gaseous eyanogen. If now when the pressure has attained its limit, at a given temperature, a quantity of cyanogen gas is forced into the apparatus, the pressure slowly falls to the initial limit with the transformation of gaseous cyanogen into solid paraeyanogen. Troost a. Hautefeuille (C. R. 66, 735, 795) have found the following values for these pressures of transformation at different temperatures :--

•				
Temp.	Pressure of tran	sform	atio	r,
502°		mm		
506	56	"	•	
55 9	123	,,		_
575	129			
587	157	"		
59 9	275	",	•	
601	318	.,,,*		
62 9	868	"	•	•
640	1 310	**		

ever, a trace of NH, is obtained.

The transformation of some paracous varieties of NH, is obtained.

The transformation of some paracous varieties of the volatilisation of a liquid in presence of the volatilisation of red phosphorus from the yellow material or vice-versa is a more complex process. If a quantity of yellow phosphorus is heated in a closed vessel (say to 500°), the mass of phosphorus being more than sufficient to volatilise in the space, a maxinum pressure is quickly attained. Aftern time the pressure gradually falls, more or lesequickly according to the temperature, till it reaches a minimum at which it remains constant. Provided there is no change of temperature, the vapour of the phosphorus is gradually converted into the red modification which condenses on the sides of the apparatus. If the quantity of phosphorns introduced into the apparatus is just sufficient to volatilise and fill the vessel with vapour at the first pressure (the heating being continued), red phosphorus begins to form after a time, and the pressure continues to fall until the minimum limit is reached as before. If, however, only sufficient ordinary phosphorus is used to till the apparatus with vapour at the lower limit of pressure, no red phosphorus is formed, however long the heating may be continued. These two pressures the maximum 18 first attained, and the final minimum limiting the transformation of yellow into red phosphorus—depend solely upon the temperature. Troost and Hantefenille (A. Ch. [5] 2, 156) found the following numbers relating to these phenomena:

Temp.	Pressure of vapour of Plimiting the trans- gtormation	
360°	·12 atms.	3.2 atnis.
440	1.75 ,,	7.5 ,
487	6.80	-
494		18 ,,
503	*	21.9 "
510	10.8	1
511	1 -	26.2
531	16 ,,	— .s
550	31 ,,	
177	56 ,,	_

The rates at which the transformation takes place as well as other phenomena exhibited du ing the change have been studied by Lemoine (A. Ch. [4] 24, 194). He gives the following numbers illustrative of the progress of the change in time:

							- 1	-
1		410)°, af 1	er				
mins.	ą h.	2h.	8h,	17h.	24lı.	32h,	4. %ì.	
•		_	2.9	_		-	_	
-				_	-	4.9	47	
• -	11:1	7.0	4.4			_	=	
	-	¥-1	4.0	3.7	3.6		-	
	mins.	mins. 3 h.	mins. \$ h. 2h. 2h. - - - - - - - - -	mins. \$\frac{1}{2}\text{h}. 2\text{h}. 8\text{h}. \frac{2}{5}\text{h}. 2\text{h}. \frac{2}{5}\text{h}. \frac{2}{5}h	mins. 2 h. 2h. 8h. 17h.	mins. \$\frac{1}{2}\text{h} \ 2\text{h} \ 17\text{h} \ 24\text{h} \ 17\text{h} \ 24\text{h} \ 17\text{h} \ 24\text{h} \ 17\text{h} \ 24\text{h} \ 17\text{h} \ 25\text{h} \ 17\text{h} \ 25\text{h} \ 17\text{h} \ 24\text{h} \ 25\text{h} \ 17\text{h} \ 24\text{h} \ 25\text{h} \ 17\text{h} \ 24\text{h} \ 25\text{h} \ 25	mins, 2 h, 2h, 8h, 17h, 24h, 32h,	mins, 2 h, 2h, 8h, 17h, 24h, 32h, 4 h, 2 h,

Lemoine (C. R. 75) 990) has given a mathematical theory of the changes tha red or yellow phosphorus undergoes when heated in a closed vessel, and has compared his formulæ with the results of experiment. Let P be the total mass be the mass of yellow phosphorus formed or existing at time t; if the red phosphorus be supposed to remain in the same state of division throughout, its tree surface will be sensibly proportional to its mass $\mathbf{r} - \mathbf{y}$. The quantity of yellow phosphorus evolved, δy_1 , in time δt is equal $a(P-y) \delta t$, and the quantity of the ordinary phosphorus, δy_2 , transformed into the red modification in the same time is $b(\mathbf{r}-y)\frac{y}{y}\delta t_i$ the total effect is therefore the difference between these two quantities, or

$$dy = \delta y_1 - \delta y_2 = a(\mathbf{P} - y) - b(\mathbf{P} - y) \frac{y}{y}$$

which may be written $\frac{dy}{dt} = f(g-y) \ (h-y)$, representing the rate of change in terms of the ordinary phosphorus existing.

For further account of Allotropic Changes

v. Allothopy.

INFLUENCE OF PRESSURE ON GASEOUS CHANGES.

Many bodies when subjected to the influence of heat in the gaseous state, undergo enarked . . changes either in their physical or chemical properties, or in both; such changes result more particularly in a diminition of molecular density or a disruption of molecular structure. Among such bodies may be eited, mercurous chloride, chloral hydrate, phosphoric chloride, hydriodic acid, nitric peroxide, hydrie sclenide, amylic bromide, and acetic acid. In the case of some of these bodies the changes in question have been proved to be the accompaniment of disruption or dissociation of their molecules (v. Dissociation); in other cases, such as nitric peroxido and acetie acid, there is no complete proof that the changes in density which these bodies suffer when heated in the gaseous stato are really occasioned by a dissociation of their molecules, or are due to the fact of their vapours not oboying the dilatation law even when sufficiently far removed from the liquid state as to place them under the category of gases. Considered from these two points of view, it is evident that the dilatation of a gas under the influence of heat may be of a twofold character, arising from two distinctly separate causes; firstly, tho expansion may be purely physical, varying or not according to the dilatation-law, and secondly, there may be expansion as the accompaniment of a chemical change, viz., separation of the gaseous, molecules into simpler groups of atoms.

"An observed variation of density at different temperatures may be produced by either of theso two causes, or by both combined, and it becomes ficrefore a matter of great importance to be able if possible to discriminate these two actions, and to say to which of them the observed results aro to be ascribed. If it could be shown that a diminution of pressure produced the same variation in the densities of certaily gases as has been observed under the influence of heat, a great point would be gained in favour of the dissociation theory in settling the cases under dispute. It would seem possible that a dis-crimination between the two possible phenomena accounting for abnormal densities might be made by introducing the element of time into such heating, required time to be effected, and flence was

of material introduced into a space v, and let y unvestigations. To make this clear, take the gas nitrogen istroxide, whose vapour density at low temperatures has been found to be approxi-mately represented by the formula N₂O₁, while at high temperatures it corresponds to NO₂ (the vapour densities being 3.18 and 1.59 respectively). Now Troost (C. R. 86, 1394) found the vapour donsity of nitrogen tetroxide at 27° and at low pressures to be as follows:

> Densitz. Pressure. 35 mm. 1.6 $1.59 \text{ (NO}_2 = 1.59).$

These results show that the same change takes place under diminished pressure as occurs under the influence of heat at ordinary atmospherio pressures; that is to say, these numbers indicate that, if the observed changes in density are due to dissociation of the molecules N2O, into the molecules NO_2 , then under a pressure p the ratio of the number of molecules of N_2O_4 to NO_2 is different from the ratio when the pressure is altered to p', temperature being the same in each case. The proof of this assertion is of considerable importance in the theory of dissociation; whether the change in dencity is or is not to be attributed to the supposed fact, that the gas N2O4 forms an exception to the dilatation-law, would seem to be capable of indisputable proof by introducing the clement of time into the experiments. If the gas N.O. is really dissociated into NO₂ under diminished pressure, 2 vols. N₂O₄ would give 4 vols. NO₂; now, by the kinetic theory of gases it is conceivable that this dissociation would take place practically instantaneously when the temperature was in: creased or the pressure was diminished, whereas on reversing tho process the molecules of NO2 would require some time before meeting with the requisite number of partners to re-form the molecules of N.O.. Such an experiment might form a crucial test of the truth of the dissociation-hypothesis for this particular gas, and it might also be applied to other eases, for it is scarcely conceivable that the coefficient of dilatation of a gas should alter in time. Natanson (W. 24, 454) has determined the ratio between the specific heats of nitrogen tetroxide by means of Kundt's dust-figure method; his results Seem to show that as the pressure decreases this gas passes from a more to a less complex molecular structure.

In the determination of the vapour densities of several bodies whose abnormal dilatations are almost undoubtedly to be ascribed to the disruption or dissociation of their molecular structures, the influence of time on the phenomenon has been several times observed, and has been made the subject of investigation by Naumann for the particular case of ammonium carbamate (v. noxt page).

Wurtz (C. R. 60, 728), when determining the vapour density of amylio bromide (B.P. 113°) between 153° and 360°, noticed that when the vapour was suddenly beated to 225° the density was 6 69, whereas in another experiment when the vapour had been maintained at this tempercture, for ten minutes the density was 3.68. Thesoresults show undoubtedly that the diminution in density, or the dissociation produced by

dhe to a chemical change and not to a variation in bringing about dissociation, or disruption, of

in the coefficient of dilatation of the grs.

Naumann (A. 160, 1) studied the influence of time on the dissociation and re-formation of ammonium carbamate; the following tables illustrate the general bearing of his experi-

Speed of dissociation of NH₃), CO, at 46°.

Pressure under the dissociation- pressure, which = 354 mm.	Increase of pressure	Time of increase			
129 mm.s 37 17 10 10 6 4 3 2 5 5 2 1 5 6 6	87 mm. 20 7 4 2 1 .5 .5 .4 1.5	5 min. 5 5 5 5 5 5 5 17			

Speed of formation a combination of 2NH_a + CO₂) at 20°.

Excess of pressure over dissociating- pressure (=824 mm.) at 20°	Decrease of pressure	Time of decrease
185 mm, 140 ", 90 ", 63 ", 45 ", 30 ", 21 ", 15 ", 10 ", 6 0", 4 ", 1 ", 0 ",	45 mm. 50 27 18 15 6 5 4 2 3 1 0	2.5 mins, 5 " 5 " 5 " 5 " 5 " 7 " 7 " 10 " "

In these two examples it is seen that, starting with a mass of solid ammonium carbamate (N. amann showed by his experiments that at all temperatures the gas evolved consists of 2NH, + CO, and suddenly increasing the temperature, a considerable time is required before the normal pressure of dissociation corresponding to that temperature is reached. In like manner, by suddealy diminishing the temperature, the recombination of the ammonia with the carbon dioxido to form the solid (NH₃)₂CO₂ does not take place instantaneously, but a considerable-time clapses before the pressure corresponding to the lower temperature is arrived at. Although analogous in some respects to the volatilisation and condensation of a liquid, the phenomena exhibited by (NH,)2CO, wher heated are characterised by their groater slowness.

In experiments relating to vapours of vary-

ing densities—such as those of Troost on acetic acid and nitric tetroxide at low pressures—it is important to determine whether diminution of pressure acts in a manner similar to that of heat

the molecules of the gas. By introducing the element of time into the experiments, and by suddonly varying the pressure, dissociation might be shown to occur in the case of nitrio tetroxide as already remarked, the process in this case being doubtless reversible; whereas with say, ozone, or a mixture of ozone and oxygen, the amount of change produced by increasing the volume, say, twenty fold, could be determined by the usual methods of analysis (cf. Aso Drs. sociation, and Equilibrium, Chemical).

Considering the three physical states in which bodies are capable of undergoing chemical change, either as gases, liquids, or solids, it is evident that there are two distinct kinds opclasses of chemical systems possible, according to the states in which the active substances exist, and which may be termed heterogeneous and homogeneous systems. The forme? name is applied to all reactions in which the active members of the system are in different physical states, a solid and a liquid, or a solid and a gas, or a liquid and a gas; as examples of each of these may be mentioned the action of acids on metals or on carbonates, the dissociation of calcie carbonate or ammonic case bamate by heat, and the oxidising action of free oxygen on solutions of stannous or ferrous salts. By homogeneous systems are to be understood those in which all the active members exist in the same physical state, either as liquids of gases; it is inconceivable that two solid bodies, however finely powdered and well mixed, could come under this category. Examples of homegeneous systems are shown in the numerous etherification processes, the oxidising action of potassic chlorate on ferrous salts, the action of oxalic acid on potassic permanganate, among li mids, and for gaseous systems, the action of iodine or sclenion on hydrogen, and the influence of light on a raixture of chlorine and hydream or on gascous hydriodic acid. Many valuable facts have been brought out by the study of heterogeneous chemical systems, especially as regards dissociation-phenomena; but the great field in which the most fundamental facts concerning chemical action will be gathered is naturally that endeacing homogeneous systems, for here the most intimate contact exists among the acting substances, affording free play to the various chemical forces at work, and the secondary physical changes which interfere with the primary chemical change are reduced to a mini-

Heterogeneous Systems. 4- Gladstone and Tribe (Pr. 19, 498) have investigated the rate at which a nore positive metal immersed ir, a solution of a sale of a less positive one displaces the latter, and the relation which exists between the rate of action and the mass of salt in the solution. Employing a solution of argentic nitrate, the displacing metal being copper, and allowing the action to continue for ten minutes under varying conditions of concentration, they found that by doubling the amount of silver salt in solution the amount of action that took place during this interval of time was trebled. Zine and cupric chloride, zinc and cupric sulphate, zinc and lead nitrate, iron and cuprio sulphate,

and other combinations, showed in every case, when the solutions were sufficiently diluted, that this 2-3 law holds good. Expressed algebraically, if y be the mass of metal dissolved, and x the concentration of the solution, then the above

2-8 law is $y = ax^{\log x}$ where x is a constant.

These experiments have been repeated and extended by Langley (C. F. 45, 663), who come firms the truth of Gladstone's law; but when the method of experimenting is modified, at by continually moving the metal about in the solution or by brushing its surface so as to keep the solution uniform throughout, Langley conciders that the rate of action is preportional solely to the amount of salt in solution. Moreover, Langley's experiments indicate that the law observed by Gladstone and Tribe arises from two causes, viz., chemical action, and gravitative action, the latter producing slow currents through the solution because of the changing densities of the original salts and of those which are produced in the change.

In studying the rate of evolution of carbon dioxide from marble by the action of acids, Bojuski and Kajander (B. 10, 34) found that The rate of action is proportional to the concentration of the acid, but varies according to the nature of the acid employed; moreover, they concluded that, for the three acids HCl, HBr, and IINO, the speed of the action is inversely proportional to the molecular weights of the acide when the eclutions are of equal degrees of concentration. Pawkewski (B. 13, 334) has continued these experiments, employing different carbonates (BaCO₃, CaCO₃, SrCO₃) with the same acid; although his results are not very regular, yet he considers them sufficient to show that the speed of the reaction is inversely proportional, not to the undecular weights of the earbonates, but to the atomic weights of the metals whose carbonates were comployed.

Of a somewhat similar nature to the experiments of Gladstone and Tribe is the work of Thorpe (C. J. 41, 287) on the behaviour of zine, magnesium, and iron, as reducing agents, with acidulated solutions of ferric sulphate. Linown weights of these three metals in the form of thin foil were introduced into acidified solutions of ferric sulphate, and the amount of reduction effected -part of the liberated hydrogen coming off as gas-under varying conditions of temperature, amount of free acid, and strength of the ferric solution, was determined. Tho reatts obtained showed that the reduction effected when a given mass of zinc dissolves in an acidified solution of ferric sulphato increases with increase of temperature, other conditions being the same. Provided a sufficiency of acid to dissolve the zinc be present, the maximum reducing action is obtained by concentrating the ferric sulphate solution, and diminishing the amount of free acid.

When magnesium is employed the reduction effected is searcely one fourth of that for zinc, while the time required for solution is comparatively very stort; by diminishing the quantity of free soid the amount of reduction effected is increased. The diminution in the rate of solution with a decrease in the quantity of free soid

was found to be much greater in the case of magnesium than in that of zinc; with zinc the rates were approximately in the ratios 1:15:2, and, nnder like conditions with magnesium, the rates were as 1:6:36. When the ferric sulphate is reduced by iron, the rate of solution becomes extremely slow, and the reducing action all pears to decrease with increase of temperature. These reduction experiments, considered as a whole, eeem to be in harmony with the the view that the reducing action of socialled nascent hydrogen is connected with the existence of atoms, as distinguished from molecules of this gas; and that any conditions which tend to prevent the mutual combination of these atoms tend also to increase the amount of reduction effected by the hydrogen.

When phosphorus oxychloride acts upon certain nitrates, it has been found that the ratio between the chlorine and phosphoric pentoxide in the residuo obtained after all action has ceased has a certain definite value. Mills (P.M. [4] 40, 134, and 44, 506), who has studied this reaction for several nitrates, has designated these ratios by the symbol α , or

rather he has taken α as = $\frac{\text{weight of 6hlorine}}{\text{weight of } P_2O_3}$

weight of Cl weight of P_2O_5 . When one nitrate fixes more chlorine, per unit of P_2O_5 , than another nitrate, Mill says that the affinity of the former is greater than that of the latter nitrate; inasmuch as this chlorine fixing action can be measured for several nitrates, the values of a, on Mill's view, represent the 'elective attractions' of the nitrates.

If a be divided by the formula-weights, Z, of the several nitrates, calculated to a uniform mass of NO₃, the following numbers (under o) are obtained:—

•	α	Σ	Q
Thallous nitrate Silver Lead Rubidium Cesium Potassium Sodium Lithium "	 8·78 5·48 5·17 2·38 2·21 1·99 1·70 1·60	265-30 139-94 165-56 147-40 195-01 101-14 86-05 69-00	30·29 31·01 32·02 61·93 88·24 50·82 50·03 42·86

These numbers show that the affinity-coefficients are directly proportional to the formula-weights of the nitrates, and that (with one exception) a and z increase and diminish in regular order. The quotients, o, therefore representathe masses of nitrates which correspond with what Mill calls a 'mit of elective attraction.'

. Chemical Systems of limited action.

When a chemical reaction is expressed symbolically either as B + O = AC + BO or A' + B' = O', it is usually understood that for the complete decomposition of the mass AB all that is necessary is to bring it into suitable contact with the definite mass of the second body on, or that the

nass A if presented to B under proper condisons will unite with it to form o'. I many sons and unite with it to form of. It many themical thanges this is true, at least within the imits of experimental error, and if a fiftient time be given; and it may be said that many processes of quantitative chomical analyses are pased on this assumption. There are, however, many instances known in which the statement loes not hold good. For example, if mol. weights if ethylic alcohol and dectic meid are mixed and heated for some time, say at 100°, only about 66 p.c. of the total action possible takes place, no matter how long the operation is allowed to continue; or again, if mol. woights of iodine and hydregen are heatedeat 440° on a sloeed vessel, even after an indefinitely long period of time there will still exist a certain raction of these elements uncombined. The limitation of these and many other similar shanges appears to be due to the fact that the products of the first action tend, under the conlitions of the experiment to re-form the original substances, and the two reactions proceed simultaneously with different degrees of intensity, lepending upon the masses of material, until a stage is reached at which a state of equilibrium is attained, the first action at this stage being balanced by the second. The mutual action of alcohol and acetic acid would thue be repre-

iented by the equations,

(1) CH₂CO₂H₄ + C₂H₂OH = CH₂CO₂C₂H₃ + H₂O,

(2) CH₃CO₂C₂H₃ + H₂O = CH₃CO₂H + C₂H₃OH.

The theory of such limited actions was formulated by Guldberg and Waage, in 1867 (Etudes sur les Affinités chimiques) and applied by them to the determination of the 'coefficients of affinity' for several restions. These shemiste concluded from the results of their experiments that in a system undergoing change the amount of action in a unit of time between wo or more active bodies in other wor's, the rate of the change-is proportional to the product of the active masses. This same assumption was made by Berthelot in 1862, based on the results of his etherification experiments; and in 1866 Harcourt and Esson showed that lor certain chemical systems the rate of change s proportional to the product of the active masses of the changing bodies. In the abovo statements the 'active masses' of the various bodiee means the number of equivalents of each present in the reacting eystem. There are instances, however in which bodies introduced into a chemical system either accelerate or etard the reaction without themselves undergoing ohango (v. post, p. 714). Guldberg and Waage assume that, in the reaction A + B = A' + B', f the masses of A and B be p and q, then the force tending to produce the change varies as tho product pq, whatever may be the kinds of mater; for two particular substances this forge is equal to κpq , where κ is the 'coefficient of affinity' depending upon the kinds of matter, and probably upon the conditions of the experinent (v. ante and also Affinity, pp. 70975). l'his, however, is not the only ferce acting; there ero others of a secondary character tending to etard or accelerate the formation of A' and B'. Asglecting these secondary forces for the praent, let the masses of A and B be p and q, and the coefficient of affinity for the reverse Vol. I.

action A' + B' = A + B be κ' , then the force tending to re-form A and B equals "p'q". When equili-brium is attained these two forces are equal, or $pq = \kappa' p'q'$, so that if the four quantities $p_1q_1p_1',q_1'$, are determined experimentally, the ratio

of the coefficients of affinity may be found.

a spressed in another way, if Fig. 79, 1991 in the horizontal of the four cubstances in the horizontal of the reaction and xpressed in another way, if P,q,P',q', be the if x be the number of equivalents of P and Q transformed into P and Q when equilibrium is reached, or no further change takes place in the system, all expressed in terms of unit volume, then p=p-x, q=q-x, f'=r'+x, and q'=q'+x; inserting these values, the equation becomes $\kappa(\mathbf{r}-x)(\mathbf{q}-x) = \kappa'(\mathbf{r}'+x)(\mathbf{q}'+x)$.

Such is the simplest representation of the theory of limited actions. The presence, however, of extraneous salts, or even the secondary actions among the four bodies themselvee. doubtlese materially influence the ultimate limit when a state of equilibrium is reached. For instance, if a body x be introduced into the system, Guldberg and Waage assume that the force produced by the action between x and a, and influencing the chango between A and B, is proportional to the product of x and s, or is equal toanx, and they term a the 'coefficient of action.' Assuming that there are coefficients of action between all the four bodies - these coefficients being a. b. c, and d, for a and a', a and B, B and a', and B and B', respectively, and a', b', c', d', for A' and A, B and A, A' an B, and B' and B, respectivelythen the total force for the reaction between A and B will be equal to

 $\kappa pq + a\hat{p}p' + bpq' + cqp' + dqq'$, and that between $\bullet A'$ and B' will be equal to $\kappa' p'q' + a'p'p + b'pq' + c'p'q + d'q'q$.

But that there may be equilibrium these forces must be equal. Writing a-a'=a, $b-b'=\beta$, &c., the equation of equilibrium becomes

 $\kappa pq = \kappa' p'q' + qpp' + \beta pq' + \gamma qp' + \delta qq'.$ If it is desired to study the rate at which the reaction progresses, then this rate is assumed to be measured by the difference between the two forces or

$$\frac{dy}{dt} = \mathbf{p}q - \mathbf{x}'p'q' - \alpha pp' - \beta qq' - \gamma p'q - dqq'.$$

Owing to their complicated character, these equations for the limit or the rate of a chomical change are of little value from an experimental point of flew; it would seem scarcely poseible to determine the numerous secondary forces Guldberg and Waage i..troduce into their for-nule. In such a case as the action between barium sulphate and potassium carbonate the secondary actions to be taken into account are between BaSO₄ and BaCO₃, BaSO₄ and K₂SO₄, K₂CO₃ and BaCO₃, K₂CO₃ and K₂SO₄, and between the water and each of the four salts. For a full discussion of this theory in its simpler form applied to experimental results sec Guldberg and Waage, J. pr. [2] 19, 69 (v. also Arrinity, p. 75).

Berthelot and Saint Gilles (A. Ch. [3] 65 385; 66, 1; 68, 225) were the first to make a complete study of the roactions between carbon acide and alcohols, ae regards the influenca exerted by variations of temperature, pressure

amounts of material, and time. They found that these reactione are characterised by three important features: (1) the combination proceeds slowly, with a velocity depending upon the influences to which the system is submitted; (2) the combination is never complete, howevelong the duration of contact; (3) the proportion of ethereal ealt formed under different conditional ways tends towards a limit.

The inverse action limiting the formation of the ethereal salt, via its decomposition by the water formed during the reaction, was found to be much less rapid than the combination. In other words, if two eystems are cuployed—one consisting of ethelic alcohol and acetic acid, the other of ethelic alcohol and acetic acid, the other of ethelic acetate and water—all in equivalent proportions, the first of these will attain the limit of equilibrium more quickly than the second under like conditions. Bertholot (A. Ch. [3] 66, 113) concluded that in the formation of the ethereal salts 'the quantities of acid and alcohol that combine at each instant are proportional to the product C, the reacting masses.' He gave the formula for expressing the rate of formation as $\frac{C_1}{dt} = m \nu \mu \left(1 - \frac{y}{t}\right)^2$, for

equivalent quantities of alcohol and acid, where the state is the limit, which for acetic acid is = 66.5.

According to the theory of mass-action, the rates of formation of ethercal salts, as well as the magnitude of the limits, ought to be incroased by an increase in either the amount of alcohol or of acid. As regards the ultimate limite, this was found to be true by Berthelot and Saint-Gilles, but for the speed of etherification they found that with nequivalents of alcohol and one of acid there was (at loast for part of the course) little or no increase over that for equivalent quantities; in fact, a diminution in the rate was observed. On the other hand, with nequivalents of acid and one of alcohol the rate of etherification was greatly accelerated. The following two tables illustrate these points (A. Ch. [3] 66, 90, 98):—

1 eq. acetic acid + n eq. alcohol. Tomp. 100°.

	n=1, acld=160, limit=100		acid=100, acid=100,		n=5, ac l=100, limit=100	
4 h.	25·8	38·8	44.0	33·8	17·5	19·3
15 ,,	47·4	71·3		53·2	31·3	34·5
33 ,,	60·6	91·1		87·1	72·2-	79·4

1 eq. alcohol + n eq. acid. Tinp. 100°.

1	n=1, acid=100, limit=100		acid = 100, acid = 100,			n=5, acid=100, limit=100	
4 h.	25·8	38·8	47·1	54.9	57·6	59·4	
15 ,,	47·4	71·3	74·4	86.7	96·6	100	
83 ,,	60·6	91·1	79·2	92.5	96·C	100	

The variation produced in the limit, or maximum amount of thereal salt formed, by employing excess of one or other of the conetituents is illustrated by the following tables (A. Ch. [3] 68, 274, 286):—

	+n eqs. alcohol.	n eqs. soft+1 eq. slophel
n /	limit	n limit •
1.0	66·5 p.o.	1.0 66.5 p.c.
1.5	77.9	67 51.9
2.0₹	82.8	•50 • 41.4
2.8	85.6	36 80.6
3.0	88.2	•33 29.3
4.0	90;2	25 22.6
5.4	92.0	•18 17-1
12.0	03.9	08 7.8
19.0	95.6″ ⋅	0.05 5.0
50 0 ·0	neutral to litmus.	. 0 .

The action of ir organic acids on alcohols has been investigated by Villiers (A. Ch. [5] 21, 72); but in these processee eccondary reactions that are liable to occur complicate matters somewhat. With a given alcohol, the speeds of etherification of the acids III, HBr, HCl, and H2SO4, were found to be widely different. Hi etherifies more quickly than HBr, and each more quickly than acctic acid; whoreas HCl acts with extreme slowness, oven much there slowly than acetic acid. H.SO, etherifies almost immediately under ordinary conditions but the epeed is diminished by dilution, as well as my lowering the temperature. The otherification limite as 100° are different for the three hydracids, and are greater than the corresponding limits at lower temperatures. The limits also depend upon the proportion of water which existe in the initial mixture, but while the limit diminishes in the case of organic acids in a continuous manner as the water increases, without actually becoming nil, the etherification by hydracids ceases completely with a certain dilution, and this limit of dilution is not fixed but rises rapidly as the temperature rises. With H2SO, the ctherification is completely stopped with a certain proportion of water, but, contrary to what occurs with the nydracids, increasing the temperature to 100° doos not cause the reaction to take place. From a consideration of the work of Berthelot and Saint-Gilles on the rate and conditions limiting the etherification of alcohols by Organic acids, it would seem natural to concludo that the application of the methods emplayed by these chemists to the various cases of isomerism among alcohols and acids would yield important results relating to the structure of euch bodies. For the purpose of discovering whether any relation exists between the rate and limit of etherification and the isomeric structure of either of the two active bodies taking part in the reaction Menschutkin (A. Ch. [5] 20, 289; 23, 14; J. pr. [2] 24, 49; 25, 193) has made an elaborate etudy of the action of organic acids on alcohols. In order to render all the results comparable with each other it was necessary to assume two standards for reference, one for tho alcohols and another for the acide. Methylio alcohol was chosen as the standard alcohol; and formic acid as the etandard acid. The two charactoristice chosen for measurement were (1) the initial speed of etherification, or the amount of action that takes place in the first hour, and (2) the final limit of the procese; these Menschutkin terms the 'etherification-data.' For the 'methylic-acetic' system Menechutkin took the limit as equal to 100; that is to eay, out of equal numbers of molecules of methylic

alcohol and acetic acid (in this case 124) only 100 molecules were converted into methylic acetate when the system reached a tate of equilibriums of these 100 molecules, 30 were formed during the first hole of action.

The following table contains the 'etherification-data' for the primary alcohols employed:—

Alcohol	•		_	Speed	Limit
Methydic, HCH, OH, Ethylic, CH, CH, OH Propylic, C, H, CH, OH Butylie, C, H, CH, OH Octylic, C, H, OH, OH	•	•		80 67·3 66·9 67·4 67·0	100 95·6 96 96·6

The influence of isomerism on etherification among the primary alcohols was investigated for the case of isobutylic alcohol; the data obtained were

Isobutylio, C₃H^s, CH₂OH, speed = 64·61 imit = 96·6. These numbers show that the limit is unaffected, but that there is a smalt decrease in the speed. The unsaturated primary alcohols showed less facility for forming others, the reaction in their case progressing much less rapidly, as the following numbers show:—

Allylic alcohol, C₂H₂CH₂OH, 51:9 85:8 Propargylic alcohol, C₂H₂CH₂OH, 29:5 7 Benzylic alcohol, C₄H₂CH₂OH, 54:6 87:3

For the secondary alcohols, the phenols, and some other alcohols, the following etherification-data were obtained:—

. •	Specd	Limit
Dimethyl carbinel (CH,)cHOH	38.2	86.9
Methylethyl , (CH,)(C,H,)CHOH.	32.5	85.2
Diethyl " (C.II.) CHOH	::4:3	84.2
Isopropylmethyl , (Cil.)(C.lis. CHOI!	27.2	85.2
Isobutylethyl (C.H.)(C.H.)(C.H.)	26-9	3
Hexylmethyl " (CH3)(Call1011	3+1	?
Ethylvinyl (C.H.)(C.H.)CHOIL	2: 3	75-1
Diellyl (C.H.) CHOH .	15 3	72
Rthylphenyl (C.H.)(C.H.)CHOH.	27.2	?
Diphenyl " (C.II.) CiloII	31.6	l è
Phenol "Calla.Oil	2.0	2.4
Paracresol. " Call Cll OH	3.7	13.7
Thymel C.H.CH.C.H.OH	1.4	13.6
a-Naphthel C. II, OII	5_	8.8
Glycol CH,OH.CH,OH.	61.7	77-1
Glycerin , (Cll_OH),CHOH .	52.3	66.2
Erythrite C.H.(OH).	34	57.6
Mannite , C. II (OII)	29.6	38

Menschutkin (J. pr. 25, 193) has also determined the initial speeds and limits for different organic acids, employing one alcohol (is butylic), and taking formic acid as the standard of reference. The following results were obtained:

Acld	Speed	Limit
Acid Acetic C,H,O, Acetic C,H,O, Propion' C,il,U, Butwic C,H,O, Caproic C,H,O, Caproic C,H,O, Caproic C,H,O, Caproic C,H,O, Caproic C,H,O, Phenylacotic C,H,O, Phenylpropionic C,H,O,O, Phenylpropionic C,H,O,O, Phenylpropionic C,H,O,O, Phenylpropionic C,H,O,O,	100 71.9 66.7 53.9 53.6 50.0 69.7	100 104·8 106·9 108·2 108·7 110·3 100·4
Methethacetic O.H.(Gi.,)(Ci.,H.,)CO,H Grotonic O.H.,O.H. Ginnamic O.H.,O.H. Ginnamic O.H.,O.H.,OO,H Trimethacetic O.H.,O. Dimethethacetic O.H.,O. Sorbie O.J.,O.	30·3 19·6 18·7 11·8 4·8 12·9	114 8 112 3 116 3 113 1 115 4 116 3

Acid					Speed	Limit
Benzoic C ₇ H _s O ₈ Vitrobenzoic C ₇ H _s (NO ₈)O ₈ Paratoluylle C ₈ H _s O ₈ Juminic C ₁₈ H ₁₈ O ₈	:	:	:	:	13·9 40·1 10·7 10·1	113-9 114-3 119-1 118-1

From the foregoing numbers it is seen that he rates of etherification of the secondary acids are much less than those of the primary acids, but that the limits show only slight fariations. The speeds of etherification of the tertiary acids are less than those of either the primary or secondary acids, but on the other hand the limits are greater. For a full discussion of the value of etherification-data as a means of determining sconcrism among alcohols and acids see Menschutkin (J. pr. [2] 26, 103; also Z. P. C. 1, 611).

The theory of limited chemical feactions has

The theory of limited chemical feactions has been formulated in a simple mannor by Van 't Hoff (B. 10, 659) for the particular case of etherification, but essentially in the same manner as Guldberg and Wange have done in their general treatment of this chemical problem. If the system initially consists of one equivalent of acetic acid, κ of alcohol, and q of water, then when the quantity ϵ of ether has been formed, there will still remain of acid $1 - \epsilon t$ of alcohol $\kappa - \epsilon$, and of water $q + \epsilon t$ consequently the rate at which ether is still being formed is expressed by $C_1(1-\epsilon)(\kappa - \epsilon)$, and the rate of decomposition of the already formed other by the water by $C_2\epsilon$ $(q + \epsilon)$. When equilibrium is attained these two expressions must be equal, or, $C_1(1-\epsilon)(\kappa - \epsilon) = C_2\epsilon(q + \epsilon)$. For equivalent quantities of acetio acid and ethylic alcohol, or $\kappa = 1$ and q = 0, Bertholot and St. Gilles found the limit to be about 66 6 p.c., or $\epsilon = \frac{2}{3}$. Inserting

these values in the equation, the ratio of the two constant C_1 and C_2 is found; or $C_1 = 4C_2$. The equation now becomes $A(\epsilon, \epsilon)(\kappa, \epsilon) = \epsilon(q + \epsilon)$ from which the maximum quantity of ether capable of being formed when various amounts of alcohol or water are employed can be calculated. For instance, if $\kappa = \infty$, i.e. if the alcohol is unlimited in amount, $\epsilon = 1$, that is, all the acid is changed into ethercal salt; if $q = \infty$, i.e. if the water is unlimited in amount, $\epsilon = 0$, or no ether is formed. These results are merely the extreme cases of what experiments have proved to be true between those limits of κ and q which have been tried.

Formation of Acclanitide.—In a study of the formation of acctanitide, according to the equation C₆H₅NH₂ + C₅H₄O₇ = C₆H₅(C₅H₅O)HN + H₂O, Menschulds P(J. 70°. 26, 208) found that, although in the processes of etheritication the final limit of the reaction attained after an indefinitely long interval of time is practically uninfluenced by change of temps as fur, in this example the limit is materially decreased as the temperature increases. The following results show this decreases—

Т	cm f .	Limit.
7	00 _e	85.05 p.c.
1	250	83.11
1	35°	82.39
1	450	81.22
1	55⊶	79.6

Another remarkable fact was noticed in this

reaction, and one which is apparently at variance with most experiments relating to the action of mass (see Berthelot's etherification experiment, ante). In any chemical system undergoing chang, comprising two or more active bodies, the rate of change is generally accelerated (v. p. 741) by an increase in the amount of any of this active bodies, and this increase in the rate is more or less proportional to the quantity factive substance added. But in the formation of acetanilido, with a constant amount of acetic acid, an increase in the quantity of aniline tetards the action, according to Menchutkin, although the final limit is increased as the theory of mass action requires. The numbers under 'speed' vicwing this fact represent the amount of action after 15 mins.

Molecules aniling with one mol. acid	Speed	g.	Limit
1 2 3 4 8	34·71 28·71 23·45 ————————————————————————————————————	•	79·68 91·65 94·61 96·17 97·22

However, when the aniline remains constant and the acetic acid is increased, the law of massaction appears in the normal way (v. also Arrinttr, p. 85).

Molecules acid with one mel, aniline	Speed	Limit
1	34·71	79:68
2	57·30	96:88
4	78·08	99:80

Division of a base between two acids.

When a mixture of two acids acts on a base, or two bases act on one acid, the two acids in the first caso being more than sufficient to combine with the base, or the two bases in the second case with the single acid, it is usually granted that the base divides itself between the two acids or the acid between the two bases in definite ratios. Or if an acid acts upon a salt in solution, as nitrie acid on potassio sulphate, a definite amount of change takes place regulting in this iustance in the formation of potassic nitrate and sulphuric acid. If the ratios in which such divisions occur were known they might afford measures of the relative affinities of the acting bodies for the particular conditions of the experiments. Such ratios have been determined for a great many acids by Ostwald, with most important results. (For an account of this work v. Affinity.)

Pattison Muir (C. J. 33, 27; 35, 311; 36, 60) has studied the conditions affecting the equilibrium of certain chemical systems wherein pps are formed, with the view of determining the relationship between the concentrations of the solutions, the ratios between the active bodies, and the influence of heat on the equilibrium ratios. An investigation some that similar to this was conducted by Marris (A. 213, 233).

Fractional Precipitation.

It has been shown (ante; and v. Affinity) that if a mixture of two acids is allowed to

act upd a single base, or of two bases on a single gild, the ratio in which the base divides itself between the two acids, or the acids between the two bases, depends upon the relative quantities or masses of the materials in the system, as well as upon the strength of the affinities acting between the saveral bodies. In like manner, if a pptant, is added to a solution, containing two or more salts of different, metals, the mass of the pptant, being less than is required for complete ppn. of all the salts-in-the solution (being,

say, the of the total necessary) then the ratios

of the quantities of the salts decomposed—or of the hydrates, carbonates, &c. formed—depends on (i) the relative masses of the substances in golution, (ii) the relative affinities of the salts or the basic powers of the oxides with reference to the pptant, and also (iii) on the fraction of the total material that is ppd.

This highly interesting subject of fractional ppn. has been as yet investigated but to a very slight extent; it would, however, seem to promise in the future a fertile field for the determination of what might be called the relative basic powers of different oxides or hydrates. If, for example, a solution contains two salts of different metals, the basic powers of whose oxides are different, and if a small faction is ppd. (say as hydrate), there will be a tendency on the part of the less basic material to accumulate in the pp. in preference to the more basic, and this tendency will be greater as the difference between the basic powers is greater. If the basic powers differ but slightly, then the increase in the ratio of the less to the more basic material will progress very slowly by repeated application of the process of fractional ppn. If in the extreme case no such difference exists under the conditions of the experiment as regards temperature and quality of the pptant. (the ratio of the basic powers may and probably does vary with the temperature), then the ratio of the Tvo materials in the small pp. will be the same as that in the original solution, and consequently, however frequently the process may be repeated on each fraction formed, no separation will be offected.

At the present time there are a number of elements known belonging to the earths, for the separation of which the only method that has yet been discovered is that of fractional ppn., or fractional fusion; in both cases the separation depends on the differences of the basic powers of the various bodies. Such, for instance, is the separation of the three elements, samarium, didymium, and lanthanum. from each other; or holmium, thulium, and erbium; or again terbium from yttrium. These separations are so extremely tedious, requiring the application of fractional ppn. to be repeated a very great number of times with but relatively infinitesimal yields of finally pure material, that it is evident that the differences in basic powers must be extremely small, more particularly in the cases of samaria-didymia, yttriaterbia, and holinia-thulia. This process for effecting the separation of these earths is repelered all the more uncertain and difficult owing to the want of facts drawn from the study of fractional pon, of other bodies bearing upon

should be conducted (v. Earths).

Chirpfski (A. Suppl. 4, 226; J. 1866. 12) has investigated the subject of fractional ppn. for the case of magnesium and calcium thlorides by phesphoric acid. This chemist employed solutions containing the two salts in varied proportione; to these solutions he added a constant quantity of pheaphoric acid insufficient for complete ppn., then ammehia was added, and he attermined the amounts of calcic and magnssie oxides in the pps. The composition of the pps. was found to vary with the ratio of the amounts of calcic and magnesic chlorides in the solutions, but to be nearly independent of the quantity of water used for dilution. By increasing the amount of calcio chloride in the solution, the magnesic chloride remaining constant, it was found that the lime passed into of magnesia decreased; with the calcium salt constant, the magnesic chloride being increased, the reverse occurred, but to a less marked degree. These variations took place in a regular manner as the composition of the solutions varied.

Mills, in conjunction with others (P. M. [5] 13, 169, 177; and Pr. 29, 181), has studied the fractional ppn., by means of sodium hydrate or carbonate, of seeral sulphates, taken in pairs under varying conditions of mass, with the visw of determining the relative facility with which one sulphate is decomposed in pressure of another when an insufficiency of a

pptant. is addedeto the solution.

In one set of experiments in which nickel and manganese sulphates whe employed, the following numbers were obtained; each solution contained 1 gram of material made up to 100 c.c., and 10 c.c. of a solution of Na CO, (.5715 gram Na CO3) were added:-

NISO.	MnSO.	NiSO. ppd.	MnSO, ppd.	Temp. C.º
·1 grm.	•9 grm.	.0953	•5850	12.9
.2 ັ	.8 ૅ	·1852	4616	13.6
•3	.7	2799	•3766	12.5
-4	-6	•3588	.2976	13
•5	∙5	.4305	.2450	13.6
•6	●•4	4788	1536	12.8
.7	-3	4991	.1089	17
•8	.2 •	.5584	.0722	17
∙9	·ī	•5841	.0363	15 2

From these numbers it is seen that the ratio of the quantities of material ppd. varies continuously, and in the same manner as the ratio of the amounts of salts employed; with equal masses of the two sulphates in solution the pp. contains much more nickel than manganese; hence it is at once inferred that the basic power of manganous hydrate or oxide is greater than that of nickel, since the less basic a material the greater its tendency to be affected by the pptant.

Extending these experiments performed in a similar manner to mixtures of nickel and cohalt sulphates, but employing sodic hydrate instead of carbonate, it was found that these two salts

the pest conducers under which the process | have almost equal degrees of precipitability; that is to say, if the two salts exist in the solution in calculate amounts they will accumulate in the pp. about equal quantities; or, with varying chantities of material, the ratio of the amounts the two salts ppd. will be approximately qual to the ratio initially in the solution; in the words, the basic powers of the two salts. are about equal. (For the theory of fractional pp in. sec Hood, P. M. 1886.)

Reduction of Oxides.

The conditions that affect the reduction of metallie oxides by hydrogen, earbon monoxide, and carbon, have been examined by Wright and Luff (C. J. 33, 1, 509; 35, 475; 37, 757), the type of the reactions being represented by the equation A + BC = An + C. The results have important practical bearings of metallurgical operations. The temperature at which roduetion commences is a function of (1) the physical conditions of the bodies experimented with, (2) and the elemical nature of the substances. With CO as the reducing agent, the temperature at which action begins in the case of cupric oxide varies from 60° to 146° according to the state of aggregation of the copper oxids; for ferric oxide the temperature ranges between 90% and 220°. The reduction by CO of copper oxide, prepared by ppn., is well marked at temperatures below 100°, but at 100° it becomes very energetie. The initial action of II on copper oxide was found to take place at temperatures ranging between 85° and 172°, and on ferric oxide between 195° and 265°. When carbon was employed as the reducing agent, the temperature of initial action varied not only with the physical nature of the metallic oxide, but also with the quality of the earbon; the temperature limits for copper oxide were 350° and 440°, and for ferric oxide 430° to 450°. By comparing the tempsratures of initial action for a given kind of me-talic oxide, it was invariably found that that reducing agent begins to act at the lowest temperature which has the greatest heat of combustion, so that the heat disturbance during its action has (algebraically) the greatest valus. Thus H always begins to act at a lower temperature that carbon, and CO at a lower temperature than H, as the following table shows for different specimens of metallic oxides :-

•	co	и	Sugar	C from
Cuprie oxide A Cuprous oxide Ferrie oxide A	60° 125 146 110 202	85° 175 172° 155 260 195	390° 4 430 440 390 450 450	350° 350 430 345 430
,, B • .	220	245	450	430

Comparing cupric and ferrie oxides prepared by analogous processes, and therefore pre-sumably in much the same physical state, it was uniformly found that the temperature of initial action of a given reducing agent is lower on oxide of copper than on oxide of fron, as the following numbers show:-

	σo	Ħ	Sugar C	O from	ľ
Oxides pre- pared by pre- oipitation. Condes pre- pared by he at in g Iron {	60° 90 125 202 220	85° 195 175 260 24ŏ	390° 450 430 450 450	350° 430 390 430 430	1

The extension of these experiments to the exides of nickel, cobalt, lead, nianganese, ferrous and ferroso-ferrio oxides, resulted in the following conclusions, among others. Differences in physical state are attended with correlative differences in the temperatures at which the actions of the roduoing agents GO, H, and C, are first manifested. For the several oxides of the same metal the temperature of the initial action of a given reducing agent is sensibly the same unless the differences in physical structure are very nurked. In no case was any exception found to the rule that the temperature of initial action of CO is lower than that of H, and that of H lower than that of C, on the same sample of metallit exide. For a large number of cases the rule holds that the greater (algebraically) the heat production during the occurrence of a reaction the lower is the temperature at which this action is first manifested.

During the investigation of the rates of action of CO and II, it was noticed that in many interces 'chemical induction' manifested itself; i.e. the reducing action of the gas on the metallic exide at a given temperature was at first slight or nil ('period of incubation'), but after a longer or shorter time the reduction commenced and proceeded at an increasing rate, until the retarding influences of the products of the action caused the rate of reduction to cease increasing, and subsequently to diminish. The 'period of incubation,' when measurable, was found to be shorter the higher the temperature.

A similar phenomenon has been observed by Bunsen and Roscoe Mt their investigation of the action of light on a mixture of chlorino and hydrogen (v. INFLUENCE OF LIGHT, post), and it is interesting to note that in a heterogeneous system consisting of a solid oxide and a gas chemical induction should also manifest itself. The question naturally arises whether or not it is a general phenomenon accompanying all chemical changes.

Homogeneous Unlimited Systems.—Considering the simplest chemical system undergoing change, that of a single body either decomposing, like ammonium nitrate when heated, or suffering polecular rearrangement, as ammonic cyanate into urea, it is evident that unless the products interfere as retarding agents the amount of change in unit of time, that is to say the rate of change, will be proportional at any time to the affount of active substance then existing. When, however, a system comprises two or more active members reacting on each other, such as an alcohol on an acid, or clydric peroxide on an acidulated solution of a soluble godide, the circumstances are much more complicated. The general experiments on the rate of chemical

change, when not limited by inverso action, prove that in such complex systems the rate of change of any one of the members is increased or dimitished by an increase or decease in the quantity of any of the other constituents, and is more or less proportional to such variation. For example, if the system comprises $A_1A_2A_3...A_n$ (different bodies reacting one with the other), the rate at which A_n changes is increased or diminiched by a similar variation in any other member, as A_n . The statement of this law of mass by Mills $\{P^i, M_n, [\delta]\}$ 1) in the words 'no matter what may be the masses of the substances reacting the cutiro mass of each takes part in the paceess,' requires to be limited by the further statement that the law applies only to homogeneous systems in the sense in which these have been before defined. It could not be asserted for instance that the entire mass of the marble in Bojuski and Kajander's experiments affects the rate of action of the acid, or that a hollow sphere of zinc disselves less rapidly in acid than a solid sphere of similar external dimensions.

Berthelot in 1862 (A. Ch.) showed that the rate of reaction of alcohol with accitic acid is proportional to the product of the two active substances. Harcourt and Esson in 1866 (Pr. 14, 470) established several formulæ representing various experimental conditions based on the same hypothesis, but the reaction they employed for verification of the theory (permanganate on oxalic acid) proved to be of so complex a character as to give but imperfect results. These chemists, however, were more successful subsequently (Pr. 15, 262) with the reaction

 $H_2O_2 + 2IH_2 = 2H_2O + I_2$. The theory of Guldberg and Waago relates more particularly to cases of limited action, but in its application to the study of the rate of change the introduction of so many 'coefficients of action' (v. ante, p. 137) renders the equation of little practical use for such investigations (But v. article APPINITY, p. 70). Except in the theory of Guldberg and Waage, the influence o the products of the change either as accelerating or retarding agents is generally overlooked in attempts to formulate chemical action; but it is easy to introduce these effects it an equation to represent the rate of change of a complex system on the hypothesis that the rate is directly pro portional to the product of all the active mem bers and is inversely proportional to the amonn of chemically inactive bodies for fined (v. BETARDA TION OF CHEMICAL CHANGE, p. 744).

In a complex system, consisting of n members undergoing change, let the masses of the initia active bodies be represented by $A_1A_2A_3...A_n$, an let the masses of these bodies that have become changed of rendered chemically inactive up to time t, be represented by $a_1, a_2, a_3, ..., a_n$; then, as cording to this hypothesis, the rate of change t any member of the system, say $A_{n,t}$ is

$$\frac{da_n}{dx^i} = \mu \frac{(A_1 - a_1)(A_2 - a_2) \dots A_n - a_n}{B \pm (\lambda' a_1 + \lambda'' a_2 \dots \lambda^n a_n)} \dots (1)$$

Where μ , and B, are constants, and λ' , λ'' are the retardation or accelerating coefficien of the products or the action, the + or - significant states according as these products all a as retarding or as accelerating agents. Whise of these actions was performed by any specific

product of the primary action could be determined experimentally, by introducing known mass of the body into the system, and com-paring the late of the change with that beeved when no more of the specified body was present than was formed during the primary reaction. Since A is the member of the system whose rate of change is the object of measurement, to the amount that remains unchanged at time t, that is $\lambda_1 - \alpha_n$, be taken as y; onen—if the initial quantities of the other members be $\nu_1, \nu_2, \ldots, \nu_n$ quivalents of A_{α} , $A_{1} = \epsilon_{1}\nu_{1}\lambda_{\alpha}$, $A_{2} = \epsilon_{2}\nu_{2}\lambda_{\alpha}$,, $A_{n} = \epsilon_{n}\nu_{n}\lambda_{\alpha}$, and $a_{1} = \epsilon_{1}a_{n}$, $a_{2} = \epsilon_{2}a_{n}$,, $a_{n} = \epsilon_{n}a_{n}$. Instructing these values in the above equation it becomes becomes

$$-\frac{dy}{dt} = \mu' \frac{y \{(\nu_1 - 1)A + y\} \{(\nu_2 - 1)A + y\} \dots \text{etc.}}{B' \mp y}$$
(2)

In this equation " and " are constants to be determined experimentally, a being the initial value of y; μ' is proportional to the rate and is dependent on the temperature (v. INFLUENCE OF HEAT ON CHEMICAL CHARGE, p. 741).

Numerical examples of this equation for a system comprising the three bodies, ferrous chloride, hydric chloride, and potassic chlorate, •have been given by $\operatorname{Hood}(P, \hat{M}, [5] 20, 414)$, but the solutions he employed were so dilute that the products of the action appeared to influence the rate inappreciably, consequently the term in the equation relating to these effects was neglected, and the equation was taken as:

$$-\frac{dy}{dt} = \mu' y \left(\nu_1 - 1 \right) \lambda + y \left((\nu_2 - 1) \lambda + y \right) \qquad (3)$$
 for the system of three bodies.

It is possible to arrange the experimental conditions in such a way that, neglecting the action of the products, the course of the change may be much simpler than is represented by equation (2). This may be done, (1) by having all the active substances present in very large excess over that one which is made the object of measurement, so that they undergo but slight diminution between the beginning and the finish of the change taking place in the body measured; or (2) by arr saging the constituents so that one or more of them, although taking part in the reaction, remains constant in amount, one constituent only diminishing in value. The equation for the rate of change of one member in cither case would be by (2)

$$\frac{dy}{dt} = -\mu y_{\mathbf{A}_1 \mathbf{A}_2 \dots \mathbf{A}_n}.$$

Where A1, A2, ... An are the masses of the chemically active constituents which remain constant or nearly so; or integrating, $y = Be^{-at}$, a being equal

to μλ, λ₂...λ_n.

Harcourt and Esson (T. 157, 117) proved the trush of this exponential formula for the action between a soluble iodide and hydric peroxide... The fundamental change in this case is represented by $H_2O_2 + 2HI = 2\Pi_1O + I_2$. By the simple device of adding a known constant amount of sodic thiosulphate to the active solution each time the liberated iodina made its appearance, the amount of hydric iodide was kept constant, while the H₂O₂ alone diminished. The successive additions of thiosulphate measured the amount of change of the hydric peroxide (or y), and the internals between each allition, or rather the appearances of free

iodine, measured the times of action. From their experiments relating to the influence of variations f temperature, and variations c. the masses of nc acting substances, Harcourt and Esson conhuded that 'whether the solution contains in Ic.c. 746 millionths of a gram of hydric sulphate or 150 times that quantity, 604 millionths of a gram of KI or 9 times that quantity, or whether HCl or hydric sodic carbonate he substituted for H2SO4, whether the temperature be 0° 2 50°, and whether the portions of change require for their accomplishment intervals of one or two minutes, or intervals of half an hour or an hour, this reaction still conforms to the law that the amount of change is at any moment proportional to the amount of changing substance.

Harcourt and Esson (T. 156, 193) had previously employed the reaction between potassic permanganate and oxalic acid for investigating the laws according to which a chemical change progresses. Although this investigation was not quite successful iu its primary object, it serves well to illustrate the anomalous results that may a obtained by the interfering action of the products formed in a peaction, or by extraneous salts. The reaction under examination may be represented at its beginning and its conclusion by the two sides of the equation:

 $K_2Mn_2O_8 + 3H_2SO_1 + 5H_2C_2O_1$

 $=K_2SO_4 + 2MuSO_4 + 10CO_2 + 8H_2O$. The reaction progresses with moderate rapidity at temperatures easily kept under control. By varying the mass of any one of the constituents a corresponding variation occurs in the rate of oxidation. The influence of H.SO, is shown in. the following table; the reaction was allowed to go on in each case for four minutes, and was then suddenly stopped by the addition of KI, the amount of change that had taken place being obtained by estimating the iodino liberated:-

Mole- nales H ₄ -O ₄	Per cent, change in 4 mins	Molecules H ₂ SO	Per cent. change in 4 min.
2 4 6 8	21·8 36 51·1 63·5	10 12 14 16 22	71.6 77.4 82.0 85.7 92.3

The principal secondary reaction in the oxida-

tion of C.H.O. by K.Mn.O. arises from the decomposition of K.Mn.O. by the MuSO.

(K.Mn.O. + 3MnSO. + 2H.O.

EK.SO. + 2H.SO. + 5MuO.); this reaction influences the z-te of oxidation in a remarkable mannet. With the materials in the proportions of K.Mn.O.:10:12SO.:5H.C.O., it was found that when no manganeus sulphate was added only eight p.c. of chemical change took place in 4 mins, but by gradually increasing the mass of MnSO, the amount of change taking place in this interval of time increased, until it reached 85 p.e. when 3MnSO, was present. Further increase of the MnSO couly slightly altered the rate of exidation. Harcourt and Esson likewise found that by varying the masses of H₂SO₄ and C₂H₂O₄, the K.Mn.O. and MnSO, remaining constant, the percentage of elemical change in a definite time (3 mins.) gradually increased thil it reached a maximum, then diminished to a minimum, and again increased on addition of more H₂C₂O₄ Experiments on the relation between the time of continuance of the action and its amount showed that after a certain interval the course of the change was represented by an hyperbola. The reason of this regularity only occurring after the action had proceeded some time was traced to the double changes, that take place, first between the MinO₂ produced and the C₂H₂O₄. Both changes are, hewever, comparatively slow; but if either of them occurred yery rapidly compared with the other, in presence of equivalent quantities of materials, the whole course of the change would doubtless he represented by an hyperbola.

doubtless be represented by an hyperbola.

Hood (P. M. [5] 6, 371; 8, 121) has studied the rate of oxidation of ferrous sulphate by potassic ohlorate, and the influence exerted on the process by variations (i) in the amounts of acid used and (ii) in the temperature. The

equation for equivalents being

= 3Fe₂(SO₄)₄ + KClO₄ + 3H₂SO₄ = 3Fe₂(SO₄)₅ + KCl + 3H₂O₇ it is ovident that the rate of change will be the product of three factors. The acid being in large excess and KClO₆:6FcSO₄ = ν :1, the rate of change by equation (3) is $\frac{dy}{at} = -\mu By(\nu - 1)A + y$ where B equals

the amount of acid; or $\log_{\epsilon} \frac{y}{(n-1)\lambda + y}$ ρ μ B(n-1)A(c-t); if, however, KClO₂:GFeSO₄ = 1:1,

then $\frac{dy}{dt} = -\mu B y^2$, or $y(a+t) = \frac{I}{\mu B}$. By a series

of determinations of y (c.c. of permanganate) at indefinite intervals of time, the constants in either of these equations (μ n and e, and μ n and e) were found for different conditions of temperature, amount of acid (n), &o., and consequently a measure was obtained of the changes produced in the rate of oxidation by such variations. Hood found that for this reaction both these formulæ hold good, and, as theory indicates, the rate of oxidation, within certain limits, is proportional, to the amount of free acid; as the amount, of acid, however, becomes comparatively very great the oxidation progresses much more rapidly than the acid increases. When HCI replaces H_sSO_s , or order to produce the same rate of oxidation the amounts must be as 36.5:80.

Ostwald (J. pr. 27, 1) has studied the interesting reaction R.CONH₂ + H₂O = R.CO.ONH₃ with reference to the accelerating influence acids have upon the rate of the change. This reaction is a striking instance of so called 'pr disposing' affinity, the reaction being a very sow one when water alone is employed. (For details of this investigation, v. the article Application of the others of the change o

The decomposition of the athereal salts, e.g. methylic acetate, by water, affords an example of chemical change somewhat analogous to that of the scotamides. The difference between the two cases is that in the former the water resolves the compound into two ethers, alcohol and acid, whereas in the latter the water is assimilated to form a more complex compound. The presence of acids greatly accelerate the decomposition of the ethereal salts, as is the case with the acetamides; the felations between speed of action and

quality of acid have been investigated by Ostweld (J. pr. 22, 449), v. Affinity.

RETABLATION AND ACCELERATION OF CHEMICAL CHANGES.—In the reaction that takes place when an alcohol and an organic acid are mixed, the amount of change is limited by the inverse action that arises between the products of the change, ethereal salt and water, which inverse action tends to the re-formation of the original alcohol and acid; it is consequently evident that the rate at which the ethorification progresses is retarded by this inverse action. In like manner if BaSO, is acted on by K_CO, the rate of the decomposition is retaided by the inverse action that occurs between the BaCO, and K_SO, which results in the formation of the original bodies.

The same may be said as regards the rate of all those reactions which are limited in oxtent by

inverse chemical changes.

There is, however, another kind of retardation possible, not arising from any secondary chemical changes taking place in the system, but of a purely physical origin. If in a homogeneous system undergoing change, such for instance as is represented by the equation A + B = AB, the chemically active bodies be considered to be in a state of continual motios, the rate of formation of AB will be proportional to the number of impaets between the A's and B's in a unit of time. It is conceivable then that if the molecules AB are not removed from the splice of action their mere presence will hamper the movements of the remaining a's and B's, and by so doing will diminish the number of impacts between them in a unit of time, that is to say, will retard their rate of combination. That retardation of a chemical change does arise by the addition of a quantity of one of the products has been shown to be true in several instances; but whether the effects are to be interpreted on a physical basis, as is done here, or on a chemical basis, cannot be decided with certainty until much more experimental evidence has been obtained. The study of the influence of elemically inactive bodies on systems undergoing change, that is to say If bodies which probably do not take part chemically in the reactions, forms a wide field for research; and there is no dowbt that the results obtained will have an important bearing on chemical science considered in its dynamical at pect.

An acceleration in the fate of a chemical change may be brought about byan increase in the amount of any one of the active constituents of the system; such an acceleration, as has been already shown, is easily explained by the law of mass-action, viz. that the total mass of each constituent takes part in the reaction.

There are instances, however, somewhat more difficult of explanation, such as the inversion of cane angars or the decomposition of methylic accetate, by acids, wherein the additions of an acid merely accelerates the change, the mass of the acid remaining the same at the finish as at the beginning of the reaction. The tendency to undergo change in these instances is merely increased by the presence of the acid, and this tendency, measured by the speed of the change, is dependent on the character of the acid employed (v. Ostwald's experiments defailed in Affinity, p. 79). The difficulties that are here

that arise in the consideration of so-called 'con-

tact actions' or catalysis.

Guldberg and Waage (Etudes), in their investigation of the rate of production of hydrogen by the mutual action of metal, and acids, found that the presence of salts in the acid solution exercised a remarkable influence on the speed, some salts accelerating, others retarding, the reaction; the salts themselves remaining unaltered. Mills and Walton (Pr. 28, 268) observed an acceleration in the rate of formation of ammonia from potassic nitrate and zine amalgam by the addition of either K,SO, or Na,SO, the increase of speed being practically the same for equal masses of the two sulphates. If a dilute acidulated solution of ferrous sulphate is oxidised by potassic chlorate at the ordinary temperature, these two bodies being present in equivalent quantities, and the free acid (H,SO₄) being much in excess, the rate of the oxidation (v. ante) is expressed by the equation $\frac{dy}{dt} = -\frac{y^2}{b}$, or

y(a+t)=b, where t is time in minutes, and y is c.c. permanganate equivalent to ferrous iron remaining at time t. Since in these equations $\frac{dy}{dt}$ b^{-1} (or the rate of change is inversely proportional to b), by performing two experiments under like conditions of temperature, dilution, amount of acid, of iron, and of chlorate, except that to one of the solutions a known mass of sulphate is added, it is easy to calculate the equations, y(a+t)=b, for each of the systems; and, by comparing the two values for b, to get a measure of the retarding action of the particular sulphate employed. In other words, the time required to exidise the iron from y' to y'' is proportional to b, and if this time for the blank experiment be taken as 1-0 minutes, the value of $\frac{100b'}{b}$ (where b corresponds to the blank

and b' to the retarded experiment) gives the number of minutes required to perform the same amount of oxidation in the presence of the added sulphate. The annexed table contains the results obtained by Hood (P. M. [5] 13, 419) in studying the retardation of various sulphates in the above manner; the temperature being 21°C. in each experiment.

The numbers show that the retardation occasioned by the presence of a chemically inactive salt in the system employed is proportional to the mass of the salt added, and that some salts of analogous character produce for equal masses the same retarding effect. Thus the potassium, sodium, and ammonium sulphytes each produce a retardation of about 10 p.s. per gram, and the two alums about 6 p.e. per gram. The differences in the effects of magnesium and zinc sulphates are, however, too great to allow of their being classed together as analogous salts from a dynamical point of view with reference to this particular case of retardation.

Considering the alkan sniphates and the alums, it is clear that, since equal masses of the several members of each group produce the same effect, the retardation produced by a molecule of one of the salts is proportional to its

	X,	50.	Na,	50 .	(1)	Н,),SO.
Weight of salt	1005'	Per oent. retar- dation for 1 gram	100½	Per cent. retar- dation for 1 gram	100)&'	Per cent. retar-dation for 1 gram
2 grams . 3 " . 4 " . 5 " . 6 " . 8 " . 10 "	120·3 130·8 143·1 153·4 167·1 195·8 221·7	10·1 10·2 10·8 10·7 11·2 11·9 12·2	120·6 131·6 113·3 119·5 166·5 190·5 212·4	10·3 10·5 10·8 10·4 11·1 11·3 11·2	120 13: 14: 15: 16: 19: 21:	2·4 4·9 6· 9 3·2	10·4 10·8 11·3 10·9 11·0 11·6 11·4
Weight of salt	100/	y r	er cent. etarda- ion for l grain	100 b	Π ₄)Α	P	o.). etarda- ion for gram
2 grams . 3 " 4 " 5 " 6 " 8 " 10 ",	112* 118* 124* 129* 138* 146* 155*	1 7 6 1 2	6·1 6·0 6·2 5·9 6·3 5·8 5·5	111° 118° 124° 129° 137° 141° 153°	3 4 7 4 5		5·9 6·1 0·1 5·9 6·2 5·6 5·3
		Mgs0.		1	Zn	50.	
Weight of salt	1000	y r	er cent. etarda- ion for l gram	100	<i>b'</i>	Ę	er cent. etarda- ion for gram
2 grains	1117 123- 128- 134 140- 150- 161-	9	7:4 7:8 7:1 6:8 6:8 6:4 6:1	110 ¹ 115 117 123 127 132 138	7	-	5·0 5·0 4·4 4·6 4·5 4·1 8·3

mass; ir other words, the greater the mass of a molecul. the more it retards the rate of the chemical change.

Judging by these facts, it would seem not improbable that chemical bodies might be cla. sifica on a dynamical basis as regards their retardation-effects. With this aim, Hood N. M. [5] 20, 444) has continued these experiments in relation to soluble chlorides, but the anomalous results obtained do not justify the assumption started with. The results show that the sulphates of the alkali metals in the oxidation of irrous chloride by KClO, preduce an equal retarling effect, about 17 p.e. per gram, but that MgSO, ZuSO,, and CdSO, (although usually classed together from a statical point of view) differ greatly in their retardation-effects; the results for the arst two sulphates are in about the ame ratio as in previous experiments with ferrous sulphate instead of ferrous chloride. Experiments with MgSO, showed that these 'retardation coefficients' vary slightly with inerease of temperature.

An examination of the effects of chlorides on this reaction showed strikingly anomalous results; no two chlorides gave the same retardation-effect. Sodic chloride practically produced no effect on the speed of the exidation, while zine, magnesium, and cadmium chlorides produced an acceleration. It seems difficult to reconcile these results with a theory that should account for the action of a chemically inactive

alt in a system undergoing change on the basis f molecular interference with the movements f the changing constituents of the system. It nay be, however, that the presence of such xtraneous bodies induces secondary reactions a the system which occasion either an acceleraion or a retardation of the principal change.

·INFLUENCE OF HEAT .- The study of the inuence exerted by heat on chemical changes, as llustrated by the phenomena of dissociation, nd by such phenomena as occur when phos-horus or sulphur are heated, forms an imortant factor in the vast problem of chemical ction. Starting with the fundamental notions, hat heat is a four of energy, and that all aternal material phenomena compriso two actor matter and motion, which it is the aim of ohemistry to investigate, with the molecular heory of the constitution of matter for a basis, t is evident that the relations between the action of heat and chemical change will be most idvantageously studied by examining in what way the rates and the limits of chemical changes becurring in homogeneous systems, are affected by heat, and by determining the influence exerted on such changes by extraneous bodies.

What is already known regalling the influence of heat in bringing about chemical combinations and decompositions would seem to indicate that such action is of a twofold charactor, more particularly in systems possessing free mobility either in the gaseous or liquid states; for, besides accelerating the motions of translation of the molecules of the system, and so mattivelying the chances of collision in a given time, and consequently increasing the rate at which the change takes place, the action of heat also accelerates the rate of chango by diminishing the stabilities of the reacting molocules, and thus increasing their tendency to undergo change. Thus, representing the molecules of gaseous iodino and hydrogen respectively as I₂ and II₂, the amount of HI produced in unit time at a given temperature, by the nutual action of H2 and I2, will depend upon the number of molecular collisions, the velocities of translation of the molecules (or the temperature of the gas), as well as upon the facility with which the molecules become separated in 211 and 21. It is eas; to understand in this way why there are definite limits of temperature within which chemical changes take place; and how some reactions which occur very rapidly at one temperature may be prevented by sufficiently cooling the reacting bodies.

Although as a general rule the action of heat is such as to induce chemical cox bindition at moderately low temperatures and decomposition at higher temperatures, instances are known of bodies being more stable, under certain conditions at high, than at low temperatures. Troost and fixutefeuille (C. R. 73, 443; 84, 946) h..vo shown that by passing SiCl, over strongly heated silicon the latter is volatilised and is again condensed on the cooler parts of the tube. This apparent volatilisation of silicon wall found to be brought about by the formation of Si_Cl_a at the higher temperature, and decomposition of this compound into the original bodies at a lower temperature (FSi_Cl_a=3SiCl_a+Si). The comparatively complex body Si_Cl_a is thus resolved

alt in a system undergoing change on the basis by diminishing the temperature into the come f molecular interference with the movements eparatively simpler bodies SiCl, and Si.

A somewhat analogous reaction is assemed to occur by litte with SeH. (C. R. 74, 930). This chemist says that if scholion is heated with hydrogen in a closed wessel, the amount of SeH, formed increases with increase of temperature up to 520°, but that any further increase in the temperature is accompanied by a decrease in the amount of SeH, formed. If two tubes are heated under like circumstances until the amount of Sell, formed is constant, and one of them is then cooled rapidly while the other is allowed to return gradually to the lower temperature, Ditte says that the second tube contains less Soil, than the first, and so much less as the cooling has been slower. On the other hand, SoH, submitted to the action of heat suffers sensible decomposition even at 150°; above 270° the amount of decomposition gradually decreases until it reaches a minimum at 520°, after which the decomposition continuously increases as temperature rises.

Chemical systems which are limited by reason of inverse actions may be represented in terms of their rates by the equations $\frac{dv_1}{dt} = f(\theta)\psi(\mathbf{A}\mathbf{B})$, and $\frac{dv}{dt} = f'(\theta)\psi(c\nu)$, in which the accelerating influence of temperature is expressed by the functions $f(\theta)$ and $f'(\theta)$, and the absolute rate by the difference, or $\frac{dv}{dt} = f(\theta)\psi(AB) - f'(\theta)\psi(CD)$. When equilibrium is attained, or no further change takes place, $f(\theta)\psi(A'B') - f'(\theta)\psi(C'D') = 0$; A', B', C', D' being the quantities of active, substances that can exist together in stable equilibrium at temperature θ° . If the temperature functions be the same in both cases, then $f(\theta) \{ \psi(A'B') - \psi(e'D') \} = 0$, or $\psi(\lambda' \mathbf{n}') = \psi(\mathbf{c}' \mathbf{n}')$, or the limits are independent of temperature. This would seem to be true between certain limits for the simpler etherification processes, as Berthelot and Saint-Gilles' experiments have shown. Menschutkin, however, has examined certain limited reactions which show a marked variation in the limits

with temperature, and seem to indicate that the ratio of $f(\theta)$ to $f(\theta)$ is not constant. From a study of certain reactions which are not affected by limiting conditions, and of other reactions which are so limited, attempts have been made to determine the decelerating action of heat, that is to say, to determine the form of the temperature-function $f(\theta)$ in the equation $\frac{dy}{dt} = f(\theta)\psi(T.a.b.c...)$.

Hood (P. M. [5] 6, 371), from experiments on the rate of oxidation of ferrous sulphate solution by $KClO_{sq}$ considered that $f(\theta) \propto \theta^2$, or that the rate of oxidation varied as the second power of the temperature; but the experiments were not sufficiently numerous to place this conclusion beyond doubt.

Warder (Am. [3] 203) studied the inflnence of heat on the rate of the action

C.H.O.C.H.3O + No.HO = No.C.H.3O. + C.H.3H.O, in dilute aqueous solutions, the temperature limits being 3.6° and 37.7° . The rosalts obtained agreed well with the formula $(7.5 + a)(62.5^{\circ} - t) = 521.4^{\circ}$; t being temperature, and a the number of gram-equivalents per litre which would (accord-

Ing to the theory of mass) react upon each other par minute in a solution kept of normal strsngth. The formula could be written as $a = \lambda + Bt^2$, indicating the rate to vary nearly as the squars of the tempsraturs. Menschatkin (J, p_1, k_2) 29, 437) employed three somewhat ballogous reactions for the study of this subject; (1) formation of sthylic acetats from acetic acid and ethylic alcohol, (2) formation of acctanilide from acetic acid and anilins, (3) formation of acetamide from acetic acid and ammonia. Molecular quantities of the materials were heated for one hour at different temperature, and the amounts of ether, anilide, and acetamide formed were determined, and talen as measures of the speaks. The following are his results:

Temp.	Ether formed	Temp.	Anilide formed	Temp.	Aceta- mide formed
90.	7.50	82°	6.08	100°	0
102	13.50	ger	8.50	110	1.27
112	19.02	102	14.59	121	4.41
122	21.78	112	21.54	130	9.02
132	82.60	122	30.71	140	21.36
142	40.65	132_	39.91	150	36.96
152	*16·82	142	47.65	152	40.66
162	52.99	152	55.49	155	50.90
172	57.45	162	61.57	160	58.67
182.5	60·99 J	171	66.39	172	72.33
212.5	63.98	182.5	68.87	182.5	78.31
-		212.5	72.19	212.5	82.83

These humbers all agree in this respect, that the difference in the amounts of action during one hour, for equal differences of temperature, gradually increase, pass through a maximum at a definite temperature, and then decrease. rsgards the goneral inferences that might be drawn from these experiments relative to the connection between action of heat and rate of changs, it must be remembered that the reactions labour under the disadvantage of being cases of only limited action, and that the products of the change no doubt retard the principal reaction, and tend to complicate matters. Besides this, its method of allowing the change to proceed in each ease for the same interval of time is objectionable, for at the higher temperatures the amounts of the products formed before the superation of one hour are so very much greater than the amounts formed at the lower temperature that their presence must exercise a considerable retarding influence on the further progress of the reaction up to the time-limit.

Unlike some of the etherification processes the limits of formation of acetanilide and acetamids are influenced very considerably by heat, as the following numbers show:—

Acta	uilide	Acet	ami ič
Temp.	Limit	Temp.	Limit
100°	80.05	125°	•75·10
125	83.11	140	78.18
135	82:39	1::5	81.46
145	81.22	195 182·5	82.82
155	79.68	212.5	84.04
182.5	78.85		
212 ·5	77.75		

In order, to determine the temperature function influencing the rate of a chemical change, Hood (P. M. [5] 20, 323) has again studied the oxidation of ferrous sulphate solution by potassic ohlorate. This reaction is well adapted for work of the kind, as it is completely under control, and can be rendered as quick or as slow as may be desired by altering such conditions as dilution, temperature, amount of free acid, de. The progress of the oxidation can also be followed with the greatest precision by means of permanganate.

Each experimental solution consisted of 5637 gram of iron as ferrous sulphate, and 3-099 grams of free H.SO, madeup to a volume of 250 c.c. To this solution 10 c.c. of a solution of KClO, were added, equal to 2057 gram being the oxidising equivalent of the iron. Fit is a solution, maintained at a constant temperature, 10 e.e. were withdrawn at indefinite intervals of time, and sitrated by permangamate, and from several such observations the constants in the equation y(4+t) = b were calculated: y being e.c. of permangamate, and t being time in minutes.

Since b is inversely proportional to the rate of change, or $\frac{dy}{dt} = -\frac{y^2}{b} = -kf(\theta)y^2$, by comparing the values of b obtained from a series of experiments in which everything remains the same except the temperature, a measure is obtained if the influence of heat on the rate of the oxidation, and consequently a means of finding the probable nature of the temperature-function $f(\theta)$.

The following table contains the results of Hood's experiments; the values for results for equation y(a+t)=0 being the means of several experiments:—

Temp. C.	·	ъ	Ratio $\frac{b_n}{b_{n+1}}$
10°	3::0.8	3327-8	
11	301⋅6	9 3025	1.100
12	274.7	2752.9	1.098
13	250	2503 •	1.029
14	227.5	2282.7	1.096
15	206.6	2055.7	1.110
16	194.3	1920-8	1.070
17	174.2	1333	1.109
18	159	15.7.4	1.098
39	147.1	1452-6	1.086
20`	134.4	1325.4	1.096
21	124	1216.8	1.089
2 2	114.9	1123	1.083
23	102.6	1002.3	1.120
24	94.8	924.5	•1·084_
2,3/	* 89.9	869	1.064
≟ 28	68.5	654.8	1.099
30	58.7	551.2	1.090
32	,50.3	465.3	1.088
•		Mean	1.093

From the numbers under $\frac{b_n}{b_{n+1}}$ it appears that this ratio has as nearly as possible a constant value, the mean of all the experiments being 1093; it would seem, therefore, that for this reaction at least the temperature function has an exponential form, and this the rate of

change may be written $\frac{dy}{dt} = -\mu a^{\theta} y^{\theta}$, θ being the temperature. Calculating the rates of oxidation on this assumption, or $\rho = \mu(1.093)^{\theta}$, where ρ is rate and θ temperature, and comparing them with the rates found by experiment from the values of b, or $\frac{b}{b\theta}$ the rate at 10°C, being taken as unity, the following numbers are obtained:—

Tenip. C.	Rate of oxidation		Calculated rate of oxidation
10°	1.00		
11	110		1 ·09
12	→ 7·21		1.19
e-irs	1.33		r 31
14	1.46		1.43
15	1.62		• 1·56
16	1.73		1.70
17	1.92		1.86
18	2.11		2.04
19	2.29		42.23
20	2.51		2.43
21	2.73	•	2.66
22	2.96		2.91
23	3.32		3.18
24	3.59		3.47
25	3.83		8.80
23	5.08		4.96
80	6.04		5.92
82	7.15		7.07

Lenestigations of many reactions must be made before it can be determined how far it may be true in general that the rate of a chemical change increases in geometrical progression as the temperature varies arithmetically. Lenoino (Etudes sur les Equilibres chimiques, 178) has expressed the opinion that the temperature function is of an exponential form; and this he considers to be correlative with the nature of the internal movements which constitute the temperature of a body.

IMPLUENCE OF LIGHT .- A survey of the principal facts that are known relating to the influence of light on memical changes, or in producing such changes, would seem to indicate the possibility of classifying these climical changes into (1) such as are only induced by the action of light, or light and heat combined, and (2) reactions which, taking place under ordinary conditions in darkness, are accelerated by the action of light. To the first class of actions belong par excellence the photographic processes (of awhich unfortunately so little is known), and such reactions as the combination of H will Cl, the dissociation of HI, or the reduction of ferric oxalate solution. From the experiments of Amato it would seem that some of these typical changes produced by light can only take I lace above a certain limit of temperature, indicating that heat as well as light is necessary. The the second class probably a great many, 'A not all, of the chomical changes that have been studied with reference to their rates, limits &c., will be found to belong when thoy have been examined in this respect, but as yet Little or nothing has been done. As an instance of the latter class of reactions may be cited the oxidation of oxalio

acid by potassic permanganate. Harcourt ana Esson (T. 156, 194) observed that the rate of this oxidation (which under ordinary conditions is modorately fuick) is greatly accelerated in direct snnlight; he amount of this acceleration was not however, determined.

Hydriodic act exposed to sunlight for one month at ordinary temperatures is decomposed to the extent of 80 p.c.; but when this gas is heated night and by for the same length of time at 265° in a dark chamber, scarcely 2 p.c. of the hydrogen is liberated.

The analogy between the chemical effects of light and heat is vory striking: both agencies act in such a way as to break down or simplify chemical structures, as well as to build up complex molecules from simpler constituents. Many instances might he cited to exemplify these statements; such as the disruption of HI into free iodine and hydrogen, the formation of HCl and of COCl, by light; and the formation, and at a higher temperature the dissociation, of HI, H₂Se, H₂O, and numberless other bodies, by heat.

There seems, however, to be one marked difference in the modes of action of heat and light. Whereas, in those chemical changes produced by heat which are termed dissociations or disruptions of molecular structures, a limit is reached depending upon the pressure and temperature to which the system is subjected, in similar changes produced by light there seems to be no limit, but the process goes on until complete decomposition is attained. This is easily understood when it is remembered that in such reactions, under suitable conditions, hoat tends to destroy as well as to re-form the molecular structures; but that when light acts in such a way as to break down complex mole-cules the inverse action has not yet been observed to occur under any conditions. For example free H and I exposed for one month to sunlight suffer no measurable change; but HI in the same interval of time, is decomposed to the extent of 80 p.c. Reactions of a limited, and perhaps reversible, character, induced by light, analogous with the dissociation pheno mena produced by heat, may yot be discovered. Light rays of different refrangibilities induce chemical changes with greatly different intensities, and probably with different effects. Lemoine (C. R. 93, 514) has shown that HI is decomposed with facility in vessels made of blue glass, but is very slowly changed in red glass vessels. For those bodies which heat alone decomposes at low temperatures, the extreme red of the spectrum appears much less efficacious than the extreme violet. In the cases of bodies which are stable at high temperatures, if the time of action be long enough, the red rays ultimately produce the same result as the giolev. Chastaing (A. Cv. [5] 11, 145) concludes that it is not necessary that white light should act more exergetically in a given way than any particular part of the solar spectrum, for it is possible that some rays produce the inverse action of others. The chemical action of the solar appetrum on binary metalloid and metallic compounds ought, he thinks, to be represented by two ourves, one reducing on the side of the violete the other of an oxidising character on the red side of the spectrum, and,

be says, there probably exists a point where would evidently place a limit to so-called instan-photochemical action is nil or equal to that staneous photography.

which takes place in darkness An elaborate study of the influence of light in producing chemical changes was made by Bunsen and Roscoe in 1857, some of the facts which their investigations brought out are of a highly important character. Draper in 1843 had observed that the action of light on a mixture of H and Cl does not begin to show itself instantaneously, and he concluded that the first action of light was to bring about a change in the Cl, probably producing an allo-tropic modification, before combination could take plane between it and hydrogen. Bunsen and Roscoe made this observation by Draper the subject of many experiments and measurements. They considered that whatever may be the cause of the resistance to combination which the gaseons mixture shows for come little time after submitting it to the action of a constant source of light, Proper's assumption is not borne out by facts.

The following measurements exemplify this resistance effect as obtained by these chemists with a ochstant source of light (T. 147, 363):-

	i	
Time in mins.	Observation	Action during 1 min.
0 •	100	
1	100.5	-5
, 2	$102 \cdot 1$	1.6
8	102.6	1 ⋅5
4	103.2	-6
5 🕈	105.3	2.1
6	119.9	14.6
7	139-1	19.2
. 8	170.2	31.1
9	• 200.6	30.4

These numbers show that about 8 mins. exposure is required before the rate of combination reaches a constant maximum. It was found that the time that clapses, from the first insolation nutil the first trace of photochemical induction becomes visible, and until the maximum action a attained varies much according to the experimental conditions. It was also found that the resistance to combination, once overcome by the influence of light, is soon restored when the gaseous mixture is allowed to stand in darkness, but that the increase of the induction from exposure to light takes place much more rapidly than the diminution of the same on darkening. The presence of a foreign gas, or of excess of Cl or H, influences the induction-effect in a remarkable manner. Thus the maximum of the induction of a normal misture was reduced from 100 to 378 by the presence of Ting of hydrogen, and ir the presence of Too and Too of expension of the presence of 100 to 9.7 and 2.7 respectively, and for 100 to 60.2. By insolating the gases eeparatoly no appreciable effect was produced on the induction effect when the gases, were afterwards mixed.

It is interesting to note that if this Archininary resistance to undergo change is a universal law in such actions as a rollbrought about by light in bodies in the fiquid or solid state, it ture, by the pressure of the restring gaseous nary resistance to undergo change is a universal

Marchand (A. Ch. [4] 80, 802) has etudied the influence a light on a mixture of oxalic acid and ferrio chlorido in aqueous solution. Such a eolution placed in darkness suffers no change, but when exposed to light it evolves CO₂ with the reduction of the ferric chloride. Heat alone has no visible effect on the mixture oven at a boiling temperature but if the solution is exposed to solar radiations and is then heated, decomposition takes place with explosive violence. Of the different parts of the greature the blue represents the parts of the spectrum, the blue raymoxercise the most energetic action, even more so than the violet rays. Some highly interesting facts have been noticed by Lemoine (C. J. 97, 1208), bearing on the chemical changes produced by light with the above mixture. He employed a number of thin vertical glass tubes, 15 mm. diameter, each containing 20 e.c. of a mixture of ferric chloride and oxalic acids the solutions were saturated with CO, and contained equivalent quantities of the reacting bodies. The evolved gas was collected over glycerine. The speed of the reaction increased in proportion to the intensity of the light, but for equal intensities of light the speed was at first approximately constant, and only began to slacken when the liquid had disengaged half the possible quantity of gas. If the two reagents are exposed separately to strong smilight and are then mixed, the decomposition goes on much more rapidly than if the mixture is exposed to light before separate insolation. The following numbers illustrate this fact; the measurements, which were made after the anno intervals of exposure, show an acceleration of obout ten p.c. in the latter case :---

Liquids not insolated of 24 52 64 83 3 100 107
Liquids insolated for nine hours 0 24 58 70 91 103 110 117 1 1-11 1-00 1-10 1-11 1-10 1-09 Latins . .

A remarkable point figured in these exporiments was that the addition of water increased the rate of action of the light. This anomalous effect may have been due to the partial docomposition of the ferric salt as well as to the fact that the upper layers of the ferric chloride absorb much of the light and provent it pene-trating far into the liquid.

According to Amato (G. 6,4, 57), many reaction: which are produced by sunlight are not real y due to this agency. Amato considers that light, only acts under certain determinate conditions of temperature, and that consequently there are limits of temperature within which light does not act in a chemical way. 'He found that "mixture of Cl and P, if cooled to -12° could be expe ed to the direct rays of the eun for hours with at combination taking place. In this experimen? care must be taken that the Cl is not exposed to the sun's rays before or ling, as insolation rendere chlorine capable of combining with hydrogen even in the dark.

INFLUENCE OF PRESSURE. - Many substances when subjected to the influence of heat in a closed vessel, such for instance as calcic carbonate, ammonic carbaniate, or paracyanogen, products. When the pressure of the evalued or, certain salts are decomposed when neatedy gases has reached a definite value no further but one phase of the change takes place at a altoration takes place. If, however, the pressure is maintained below this limit, by allewing the gases to escape, complete decomposition rosults. The consideration of the influence of gaseous pressure in such instances belongs to dissociation (q. v.). There are, however, a few chemical changes known, other than those of disconttion, which occur only under considerable pressures, and others again which are prevented, or at least greatly retarded, by pressure.

Cailletet (C. R. 58, 395) found that, sepresenting the amount of action between zinc and HClAq of a definite strength under ordinary atmospheric pressure by 10, the action was reduced to 4.7 ander a pressure of 60 atmospheres, and under 120 atmospheres the amount of action in the same time was only 1. The amount of action between HNO, and CaCO, under pressures of 1 and 150 atmospheres he found to be as 11.09:1.

Beketoff (C. R. 48, 442) reduced colntions of silver nitrate and sulphate, and ammoniacal silver chloride, by hydrogen, under pressure-reactions which do not take place at atmospheric

pressuro.

By mere mechanical preseuro Spring (B. 17, 1218) caused several of the metals, such as copper and lead, to combine with sulphur, and also brought about the formation of many,

CONTACT-ACTIONS. — Catalysis. Cyclical Actions .- Catalysis, or contact-action, is the namo given to a cumerous class of chemical changes that are induced in certain chemical systems by a substance which does not itself undergo any permanent alteration, but which by its mere presence under suitable conditions brings about a re-arrangement among the molecules of the bodies with which it ie placed in contact. The naterial which acts in "his manner, without apparently being affected itself by the changes it nduces, has been termed a oatalytic or contact-

According to the theory of Berzelius, who was the first to study the class of reactions, such bodies are possessed At a peculiar property or power which he termed 'catalytic force,' or the power to bring aboy, chemical changes. Berzelius assumed this catalytic force to be of the character of an electrical force. It seems simpler, however, to regard such actions as being merely manifestations of the same property or power that is exhibited by all forms of matter undergoing chemical change, or the manifestation of the affinities of one kind of maker for another. It is reasonable to suppose that in every chemical system there is a tendency to undergo change of some definite character, such for instance as hydrogen and oxygen to unite, cane styar and water to form glucose, potassio chlorate to give off oxygen, &c. The conditions under which the system exists may be such that the afinities are in a state of stable cquibrium among themselvet. Every system may be regarded as having a weak point, or paint of least resistance, at which an alteration will most easily take place. For instance, in the reduction of certain metallic oxides, the oxides are first reduced to lower oxides and then to the metal; H₂O, &c.

lower temperature, or more easily, than another phase. It would seem probable, therefore, that if a suitable material were introduced into a chemical system, i might so react with certain constituonts of the system as to upset the previous equilibrium to such an extent that what was before merely a tendency to undergo change would become an actual change, beginning at the point which before the Etfoduction of the catalytic agent was the weakest point of the system. The catalytic agent may be regarded as tending to form, with one of the constituents of the system, a compound too unstable to exist under the conditions, which compound immodiately breaks up, leaving the so-called catalytic agent in its original condition, free to react with a fresh portion of the system.

Contact-action would seem to be rather an illchosen term for this class of reactions, since all chemical Combinations timply contact. It is also well known that many soluble salts if placed in contact with insoluble salts or pps. adhero tenaciously to these. A striking inetance of this kind of contact-action is exhibited by metastannic acid. If a small quantity of this powder be chaken np with a highly ferruginoue solution of aluminium sulphate, the ferric oxide in solution is scized upon by the insoluble metastannic acid, leaving a solution of aluminium sulphate in which scarcely a trace of iron can be-

detected.

From the evidence that exists relating to what is strictly known as catalytic action, if a word were necessary to distinguish this kind of change from ordinary chemical reactions, cyclical action or cyclical change would seem to be near the mark.

The instances that are known among gases in which the presence of a body brings about chemical action in an otherwise stable gaseous mixture seem to be explained by assuming that contact action emercly causes a condensation of the gases upon the surface of the material that brings about their union. Faraday (T. 1834. 55) found that if a plate of perfectly clean platinum is brought into a mixture of hydrogen and oxygen, combination of the gases begins to take place, at first slowly, but at a gradually increasing rate, until combination occurs with explosive violence. This combination was considered by Faraday to he due to the condensation of the gases upon the motallie surface, whereby the molecules of oxygen and hydrogen were brought into such close contact that chemical union took place. The presence of small quantities of CO or CS₂ prevents the combination of the oxygen and hydrogen by aid of a platinum surface, although the metal is not found to lose its power if afterwards plunged into a pure mixture of the gases. Small quantities of such gases as H, S or HCl, however, so alter the platinum-surface that the motal is now incapable of effecting the combination of H with O. Other substances such as charcoal, pumice, rock crystal, &c., act in a similar manner to, but less rapidly than, platinum. Platinum also orings about the combination of SO, and O to form SO, of NH, and O to form HNO, and

etermining the vapour density of tertiary mylacetate, found that the dissociation-phenonena exhibited by the vapour of this body are nfluenced in a stilking manner by the presence of many finely divide I substances such as silica, nagnesia, calcium sulphate, a.g. The effects varied with the chemical, as well as with the physical, characters of the substances placed in contact with the vapour. To such a slight extent have these contact actions been studied that it seems as yet impossible to interpret them in the same way as those chemical actions which are here termed cyclical. Taraday's theory for the action of platinum in bringing about the union of daygen and hydrogen by a mere condensation of the gases would seem to be the most resonable explanation in the face of the facts at present known.

Examples of catalytic actions are known among liquids, which may be explained on the among induces, which may be explained on the theory of cyclical cleange; tuch are the evolution of oxygen from a solution of a hypochlorite when warmed with cobaltous or manganic oxide, and the decomposition of hydrogen peroxide by manganess dioxide, finely divided platinum or silver, or, by oxide of silver. The last case is remarkable, for here oxygen is evolved both from the silver oxide and from the hydrogen peroxide; to explain this, and a number of analogous reactions, Brodio (T. 1-10, 759) assumed that atoms of the same body may is an attraction for each other or be in a stato of polarisation. Brodio expressed the

reaction of Ag₂Q with H₂O₂ thus, H₂OO + Ag₂O = $H_2O + \overline{OO} + Ag_2$

The change of cane sugar and water into glucose, and of ethercal salts and mater into acids and alcohols, in the presence of acids which themselves remain _Achanged, are other instances of catalytic action probably of a

yelical character (see AFFINITY, pp. 71 et seq.).

If potassic chlore e is heated alone it melts at about 345°C, and on increasing the temperature to about 370°C. oxygen begins to be produced. Many substances in a fine state of division when mixed with this salt cause an ovolution of oxygen nuch below the temperature at which the chlorate decomposes when heated alone, and without the salt entering into a state of fusion. The substances which facilitate this decomposition do not themselves appear to undergo any chemical changs. It is probable that the theory propounded by Mercer (B. A. 1812. 32) to explain analogous chemical changes is the true one, viz. that the material which facilitates the decomposition has a tendency to pass into a higher state of oxidation, and that an nnstable compound is formed but is decomposed at the temperature of the experiment. On this hypothesis the potassium ohlorate is regarded as being decomposed at the lower temperature by the double effect of heat and the affinity of the contact substance as MnO₂—for the oxygen of the chlorate. Heated by itself, potassium chlorate passes through an intermediate stage. in its decomposition with the formation of perchlorate; this intermediate stage is represented according to some chemists by the equation $2KClO_2 = KClO_4 + KCl + O_{22}$ but it appears to be more concept, expressed by the

Konowadow (B. 17, 1360; 18, 2808), when requation 10KClO, =6KClO, +4KCl+8O (Teed, primining the vapour density of tertiary (C. M. 22, 248). If, however, manganese decide via the chlorate, no perchlorate is no exhibited by the vapour of this body are an argument in favour of Mercer's view, by supposing that KClO, when decomposed by itself posing that KOIO, when decomposed by itself forms KCl and O₃, and that the nascent ozone oxidises a second molecule of chibrate to perchlorate, thereas in the presence of MnO₂. In ligher but tery unstable exide of manganese is formed, and is almost simultaneously decomposed. The oxides which noost markedly facilitate the decomposition of potassium chlorate are as a rnle those the metals of which form soveral oxides. It is a well-known fact that the oxygen prepared from KClO, either by heating the salt alone or mixer with MnO, liberates iodine from an alkaline iodide; this is usually considered to be due to a trace of free chlorine; it day, however, be occas much by a little ozon, that esca decomposition by the MnO, or the KClO,

Contact chemical action, whatever be its true cause, plays a highly important part in several industrial op rations, as the inversion of cane sugar, the conversion of starchy matters into glucose, the decolorisation of sugar solutions by charcoal, and probably in the purification of waters by filtre fon through porous media. Formerly the great industrial processes of ferments. tion in the formation of alcoholic liquors were referred to this cause, but it beems now certain that such changes are phenomena connected with organic life and not with those of unorganiscd matter.

(For a theoretical consideration of catalysis

sec.Mendeléeff, B. 19, 456.)

A compleration of the facts that are known relating to chemical change shows that in the study of the subject not only must the kind o. quality, and the mass, of the reacting matter, be taken into account, but attention must also be given to the intrinsic fg. &cs that come into play, as well as to the votion of molecular or atomic motions. It is not at present so much the relative values of these forces that one desires to know, in whatever way they may be measured, as the circumstance under which the forces act, or are modified in their acti u. It has been seen that al vatomic structures are possessed of rela-tive degrees of stability, as is hown when sub-mitted to the action of physical agencies, or when they play an active part in chemical systems, this stability being due to the interaction of the affinities helding the structure together. These forces or affinities offer different degrees of resistance, to the action of different agencies, and it would's. I to be only by the study of such influ Aces that a rational conception of the nature of chemical action will be arrived at.

That the ultimate constituents of matterthe atoms or molecules - are in continual moton, the interdiffusion of gases, and of salts in solution, seems to prove; and the fact that, in a chemical system undergoing change, such chango is more of less gradual, taking fractions of seconds or while years to be accomplished, seems to offer conclusive proof that the atomic constituents are in a centin pl motion of interdiffusion. But whether or not the charge zoing on in a chemical system is brought abo '> by simple

scetic acid, when mixed in equivalent proportions react upon each other at ordinary temperatures with extreme slowness; in fact, it tries months to accomplish what at 100° requires only minutes, and yet it is highly probable that very many more fullisions occur between the alcohol and the acid molecules than the rate of change would lead us to conclude. It man bo that in these and similar cares the modules of the two constituents of the system must be moving with a definite velocity if chemical action is to occur. But the kinetic theory of gases teaches that in a space of uniform temperature some molecules have high and others low velocities, and that the ratio between the run. For of molecult having high and those having fow velocities vara thyith the temperature; consequently in chemical whange which occur may be but a process of selection among the molecules according to the velocities they possess, those with velocities below a certain limit colliding, but not reacting chemically with, each ther.

As chemical reactions are generally formula lated, the phenomena of change are for the most part at present viewed only in the light of the distribution of certain masses of matter of various kinds, and no cognisance is taken of the changes in the energies of the systems as these pass in m the initial to the final states. In the blank that is at present occupied by the sign '= lie all the real phenomena of the science of matter. Attempts have been made to fill up this blank by the investigation and measurement of the hoat-disturbances that arise when a ohemical system passes from the state reprosented by one side of the equation to that represented by the other side. By virtue of the inherent forces or affinities, as well as by the particular motions of the ultimate particles or atoms of matter, all substances may be looked upon as possessing a certain definite amount of energy, potential as well as kinetical, and consequently as capable of performing a definite amount of work. The tendency of the constituents of a system is invariably towards a state the attainment of which involves a degradation of energy; in other Ards the total energy of the system tends to te a from a higher to a lower level. For the general inferences that have been drawn from the study of thermal phenomena bearing on the applications of the laws of energy to chemical change reference must be made to the section on THERMAL PHENOMENA of the article PHYSICAL METHODS USED IN CHEMISTRY.)

It is much to be desired that a classification of the elements, or, what seems many possible, of their compounds, should be attempted, 2 sed | [5] xviii.).

collisions among the moving molecules cannot report some particular dynamical properties be asserted. For instance, anhydrous alcohol and which should include not only the conception of mass but also the conceptions of time and work; it is evident, however, that the difficulty lies in the kind, of phenomena to be observed and measured; Mills 47. M. [5] 1) has propounded certain ideas regating to chemical phenomena, making motion the basis of the science; and he considers that chemical substances should be valued not for what they are conceived as being, but for what they are capable of doing. Doubt-less, however, the being as well as the doing must be considered together. The masses of various bodies necessary for the performance of nnit of work Mills terms the dynamic equivalents, for the 'bergmannics,' of the respective bodies; these may vary according to the sort of doing, or work, the several substances : .. employed to effect; such as the power of various acids to invert sugar, or to decoripose ethereal salts, the precipitability of salts, the coefficients of diffusion, &c., &c. For many valuable determinations of dynamical effolis of substances in inducing or accelerating chemical changes, see the work of Ostwald. For a full account of this work v. the article Affinity. (In connexion with this article, v. the articles: AFFINITY; ALLOTROPY; CHEMICAL AND PHYSICAL PROPERTIES OF BODIES, CONNEXIONS BETWEEN; COMBINATION, CHEMICAL; DISSOCIATION; Egg ALBRIUM, CHEMI-CAL; ISOMERISM.)

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